# **EXHIBIT 3**



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## (12) United States Patent

## Ciaramella et al.

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## (54) BETACORONAVIRUS MRNA VACCINE

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## (58) Field of Classification Search

None

See application file for complete search history.

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## (57) ABSTRACT

The disclosure relates to respiratory virus ribonucleic acid (RNA) vaccines and combination vaccines, as well as methods of using the vaccines and compositions comprising the vaccines.

## 21 Claims, 24 Drawing Sheets

Specification includes a Sequence Listing.

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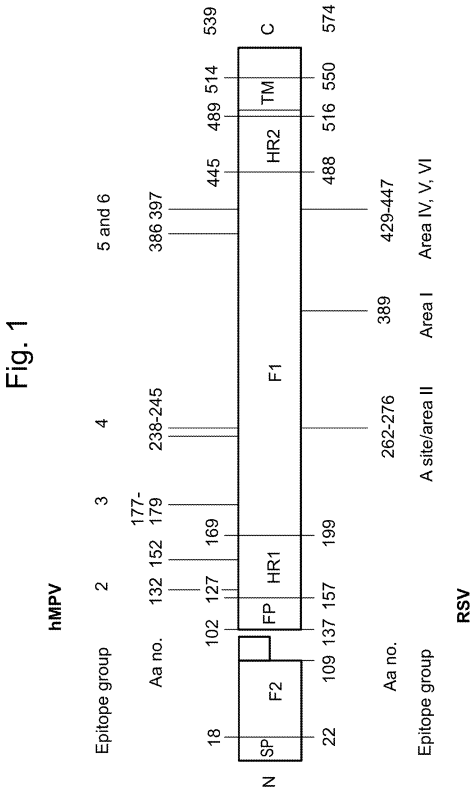
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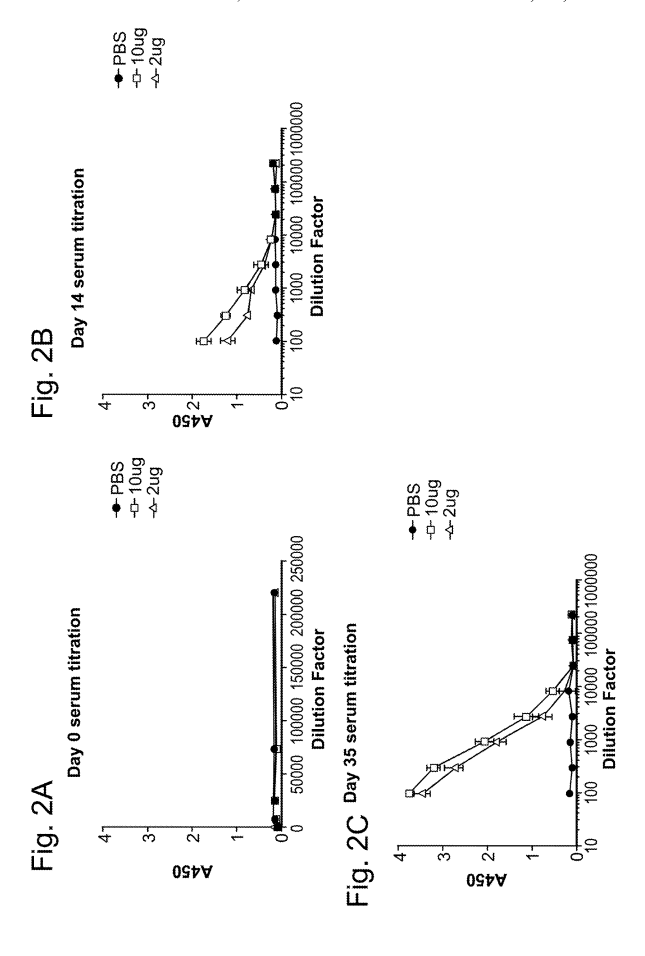
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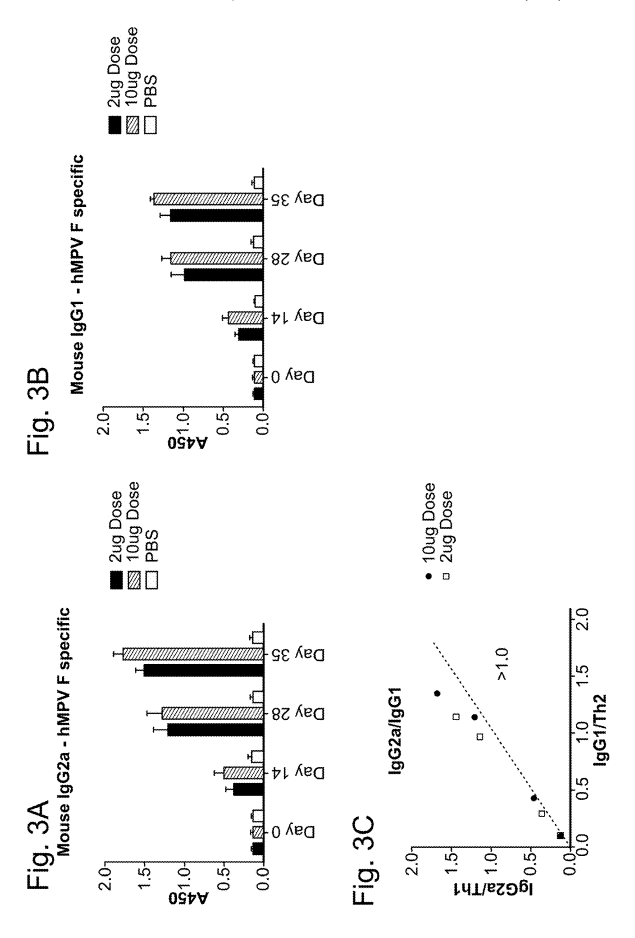
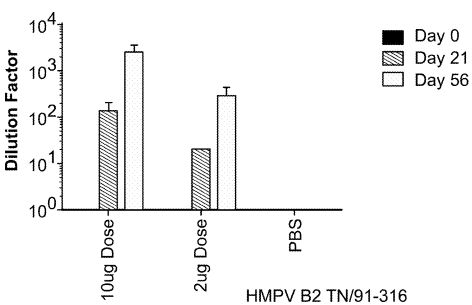
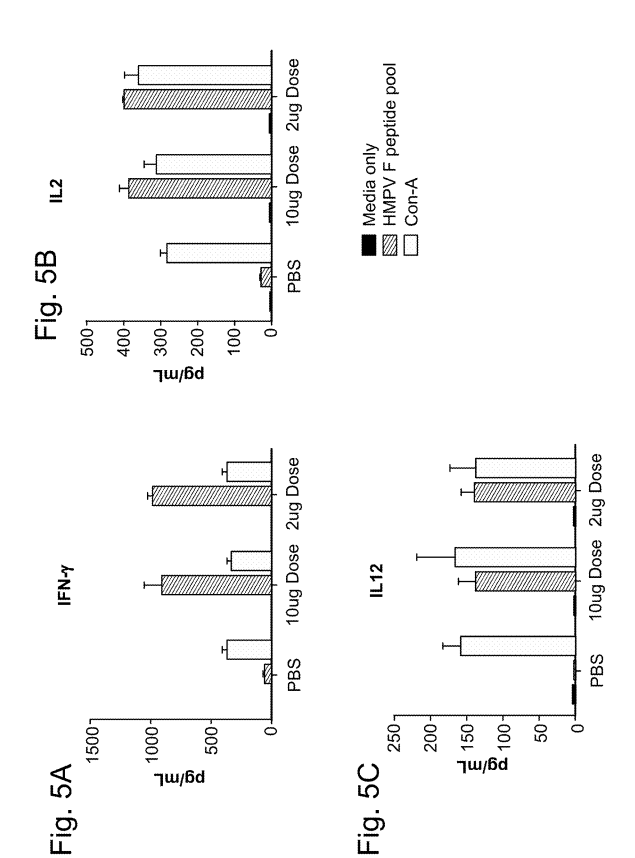
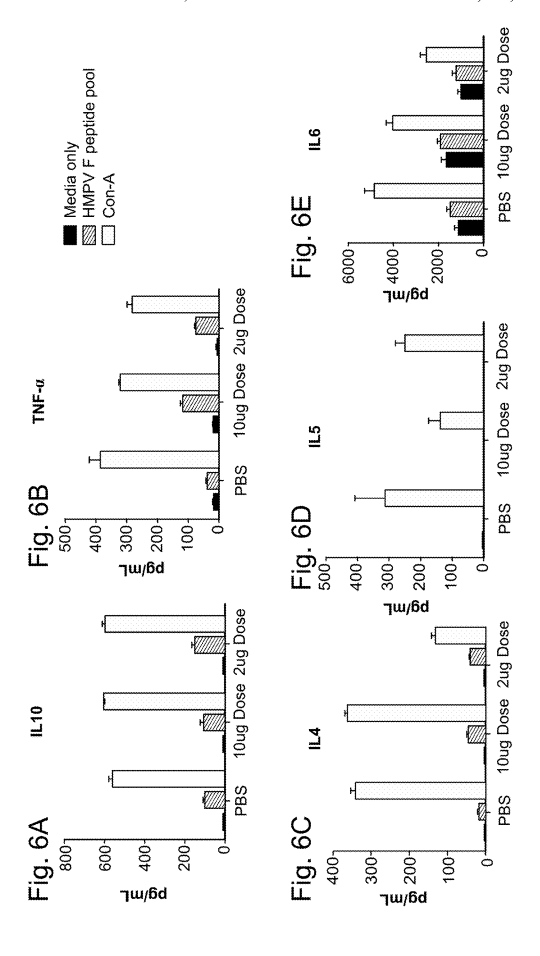


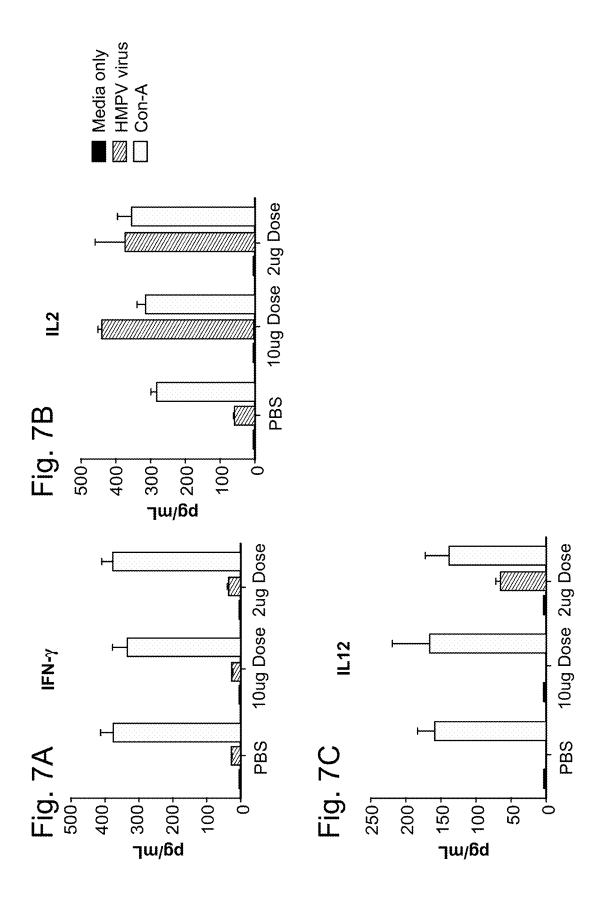
Fig. 4











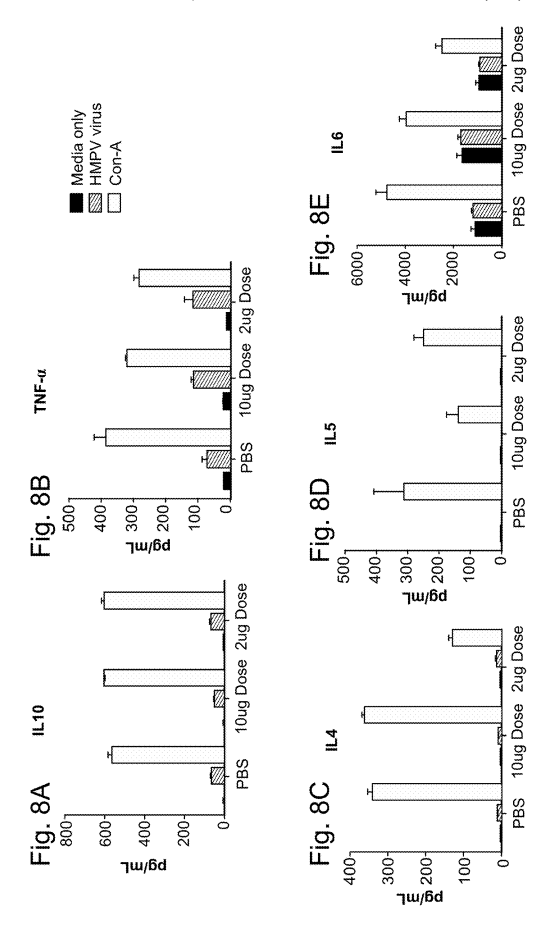


Fig. 9A

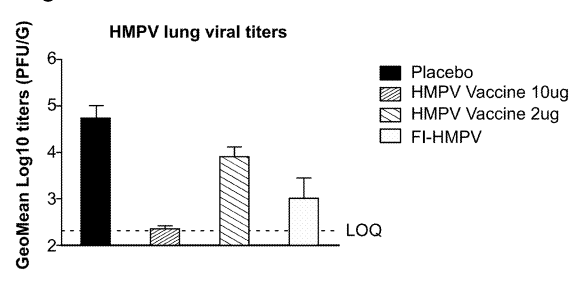
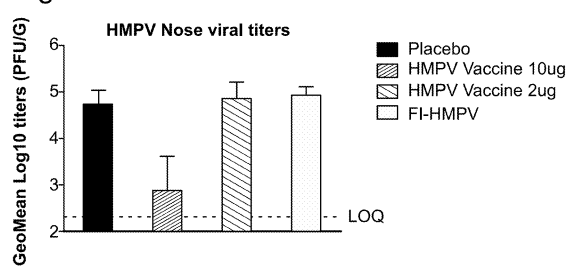


Fig. 9B



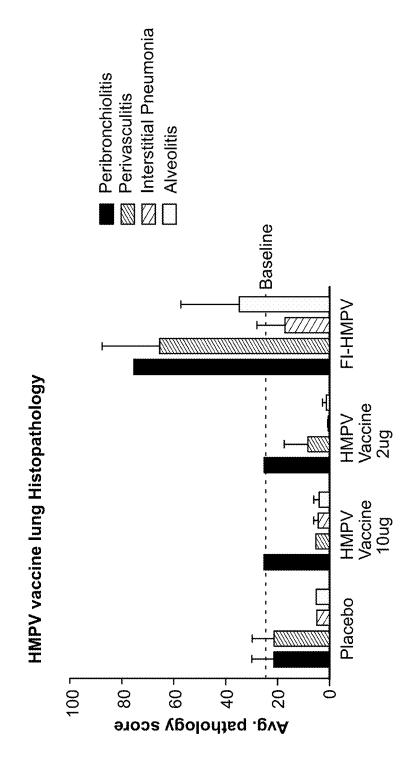
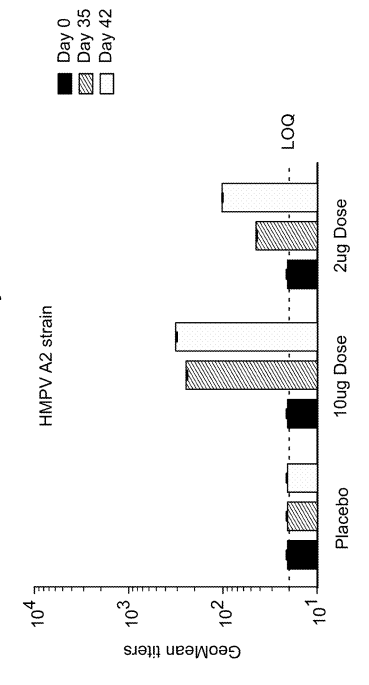
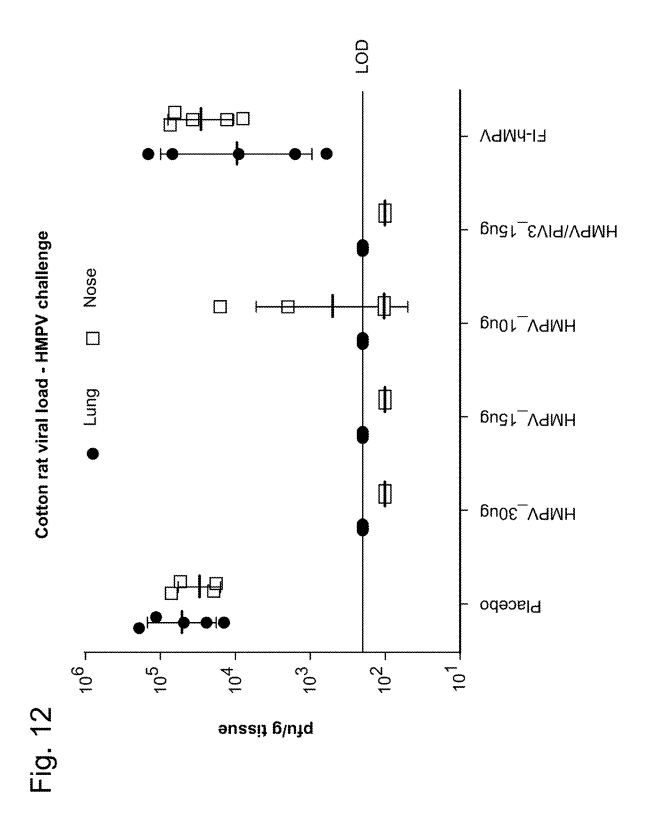


Fig. 10

Fig. 11
HMPV neutralization antibody titers in cotton rats





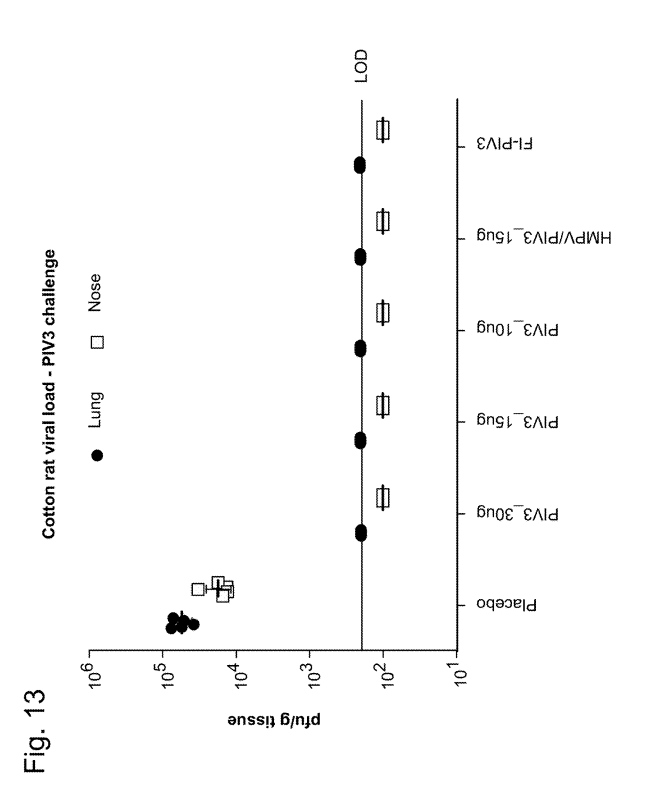
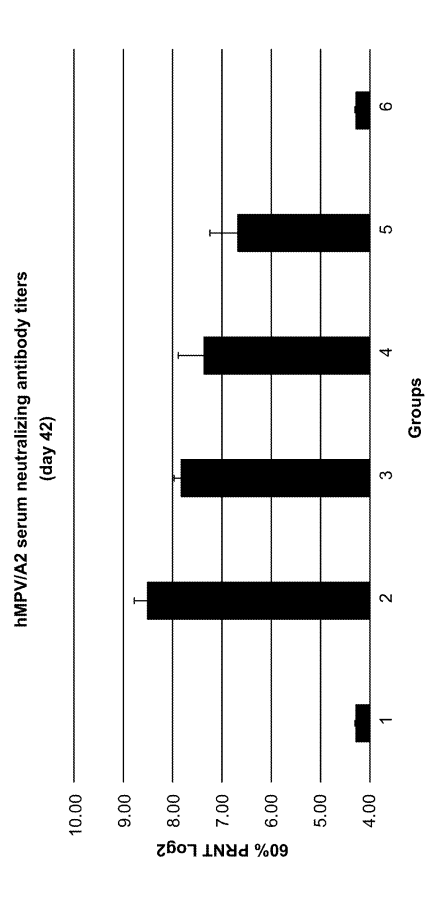


Fig. 14



8.00

7.00

6.00

**60% PRNT Log2** 

5.00

12

7

9

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 $\infty$ 

Groups

Fig. 16 Cotton rat lung histopathology

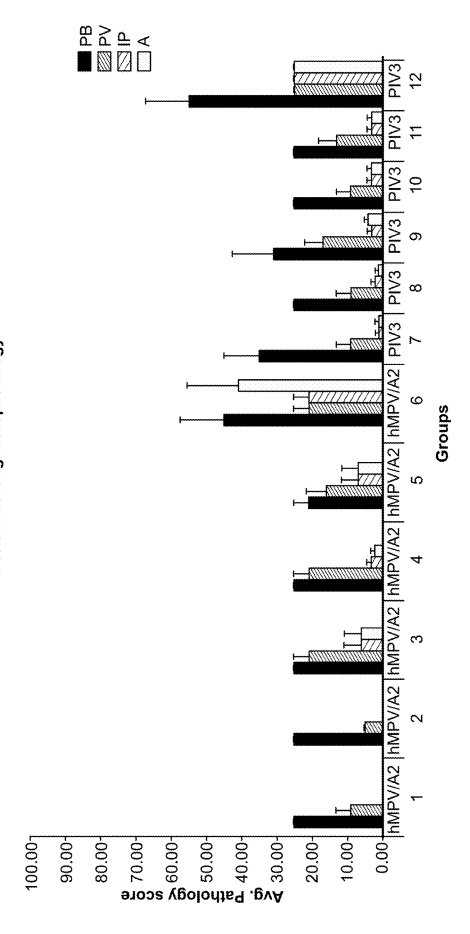
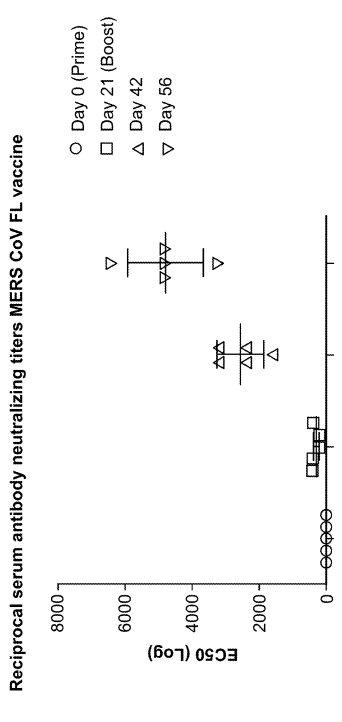


Fig. 17



FL Spike protein S2 spike protein Reciprocal serum antibody neutralizing titers MERS CoV vaccine 99 Day 21(Boost) 0 (Prime) 10000₁ 1000= 100-EC20 (Fod)

Fig. 19/

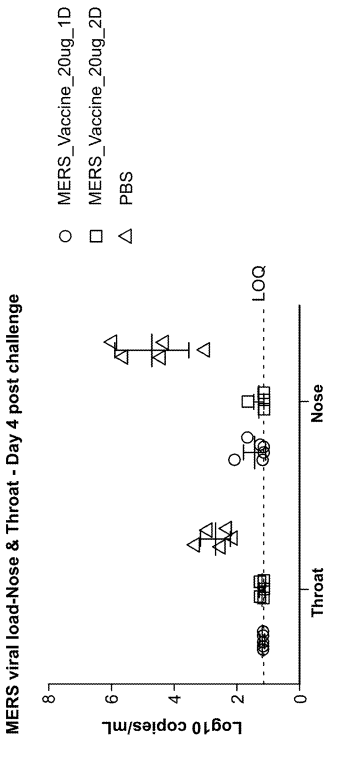
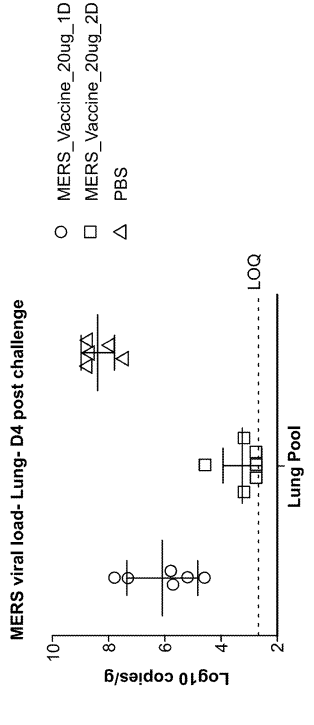


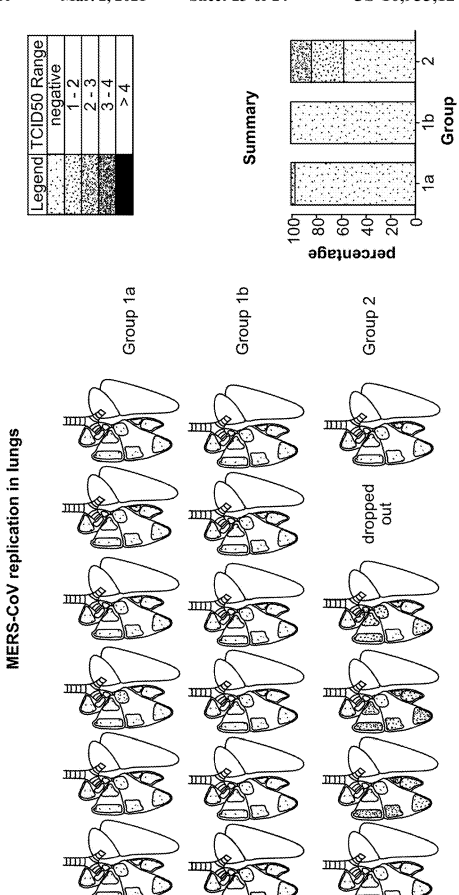
Fig. 19E



MERS\_Vaccine\_20ug\_1D MERS\_Vaccine\_20ug\_2D PBS 0  $\triangleleft$ MERS viral load- Lung- D4 post challenge Lung Pool 107 8 6 Log10 copies/g

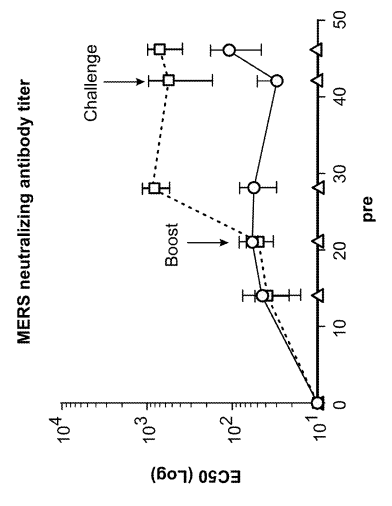
egend PCR Range Summary Group 5-6 percentage 2 6 6 8 Group 1a Group 1b Group 2 MERS-CoV RNA loads in lungs dropped out Ш

Fig. 20A



-O- MERS\_20ug\_1Dose -□- MERS\_20ug\_2Doses -△- Placebo

Fig. 21



## BETACORONAVIRUS MRNA VACCINE

## RELATED APPLICATIONS

This application is a division of U.S. application Ser. No. 5 16/805,587, filed Feb. 28, 2020, now U.S. Pat. No. 10,702, 600, which is a continuation of U.S. application Ser. No. 16/368,270, filed Mar. 28, 2019, now U.S. Pat. No. 10,702, 599, which is a continuation of Ser. No. 16/040,981, filed Jul. 20, 2018, now U.S. Pat. No. 10,272,150, which is a 10 continuation of U.S. application Ser. No. 15/674,599, filed Aug. 11, 2017, now U.S. Pat. No. 10,064,934, which is a continuation of International application number PCT/ US2016/058327, filed Oct. 21, 2016, which claims the benefit under 35 U.S.C. § 119(e) of U.S. provisional appli- 15 cation No. 62/244,802, filed Oct. 22, 2015, U.S. provisional application No. 62/247,297, filed Oct. 28, 2015, U.S. provisional application No. 62/244,946, filed Oct. 22, 2015, U.S. provisional application No. 62/247,362, filed Oct. 28, 2015, U.S. provisional application No. 62/244,813, filed 20 Oct. 22, 2015, U.S. provisional application No. 62/247,394, filed Oct. 28, 2015, U.S. provisional application No. 62/244, 837, filed Oct. 22, 2015, U.S. provisional application No. 62/247,483, filed Oct. 28, 2015, and U.S. provisional application No. 62/245,031, filed Oct. 22, 2015, each of which is 25 incorporated by reference herein in its entirety.

## BACKGROUND

Respiratory disease is a medical term that encompasses 30 pathological conditions affecting the organs and tissues that make gas exchange possible in higher organisms, and includes conditions of the upper respiratory tract, trachea, bronchi, bronchioles, alveoli, pleura and pleural cavity, and the nerves and muscles of breathing. Respiratory diseases 35 range from mild and self-limiting, such as the common cold, to life-threatening entities like bacterial pneumonia, pulmonary embolism, acute asthma and lung cancer. Respiratory disease is a common and significant cause of illness and death around the world. In the US, approximately 1 billion 40 "common colds" occur each year. Respiratory conditions are among the most frequent reasons for hospital stays among children.

The human *Metapneumovirus* (hMPV) is a negative-sense, single-stranded RNA virus of the genus *Pneumoviri-nae* and of the family Paramyxoviridae and is closely related to the avian *Metapneumovirus* (AMPV) subgroup C. It was isolated for the first time in 2001 in the Netherlands by using the RAP-PCR (RNA arbitrarily primed PCR) technique for identification of unknown viruses growing in cultured cells. hPMV is second only to RSV as an important cause of viral lower respiratory tract illness (LRI) in young children. The seasonal epidemiology of hMPV appears to be similar to that of RSV, but the incidence of infection and illness appears to be substantially lower.

Parainfluenza virus type 3 (PIV3), like hMPV, is also a negative-sense, single-stranded sense RNA virus of the genus *Pneumovirinae* and of the family Paramyxoviridae and is a major cause of ubiquitous acute respiratory infections of infancy and early childhood. Its incidence peaks 60 around 4-12 months of age, and the virus is responsible for 3-10% of hospitalizations, mainly for bronchiolitis and pneumonia. PIV3 can be fatal, and in some instances is associated with neurologic diseases, such as febrile seizures. It can also result in airway remodeling, a significant cause of 65 morbidity. In developing regions of the world, infants and young children are at the highest risk of mortality, either

2

from primary PIV3 viral infection or a secondary consequences, such as bacterial infections. Human parainfluenza viruses (hPIV) types 1, 2 and 3 (hPIV1, hPIV2 and hPIV3, respectively), also like hMPV, are second only to RSV as important causes of viral LRI in young children.

RSV, too, is a negative-sense, single-stranded RNA virus of the genus Pneumovirinae and of the family Paramyxoviridae. Symptoms in adults typically resemble a sinus infection or the common cold, although the infection may be asymptomatic. In older adults (e.g., >60 years), RSV infection may progress to bronchiolitis or pneumonia. Symptoms in children are often more severe, including bronchiolitis and pneumonia. It is estimated that in the United States, most children are infected with RSV by the age of three. The RSV virion consists of an internal nucleocapsid comprised of the viral RNA bound to nucleoprotein (N), phosphoprotein (P), and large polymerase protein (L). The nucleocapsid is surrounded by matrix protein (M) and is encapsulated by a lipid bilayer into which the viral fusion (F) and attachment (G) proteins as well as the small hydrophobic protein (SH) are incorporated. The viral genome also encodes two nonstructural proteins (NS1 and NS2), which inhibit type I interferon activity as well as the M-2 protein.

The continuing health problems associated with hMPV, PIV3 and RSV are of concern internationally, reinforcing the importance of developing effective and safe vaccine candidates against these virus.

Despite decades of research, no vaccines currently exist (Sato and Wright, *Pediatr. Infect. Dis. J.* 2008; 27(10 Suppl): S123-5). Recombinant technology, however, has been used to target the formation of vaccines for hPIV-1, 2 and 3 serotypes, for example, and has taken the form of several live-attenuated intranasal vaccines. Two vaccines in particular were found to be immunogenic and well tolerated against hPIV-3 in phase I trials. hPIV1 and hPIV2 vaccine candidates remain less advanced (Durbin and Karron, Clinical infectious diseases: an official publication of the Infectious Diseases Society of America 2003; 37(12):1668-77).

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In additional to hMPV, PIV, RSV and MeV, Betacoronaviruses are known to cause respiratory illnesses. Betacoronaviruses (BetaCoVs) are one of four genera of coronaviruses of the subfamily Coronavirinae in the family Coronaviridae, of the order Nidovirales. They are enveloped, positive-sense, single-stranded RNA viruses of zoonotic origin. The coronavirus genera are each composed of varying viral lineages, with the *Betacoronavirus* genus containing four such lineages. The BetaCoVs of the greatest clinical importance concerning humans are OC43 and HKU1 of the A lineage, SARS-CoV of the B lineage, and MERS-CoV of the C lineage. MERS-CoV is the first *Betacoronavirus* belonging to lineage C that is known to infect humans.

The Middle East respiratory syndrome coronavirus (MERS-CoV), or EMC/2012 (HCoV-EMC/2012), initially referred to as novel coronavirus 2012 or simply novel coronavirus, was first reported in 2012 after genome sequencing of a virus isolated from sputum samples from a person who fell ill during a 2012 outbreak of a new flu. As

of July 2015, MERS-CoV cases have been reported in over 21 countries. The outbreaks of MERS-CoV have raised serious concerns world-wide, reinforcing the importance of developing effective and safe vaccine candidates against MERS-CoV.

Severe acute respiratory syndrome (SARS) emerged in China in 2002 and spread to other countries before brought under control. Because of a concern for reemergence or a deliberate release of the SARS coronavirus, vaccine development was initiated.

Deoxyribonucleic acid (DNA) vaccination is one technique used to stimulate humoral and cellular immune responses to foreign antigens, such as hMPV antigens and/or PIV antigens and/or RSV antigens. The direct injection of genetically engineered DNA (e.g., naked plasmid DNA) into a living host results in a small number of its cells directly producing an antigen, resulting in a protective immunological response. With this technique, however, comes potential problems, including the possibility of insertional mutagenesis, which could lead to the activation of oncogenes or the inhibition of tumor suppressor genes.

#### **SUMMARY**

Provided herein are ribonucleic acid (RNA) vaccines that build on the knowledge that RNA (e.g., messenger RNA (mRNA)) can safely direct the body's cellular machinery to produce nearly any protein of interest, from native proteins to antibodies and other entirely novel protein constructs that 30 can have the rapeutic activity inside and outside of cells. The RNA (e.g., mRNA) vaccines of the present disclosure may be used to induce a balanced immune response against  $hMPV,\,PIV,\,RSV,\,MeV,\,and/or\,\,BetaCoV$  (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, 35 HCoV-NL, HCoV-NH and/or HCoV-HKU1), or any combination of two or more of the foregoing viruses, comprising both cellular and humoral immunity, without risking the possibility of insertional mutagenesis, for example. hMPV, PIV, RSV, MeV, BetaCoV (e.g., MERS-CoV, SARS-CoV, 40 HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1) and combinations thereof are referred to herein as "respiratory viruses." Thus, the term "respiratory virus RNA vaccines" encompasses hMPV RNA vaccines, PIV RNA vaccines, RSV RNA vaccines, MeV 45 RNA vaccines, BetaCoV RNA vaccines, and any combination of two or more of hMPV RNA vaccines, PIV RNA vaccines, RSV RNA vaccines, MeV RNA vaccines, and BetaCoV RNA vaccines.

The RNA (e.g., mRNA) vaccines may be utilized in 50 various settings depending on the prevalence of the infection or the degree or level of unmet medical need. The RNA (e.g. mRNA) vaccines may be utilized to treat and/or prevent a hMPV, PIV, RSV, MeV, a BetaCoV (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, 55 HCoV-NL, HCoV-NH, HCoV-HKU1), or any combination of two or more of the foregoing viruses, of various genotypes, strains, and isolates. The RNA (e.g., mRNA) vaccines have superior properties in that they produce much larger antibody titers and produce responses earlier than commer- 60 cially available anti-viral therapeutic treatments. While not wishing to be bound by theory, it is believed that the RNA (e.g., mRNA) vaccines, as mRNA polynucleotides, are better designed to produce the appropriate protein conformation upon translation as the RNA (e.g., mRNA) vaccines 65 co-opt natural cellular machinery. Unlike traditional vaccines, which are manufactured ex vivo and may trigger

4

unwanted cellular responses, RNA (e.g., mRNA) vaccines are presented to the cellular system in a more native fashion.

In some aspects the invention is a respiratory virus vaccine, comprising at least one RNA polynucleotide having an open reading frame encoding at least one respiratory virus antigenic polypeptide, formulated in a cationic lipid nanoparticle.

Surprisingly, in some aspects it has also been shown that efficacy of mRNA vaccines can be significantly enhanced when combined with a flagellin adjuvant, in particular, when one or more antigen-encoding mRNAs is combined with an mRNA encoding flagellin.

RNA (e.g., mRNA) vaccines combined with the flagellin adjuvant (e.g., mRNA-encoded flagellin adjuvant) have superior properties in that they may produce much larger antibody titers and produce responses earlier than commercially available vaccine formulations. While not wishing to be bound by theory, it is believed that the RNA (e.g., mRNA) vaccines, for example, as mRNA polynucleotides, are better designed to produce the appropriate protein conformation upon translation, for both the antigen and the adjuvant, as the RNA (e.g., mRNA) vaccines co-opt natural cellular machinery. Unlike traditional vaccines, which are manufactured ex vivo and may trigger unwanted cellular responses, RNA (e.g., mRNA) vaccines are presented to the cellular system in a more native fashion.

Some embodiments of the present disclosure provide RNA (e.g., mRNA) vaccines that include at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one antigenic polypeptide or an immunogenic fragment thereof (e.g., an immunogenic fragment capable of inducing an immune response to the antigenic polypeptide) and at least one RNA (e.g., mRNA polynucleotide) having an open reading frame encoding a flagellin adjuvant.

In some embodiments, at least one flagellin polypeptide (e.g., encoded flagellin polypeptide) is a flagellin protein. In some embodiments, at least one flagellin polypeptide (e.g., encoded flagellin polypeptide) is an immunogenic flagellin fragment. In some embodiments, at least one flagellin polypeptide and at least one antigenic polypeptide are encoded by a single RNA (e.g., mRNA) polynucleotide. In other embodiments, at least one flagellin polypeptide and at least one antigenic polypeptide are each encoded by a different RNA polynucleotide.

In some embodiments at least one flagellin polypeptide has at least 80%, at least 85%, at least 90%, or at least 95% identity to a flagellin polypeptide having a sequence identified by any one of SEQ ID NO: 54-56.

Provided herein, in some embodiments, is a ribonucleic acid (RNA) (e.g., mRNA) vaccine, comprising at least one (e.g., at least 2, 3, 4 or 5) RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one (e.g., at least 2, 3, 4 or 5) hMPV, PIV, RSV, MeV, or a BetaCoV (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH, HCoV-HKU1) antigenic polypeptide, or any combination of two or more of the foregoing antigenic polypeptides. Herein, use of the term "antigenic polypeptide" encompasses immunogenic fragments of the antigenic polypeptide (an immunogenic fragment that is induces (or is capable of inducing) an immune response to hMPV, PIV, RSV, MeV, or a BetaCoV), unless otherwise stated.

Also provided herein, in some embodiments, is a RNA (e.g., mRNA) vaccine comprising at least one (e.g., at least 2, 3, 4 or 5) RNA polynucleotide having an open reading frame encoding at least one (e.g., at least 2, 3, 4 or 5) hMPV,

PIV, RSV, MeV, and/or a BetaCoV (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH, HCoV-HKU1) antigenic polypeptide or an immunogenic fragment thereof, linked to a signal peptide.

Further provided herein, in some embodiments, is a nucleic acid (e.g., DNA) encoding at least one (e.g., at least 2, 3, 4 or 5) hMPV, PIV, RSV, MeV, and/or a BetaCoV (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH, HCoV-HKU1) RNA (e.g., mRNA) polynucleotide.

Further still, provided herein, in some embodiments, is a method of inducing an immune response in a subject, the method comprising administering to the subject a vaccine comprising at least one (e.g., at least 2, 3, 4 or 5) RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one (e.g., at least 2, 3, 4 or 5) hMPV, PIV, RSV, MeV, and/or a BetaCoV (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, 20 HCoV-NH, HCoV-HKU1) antigenic polypeptide, or any combination of two or more of the foregoing antigenic polypeptides.

#### hMPV/PIV3/RSV

In some embodiments, a RNA (e.g., mRNA) vaccine 25 comprises at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one hMPV, PIV3 or RSV antigenic polypeptide. In some embodiments, at least one antigenic polypeptide is a hMPV, PIV3 or RSV polyprotein. In some embodiments, at least one antigenic 30 polypeptide is major surface glycoprotein G or an immunogenic fragment thereof. In some embodiments, at least one antigenic polypeptide is Fusion (F) glycoprotein (e.g., Fusion glycoprotein F0, F1 or F2) or an immunogenic fragment thereof. In some embodiments, at least one anti- 35 genic polypeptide is major surface glycoprotein G or an immunogenic fragment thereof and F glycoprotein or an immunogenic fragment thereof. In some embodiments, the antigenic polypeptide is nucleoprotein (N) or an immunogenic fragment thereof, phosphoprotein (P) or an immuno- 40 genic fragment thereof, large polymerase protein (L) or an immunogenic fragment thereof, matrix protein (M) or an immunogenic fragment thereof, small hydrophobic protein (SH) or an immunogenic fragment thereof nonstructural protein 1 (NS1) or an immunogenic fragment thereof, or 45 nonstructural protein 2 (NS2) and an immunogenic fragment thereof.

In some embodiments, at least one hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 (Table 3; see also amino acid 50 sequences of Table 4). In some embodiments, the amino acid sequence of the hMPV antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 5-8 (Table 55 3; see also amino acid sequences of Table 4).

In some embodiments, at least one hMPV antigenic polypeptide is encoded by a nucleic acid sequence identified by any one of SEQ ID NO: 1-4 (Table 2).

In some embodiments, at least one hMPV RNA (e.g., 60 mRNA) polynucleotide is encoded by a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 1-4 (Table 2). In some embodiments, at least one hMPV RNA (e.g., mRNA) polynucleotide comprises a nucleic acid sequence, or a fragment of a 65 nucleotide sequence, identified by any one of SEQ ID NO: 57-60 (Table 2).

6

In some embodiments, at least one antigenic polypeptide is obtained from hMPV strain CAN98-75 (CAN75) or the hMPV strain CAN97-83 (CAN83).

In some embodiments, at least one PIV3 antigenic polypeptide comprises hemagglutinin-neuraminidase, Fusion (F) glycoprotein, matrix protein (M), nucleocapsid protein (N), viral replicase (L), non-structural V protein, or an immunogenic fragment thereof.

In some embodiments, at least one PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 (Table 6; see also amino acid sequences of Table 7). In some embodiments, the amino acid sequence of the PIV3 antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 12-13 (Table 6; see also amino acid sequences of Table 7).

In some embodiments, at least one PIV3 antigenic polypeptide is encoded by a nucleic acid sequence identified by any one of SEQ ID NO: 9-12 (Table 5; see also nucleic acid sequences of Table 7).

In some embodiments, at least one PIV3 RNA (e.g., mRNA) polynucleotide is encoded by a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 9-12 (Table 5; see also nucleic acid sequences of Table 7). In some embodiments, at least one PIV3 RNA (e.g., mRNA) polynucleotide comprises a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 61-64 (Table 5)

In some embodiments, at least one antigenic polypeptide is obtained from PIV3 strain HPIV3/Homo sapiens/PER/FLA4815/2008.

In some embodiments, at least one RSV antigenic polypeptide comprises at least one antigenic polypeptide that comprises glycoprotein G, glycoprotein F, or an immunogenic fragment thereof. In some embodiments, at least one RSV antigenic polypeptide comprises at least one antigenic polypeptide that comprises glycoprotein F and at least one or at least two antigenic polypeptide selected from G, M, N, P, L, SH, M2, NS1 and NS2.

MeV

In some embodiments, a RNA (e.g., mRNA) vaccine comprises at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one MeV antigenic polypeptide. In some embodiments, at least one antigenic polypeptide is a hemagglutinin (HA) protein or an immunogenic fragment thereof. The HA protein may be from MeV strain D3 or B8, for example. In some embodiments, at least one antigenic polypeptide is a Fusion (F) protein or an immunogenic fragment thereof. The F protein may be from MeV strain D3 or B8, for example. In some embodiments, a MeV RNA (e.g., mRNA) vaccines comprises a least one RNA polynucleotide encoding a HA protein and a F protein. The HA and F proteins may be from MeV strain D3 or B8, for example.

In some embodiments, at least one MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 (Table 14). In some embodiments, the amino acid sequence of the MeV antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 47-50 (Table 14).

In some embodiments, at least one MeV antigenic polypeptide is encoded by a nucleic acid sequence of SEQ ID NO: 35-46 (Table 13).

In some embodiments, at least one MeV RNA (e.g., mRNA) polynucleotide is encoded by a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 35-46 (Table 13). In some embodiments, at least one MeV RNA (e.g., mRNA) polynucleotide comprises a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 69-80 (Table 13).

In some embodiments, at least one antigenic polypeptide is obtained from MeV strain B3/B3.1, C2, D4, D6, D7, D8, 10 G3, H1, Moraten, Rubeovax, MVi/New Jersey.USA/45.05, MVi/Texas.USA/4.07, AIK-C, MVi/New York.USA/26.09/3, MVi/California.USA/16.03, MVi/Virginia.USA/15.09, MVi/California.USA/8.04, or MVi/Pennsylvania.USA/20.09.

## BetaCoV

In some embodiments, a RNA (e.g., mRNA) vaccine comprises at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one Beta-CoV antigenic polypeptide. In some embodiments, the Beta-20 CoV is MERS-CoV. In some embodiments, the BetaCoV is SARS-CoV. In some embodiments, the BetaCoV is HCoV-OC43. In some embodiments, the BetaCoV is HCoV-229E. In some embodiments, the BetaCoV is HCoV-NL63. In some embodiments, the BetaCoV is HCoV-HKU1. In some 25 embodiments, at least one antigenic polypeptide is a Betacoronavirus structural protein. For example, a Betacoronavirus structural protein may be spike protein (S), envelope protein (E), nucleocapsid protein (N), membrane protein (M) or an immunogenic fragment thereof. In some embodi- 30 ments, a Betacoronavirus structural protein is a spike protein (S). In some embodiments, a Betacoronavirus structural protein is a S1 subunit or a S2 subunit of spike protein (S) or an immunogenic fragment thereof.

BetaCoV RNA (e.g., mRNA) polynucleotides of the vac- 35 cines provided herein may encode viral protein components of Betacoronaviruses, for example, accessory proteins, replicase proteins and the like are encompassed by the present disclosure. RNA (e.g., mRNA) vaccines may include RNA polynucleotides encoding at least one accessory protein 40 (e.g., protein 3, protein 4a, protein 4b, protein 5), at least one replicase protein (e.g., protein 1a, protein 1b), or a combination of at least one accessory protein and at least one replicase protein. The present disclosure also encompasses RNA (e.g., mRNA) vaccines comprising RNA (e.g., mRNA) 45 polynucleotides encoding an accessory protein and/or a replicase protein in combination with at least one structural protein. Due to their surface expression properties, vaccines featuring RNA polynucleotides encoding structural proteins are believed to have preferred immunogenic activity and, 50 hence, may be most suitable for use in the vaccines of the present disclosure.

Some embodiments of the present disclosure provide *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH, 55 HCoV-HKU1 or a combination thereof) vaccines that include at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH, HCoV- 60 HKU1) antigenic polypeptide. Also provided herein are pan-*Betacoronavirus* vaccines. Thus, a *Betacoronavirus* vaccine comprising a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding any one, two, three or four of MERS-CoV, SARS-CoV, HCoV-OC43, HCoV- 65 229E, HCoV-NL63, and HCoV-HKU1, for example, may be effective against any one of, any combination of, or all of,

8

MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1. Other Betacoronaviruses are encompassed by the present disclosure.

In some embodiments, at least one antigenic polypeptide is a MERS-CoV structural protein. For example, a MERS-CoV structural protein may be spike protein (S), envelope protein (E), nucleocapsid protein (N), membrane protein (M) or an immunogenic fragment thereof. In some embodiments, the MERS-CoV structural protein is a spike protein (S) (see, e.g., Coleman C M et al. *Vaccine* 2014; 32:3169-74, incorporated herein by reference). In some embodiments, the MERS-CoV structural protein is a S1 subunit or a S2 subunit of spike protein (S) or an immunogenic fragment thereof (Li J et al. *Viral Immunol* 2013; 26(2):126-32; He Y et al. *Biochem Biophys Res Commun* 2004; 324(2):773-81, each of which is incorporated herein by reference).

In some embodiments, at least one MERS-CoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-28 or 33 (Table 11). In some embodiments, the amino acid sequence of the MERS-CoV antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 24-28 or 33 (Table 11).

In some embodiments, at least one MERS-CoV antigenic polypeptide is encoded by a nucleic acid sequence identified by any one of SEQ ID NO: 20-23 (Table 10).

In some embodiments, at least one MERS-CoV RNA (e.g., mRNA) polynucleotide is encoded by a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 20-23 (Table 10). In some embodiments, at least one MERS-CoV RNA (e.g., mRNA) polynucleotide comprises a nucleic acid sequence, or a fragment of a nucleotide sequence, identified by any one of SEQ ID NO: 65-68 (Table 10).

In some embodiments, at least one antigenic polypeptide is obtained from MERS-CoV strain Riyadh\_14\_2013, 2cEMC/2012, or Hasa\_1\_2013.

In some embodiments, at least one antigenic polypeptide is a SARS-CoV structural protein. For example, a SARS-CoV structural protein may be spike protein (S), envelope protein (E), nucleocapsid protein (N), membrane protein (M) or an immunogenic fragment thereof. In some embodiments, the SARS-CoV structural protein is a spike protein (S). In some embodiments, the SARS-CoV structural protein is a S1 subunit or a S2 subunit of spike protein (S) or an immunogenic fragment thereof.

In some embodiments, at least one SARS-CoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 29, 32 or 34 (Table 11). In some embodiments, the amino acid sequence of the SARS-CoV antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 29, 32 or 34 (Table 11).

In some embodiments, at least one antigenic polypeptide is a HCoV-OC43 structural protein. For example, a HCoV-OC43 structural protein may be spike protein (S), envelope protein (E), nucleocapsid protein (N), membrane protein (M) or an immunogenic fragment thereof. In some embodiments, the HCoV-OC43 structural protein is a spike protein (S). In some embodiments, the HCoV-OC43 structural protein is a S1 subunit or a S2 subunit of spike protein (S) or an immunogenic fragment thereof.

In some embodiments, at least one HCoV-OC43 antigenic polypeptide comprises an amino acid sequence identified by

any one of SEQ ID NO: 30 (Table 11). In some embodiments, the amino acid sequence of the HCoV-OC43 antigenic polypeptide is, or is a fragment of, or is a homolog or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any 5 one of SEQ ID NO: 30 (Table 11).

In some embodiments, an antigenic polypeptide is a HCoV-HKU1 structural protein. For example, a HCoV-HKU1 structural protein may be spike protein (S), envelope protein (E), nucleocapsid protein (N), membrane protein (M) or an immunogenic fragment thereof. In some embodiments, the HCoV-HKU1 structural protein is a spike protein (S). In some embodiments, the HCoV-HKU1 structural protein is a S1 subunit or a S2 subunit of spike protein (S) or an immunogenic fragment thereof.

In some embodiments, at least one HCoV-HKU1 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 31 (Table 11). In some embodiments, the amino acid sequence of the HCoV-HKU1 antigenic polypeptide is, or is a fragment of, or is a homolog 20 or variant having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to, the amino acid sequence identified by any one of SEQ ID NO: 31 (Table 11).

In some embodiments, an open reading frame of a RNA (e.g., mRNA) vaccine is codon-optimized. In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and is codon optimized mRNA.

In some embodiments, a RNA (e.g., mRNA) vaccine further comprising an adjuvant.

Tables 4, 7, 12 and 15 provide National Center for Biotechnology Information (NCBI) accession numbers of interest. It should be understood that the phrase "an amino 35 acid sequence of Tables 4, 7, 12 and 15" refers to an amino acid sequence identified by one or more NCBI accession numbers listed in Tables 4, 7, 12 and 15. Each of the amino acid sequences, and variants having greater than 95% identity or greater than 98% identity to each of the amino acid 40 sequences encompassed by the accession numbers of Tables 4, 7, 12 and 15 are included within the constructs (polynucleotides/polypeptides) of the present disclosure.

In some embodiments, at least one mRNA polynucleotide is encoded by a nucleic acid having a sequence identified by 45 any one of SEQ ID NO: 1-4, 9-12, 20-23, or 35-46 (Tables 2, 5, 10 and 13; see also nucleic acid sequences of Table 7) and having less than 80% identity to wild-type mRNA sequence. In some embodiments, at least one mRNA polynucleotide is encoded by a nucleic acid having a sequence 50 identified by any one of SEQ ID NO: 1-4, 9-12, 20-23, or 35-46 (Tables 2, 5, 10 and 13; see also nucleic acid sequences of Table 7) and having less than 75%, 85% or 95% identity to a wild-type mRNA sequence. In some embodiments, at least one mRNA polynucleotide is encoded 55 by a nucleic acid having a sequence identified by any one of SEQ ID NO: 1-4, 9-12, 20-23, or 35-46 (Tables 2, 5, 10 and 13; see also nucleic acid sequences of Table 7) and having less than 50-80%, 60-80%, 40-80%, 30-80%, 70-80%, 75-80% or 78-80% identity to wild-type mRNA sequence. 60 In some embodiments, at least one mRNA polynucleotide is encoded by a nucleic acid having a sequence identified by any one of SEQ ID NO: 1-4, 9-12, 20-23, or 35-46 (Tables 2, 5, 10 and 13; see also nucleic acid sequences of Table 7) and having less than 40-85%, 50-85%, 60-85%, 30-85%, 65 70-85%, 75-85% or 80-85% identity to wild-type mRNA sequence. In some embodiments, at least one mRNA poly10

nucleotide is encoded by a nucleic acid having a sequence identified by any one of SEQ ID NO: 1-4, 9-12, 20-23, or 35-46 (Tables 2, 5, 10 and 13; see also nucleic acid sequences of Table 7) and having less than 40-90%, 50-90%, 60-90%, 30-90%, 70-90%, 75-90%, 80-90%, or 85-90% identity to wild-type mRNA sequence.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and having at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) identity to wild-type mRNA sequence, but does not include wild-type mRNA sequence.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and has less than 95%, 90%, 85%, 80% or 75% identity to wild-type mRNA sequence. In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and has 30-80%, 40-80%, 50-80%, 60-80%, 70-80%, 75-80% or 78-80%, 30-85%, 40-85%, 50-805%, 60-85%, 70-85%, 75-85% or 78-85%, 30-90%, 40-90%, 50-90%, 60-90%, 70-90%, 75-90%, 80-90% or 85-90% identity to wild-type 30 mRNA sequence.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15). In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having 95%-99% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15).

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and having membrane fusion activity. In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide having 95%-99% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, or 47-50 (Tables 3, 6, 11 and 14; see also amino acid sequences of Tables 4, 7, 12 and 15) and having membrane fusion activity.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides) that attaches to cell receptors.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide (e.g., at least one

hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides) that causes fusion of viral and cellular membranes.

In some embodiments, at least one RNA polynucleotide encodes at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least 12

ionizable cationic lipid and the non-cationic lipid is a neutral lipid, and the sterol is a cholesterol. In some embodiments, a cationic lipid is selected from the group consisting of 2.2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), di((Z)-non-2-en-1-v1) 9-((4-(dimethylamino)butanovl)oxy)heptadecanedioate (L319). (12Z,15Z)—N,N-dimethyl-2-nonylhenicosa-12,15-dien-1amine (L608), and N,N-dimethyl-1-[(1S,2R)-2-octylcyclopropyl]heptadecan-8-amine (L530).

In some embodiments, the lipid is

one MeV antigenic polypeptide, or at least one BetaCoV 25 antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides) that is responsible for binding of the virus to a cell being 30 infected.

Some embodiments of the present disclosure provide a vaccine that includes at least one ribonucleic acid (RNA) (e.g., mRNA) polynucleotide having an open reading frame 35 encoding at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides), at least one 5' terminal cap and at least one chemical modification, formulated within a lipid nanoparticle.

In some embodiments, a 5' terminal cap is 7mG(5')ppp (5')NlmpNp.

In some embodiments, at least one chemical modification is selected from pseudouridine, N1-methylpseudouridine, 50 N1-ethylpseudouridine, 2-thiouridine, 4'-thiouridine. 5-methylcytosine, 5-methyluridine, 2-thio-1-methyl-1deaza-pseudouridine, 2-thio-1-methyl-pseudouridine, 2-thio-5-aza-uridine, 2-thio-dihydropseudouridine, 2-thiodihydrouridine, 2-thio-pseudouridine, 4-methoxy-2-thio-55 pseudouridine, 4-methoxy-pseudouridine, 4-thio-1-methyl-4-thio-pseudouridine, pseudouridine, 5-aza-uridine, dihydropseudouridine, 5-methoxyuridine and 2'-O-methyl uridine. In some embodiments, the chemical modification is in the 5-position of the uracil. In some embodiments, the chemical modification is a N1-methylpseudouridine. In some embodiments, the chemical modification is a N1-ethylpseudouridine.

In some embodiments, a lipid nanoparticle comprises a 65 cationic lipid, a PEG-modified lipid, a sterol and a noncationic lipid. In some embodiments, a cationic lipid is an

In some embodiments, the lipid is

In some embodiments, a lipid nanoparticle comprises compounds of Formula (I) and/or Formula (II), discussed

In some embodiments, a respiratory virus RNA (e.g., antigenic polypeptide, e.g., selected from MERS-CoV, 40 mRNA) vaccine is formulated in a lipid nanoparticle that comprises a compound selected from Compounds 3, 18, 20, 25, 26, 29, 30, 60, 108-112 and 122, described below.

Some embodiments of the present disclosure provide a vaccine that includes at least one RNA (e.g., mRNA) poly-45 nucleotide having an open reading frame encoding at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides), wherein at least 80% (e.g., 85%, 90%, 95%, 98%, 99%) of the uracil in the open reading frame have a chemical modification, optionally wherein the vaccine is formulated in a lipid nanoparticle (e.g., a lipid nanoparticle comprises a cationic lipid, a PEG-modified lipid, a sterol and a non-cationic lipid).

In some embodiments, 100% of the uracil in the open reading frame have a chemical modification. In some embodiments, a chemical modification is in the 5-position of the uracil. In some embodiments, a chemical modification is a N1-methyl pseudouridine. In some embodiments, 100% of the uracil in the open reading frame have a N1-methyl pseudouridine in the 5-position of the uracil.

In some embodiments, an open reading frame of a RNA (e.g., mRNA) polynucleotide encodes at least two antigenic

polypeptides (e.g., at least two hMPV antigenic polypeptides, at least two PIV3 antigenic polypeptides, at least two RSV antigenic polypeptides, at least two MeV antigenic polypeptides, or at least two BetaCoV antigenic polypeptides, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides). In some embodiments, the open reading frame encodes at least five or at least ten antigenic polypeptides. In some embodiments, the open reading frame encodes at least 100 antigenic polypeptides. In some embodiments, the open reading frame encodes at least 100 antigenic polypeptides.

In some embodiments, a vaccine comprises at least two RNA (e.g., mRNA) polynucleotides, each having an open reading frame encoding at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at 20 least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides). In some embodiments, the vaccine comprises 25 at least five or at least ten RNA (e.g., mRNA) polynucleotides, each having an open reading frame encoding at least one antigenic polypeptide or an immunogenic fragment thereof. In some embodiments, the vaccine comprises at least 100 RNA (e.g., mRNA) polynucleotides, each having an open reading frame encoding at least one antigenic polypeptide. In some embodiments, the vaccine comprises 2-100 RNA (e.g., mRNA) polynucleotides, each having an open reading frame encoding at least one antigenic polypeptide.

In some embodiments, at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV antigenic polypeptide, e.g., selected from 40 MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides) is fused to a signal peptide. In some embodiments, the signal peptide is selected from: a HuIgGk signal 45 peptide (METPAQLLFLLLLWLPDTTG; SEQ ID NO: 15); IgE heavy chain epsilon-1 signal peptide (MDWTWIL-FLVAAATRVHS; SEQ ID NO: 16); Japanese encephalitis PRM signal sequence (MLGSNSGQRVVFTILLLLVA-PAYS; SEQ ID NO: 17), VSVg protein signal sequence 50 (MKCLLYLAFLFIGVNCA; SEQ ID NO: 18) and Japanese encephalitis JEV signal sequence (MWLVSLAIVTA-CAGA; SEQ ID NO: 19).

In some embodiments, the signal peptide is fused to the N-terminus of at least one antigenic polypeptide. In some 55 embodiments, a signal peptide is fused to the C-terminus of at least one antigenic polypeptide.

In some embodiments, at least one antigenic polypeptide (e.g., at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic 60 polypeptide, at least one MeV antigenic polypeptide, or at least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic 65 polypeptides) comprises a mutated N-linked glycosylation site

14

Also provided herein is a RNA (e.g., mRNA) vaccine of any one of the foregoing paragraphs (e.g., a hMPV vaccine, a PIV3 vaccine, a RSV vaccine, a MeV vaccine, or a BetaCoV vaccine, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing vaccines), formulated in a nanoparticle (e.g., a lipid nanoparticle).

In some embodiments, the nanoparticle has a mean diameter of 50-200 nm. In some embodiments, the nanoparticle is a lipid nanoparticle. In some embodiments, the lipid nanoparticle comprises a cationic lipid, a PEG-modified lipid, a sterol and a non-cationic lipid. In some embodiments, the lipid nanoparticle comprises a molar ratio of about 20-60% cationic lipid, 0.5-15% PEG-modified lipid, 25-55% sterol, and 25% non-cationic lipid. In some embodiments, the cationic lipid is an ionizable cationic lipid and the non-cationic lipid is a neutral lipid, and the sterol is a cholesterol. In some embodiments, the cationic lipid is selected from 2.2-dilinoleyl-4-dimethylaminoethyl[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (I.319).

In some embodiments, a lipid nanoparticle comprises compounds of Formula (I) and/or Formula (II), as discussed below.

In some embodiments, a lipid nanoparticle comprises Compounds 3, 18, 20, 25, 26, 29, 30, 60, 108-112, or 122, as discussed below.

In some embodiments, the nanoparticle has a polydispersity value of less than 0.4 (e.g., less than 0.3, 0.2 or 0.1).

In some embodiments, the nanoparticle has a net neutral charge at a neutral pH value.

In some embodiments, the respiratory virus vaccine is multivalent.

Some embodiments of the present disclosure provide methods of inducing an antigen specific immune response in a subject, comprising administering to the subject any of the RNA (e.g., mRNA) vaccine as provided herein in an amount effective to produce an antigen-specific immune response. In some embodiments, the RNA (e.g., mRNA) vaccine is a hMPV vaccine, a PIV3 vaccine, a RSV vaccine, a MeV vaccine, or a BetaCoV vaccine, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1 vaccines. In some embodiments, the RNA (e.g., mRNA) vaccine is a combination vaccine comprising a combination of any two or more of the foregoing vaccines.

In some embodiments, an antigen-specific immune response comprises a T cell response or a B cell response.

In some embodiments, a method of producing an antigen-specific immune response comprises administering to a subject a single dose (no booster dose) of a RNA (e.g., mRNA) vaccine of the present disclosure. In some embodiments, the RNA (e.g., mRNA) vaccine is a hMPV vaccine, a PIV3 vaccine, a RSV vaccine, a MeV vaccine, or a BetaCoV vaccine, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1 vaccines. In some embodiments, the RNA (e.g., mRNA) vaccine is a combination vaccine comprising a combination of any two or more of the foregoing vaccines.

In some embodiments, a method further comprises administering to the subject a second (booster) dose of a RNA (e.g., mRNA) vaccine. Additional doses of a RNA (e.g., mRNA) vaccine may be administered.

15 In some embodiments, the subjects exhibit a seroconver-

sion rate of at least 80% (e.g., at least 85%, at least 90%, or at least 95%) following the first dose or the second (booster)

dose of the vaccine. Seroconversion is the time period

detectable in the blood. After seroconversion has occurred,

a virus can be detected in blood tests for the antibody.

During an infection or immunization, antigens enter the

blood, and the immune system begins to produce antibodies

may not be detectable, but antibodies are considered absent. During seroconversion, antibodies are present but not yet

detectable. Any time after seroconversion, the antibodies can be detected in the blood, indicating a prior or current

in response. Before seroconversion, the antigen itself may or 10

during which a specific antibody develops and becomes 5

In some embodiments, a RNA (e.g., mRNA) vaccine is administered to a subject by intradermal or intramuscular injection.

Some embodiments, of the present disclosure provide methods of inducing an antigen specific immune response in 20 a subject, including administering to a subject a RNA (e.g., mRNA) vaccine in an effective amount to produce an antigen specific immune response in a subject. Antigenspecific immune responses in a subject may be determined, in some embodiments, by assaying for antibody titer (for 25 titer of an antibody that binds to a hMPV, PIV3, RSV, MeV and/or BetaCoV antigenic polypeptide) following administration to the subject of any of the RNA (e.g., mRNA) vaccines of the present disclosure. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the 30 subject is increased by at least 1 log relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased by 1-3 log relative to a control.

In some embodiments, the anti-antigenic polypeptide 35 antibody titer produced in a subject is increased at least 2 times relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased at least 5 times relative to a control. In some embodiments, the anti-antigenic polypeptide antibody 40 titer produced in the subject is increased at least 10 times relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased 2-10 times relative to a control.

In some embodiments, the control is an anti-antigenic 45 polypeptide antibody titer produced in a subject who has not been administered a RNA (e.g., mRNA) vaccine of the present disclosure. In some embodiments, the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a live attenuated or inacti- 50 vated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine (see, e.g., Ren J. et al. J of Gen. Virol. 2015; 96: 1515-1520), or wherein the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a recombinant or purified hMPV, PIV3, RSV, MeV 55 and/or BetaCoV protein vaccine. In some embodiments, the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a hMPV, PIV3, RSV, MeV and/or BetaCoV virus-like particle (VLP) vaccine (see, e.g., Cox R G et al., J Virol. 2014 June; 88(11): 60

A RNA (e.g., mRNA) vaccine of the present disclosure is administered to a subject in an effective amount (an amount effective to induce an immune response). In some embodiments, the effective amount is a dose equivalent to an at least 65 2-fold, at least 4-fold, at least 10-fold, at least 100-fold, at least 1000-fold reduction in the standard of care dose of a

recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, wherein the anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, a purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, a live attenuated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine, an inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine, or a hMPV, PIV3, RSV, MeV and/or BetaCoV VLP vaccine. In some embodiments, the effective amount is a dose equivalent to 2-1000-fold reduction in the standard of care dose of a recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, wherein the anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, a purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, a live attenuated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine, an inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine, or a hMPV, PIV3, RSV, MeV and/or BetaCoV VLP vaccine.

16

In some embodiments, the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a virus-like particle (VLP) vaccine comprising structural proteins of hMPV, PIV3, RSV, MeV and/or BetaCoV.

In some embodiments, the RNA (e.g., mRNA) vaccine is formulated in an effective amount to produce an antigen specific immune response in a subject.

In some embodiments, the effective amount is a total dose of 25  $\mu g$  to 1000  $\mu g$ , or 50  $\mu g$  to 1000  $\mu g$ . In some embodiments, the effective amount is a total dose of 100 µg. In some embodiments, the effective amount is a dose of 25 μg administered to the subject a total of two times. In some embodiments, the effective amount is a dose of 100 µg administered to the subject a total of two times. In some embodiments, the effective amount is a dose of 400 µg administered to the subject a total of two times. In some embodiments, the effective amount is a dose of 500 µg administered to the subject a total of two times.

In some embodiments, the efficacy (or effectiveness) of a RNA (e.g., mRNA) vaccine is greater than 60%. In some embodiments, the RNA (e.g., mRNA) polynucleotide of the vaccine at least one hMPV antigenic polypeptide, at least one PIV3 antigenic polypeptide, at least one RSV antigenic polypeptide, at least one MeV antigenic polypeptide, at least one BetaCoV antigenic polypeptide, e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1, or any combination of two or more of the foregoing antigenic polypeptides.

Vaccine efficacy may be assessed using standard analyses (see, e.g., Weinberg et al., J Infect Dis. 2010 Jun. 1; 201(11):1607-10). For example, vaccine efficacy may be measured by double-blind, randomized, clinical controlled trials. Vaccine efficacy may be expressed as a proportionate reduction in disease attack rate (AR) between the unvaccinated (ARU) and vaccinated (ARV) study cohorts and can be calculated from the relative risk (RR) of disease among the vaccinated group with use of the following formulas:

Efficacy=(ARU-ARV)/ARU×100; and

Likewise, vaccine effectiveness may be assessed using standard analyses (see, e.g., Weinberg et al., J Infect Dis. 2010 Jun. 1; 201(11):1607-10). Vaccine effectiveness is an assessment of how a vaccine (which may have already proven to have high vaccine efficacy) reduces disease in a population. This measure can assess the net balance of benefits and adverse effects of a vaccination program, not just the vaccine itself, under natural field conditions rather than in a controlled clinical trial. Vaccine effectiveness is proportional to vaccine efficacy (potency) but is also affected by how well target groups in the population are immunized, as well as by other non-vaccine-related factors that influence the 'real-world' outcomes of hospitalizations, ambulatory visits, or costs. For example, a retrospective case control analysis may be used, in which the rates of vaccination among a set of infected cases and appropriate controls are compared. Vaccine effectiveness may be expressed as a rate difference, with use of the odds ratio (OR) for developing infection despite vaccination:

Effectiveness=(1-OR)×100.

In some embodiments, the efficacy (or effectiveness) of a RNA (e.g., mRNA) vaccine is at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, or at least 90%.

In some embodiments, the vaccine immunizes the subject against hMPV, PIV3, RSV, MeV, BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1), or any combination of two or more of the foregoing viruses for 30 up to 2 years. In some embodiments, the vaccine immunizes the subject against hMPV, PIV3, RSV, MeV, BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1), or any combination of two or more of the 35 nucleic acid vaccines are unmodified. foregoing viruses for more than 2 years, more than 3 years, more than 4 years, or for 5-10 years.

In some embodiments, the subject is about 5 years old or younger. For example, the subject may be between the ages of about 1 year and about 5 years (e.g., about 1, 2, 3, 5 or 40 5 years), or between the ages of about 6 months and about 1 year (e.g., about 6, 7, 8, 9, 10, 11 or 12 months). In some embodiments, the subject is about 12 months or younger (e.g., 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2 months or 1 month). In some embodiments, the subject is about 6 months or 45 younger.

In some embodiments, the subject was born full term (e.g., about 37-42 weeks). In some embodiments, the subject was born prematurely, for example, at about 36 weeks of gestation or earlier (e.g., about 36, 35, 34, 33, 32, 31, 30, 29, 50 28, 27, 26 or 25 weeks). For example, the subject may have been born at about 32 weeks of gestation or earlier. In some embodiments, the subject was born prematurely between about 32 weeks and about 36 weeks of gestation. In such subjects, a RNA (e.g., mRNA) vaccine may be administered 55 later in life, for example, at the age of about 6 months to about 5 years, or older.

In some embodiments, the subject is pregnant (e.g., in the first, second or third trimester) when administered an RNA (e.g., mRNA) vaccine. Viruses such as hMPV, PIV3 and 60 RSV causes infections of the lower respiratory tract, mainly in infants and young children. One-third of RSV related deaths, for example, occur in the first year of life, with 99 percent of these deaths occurring in low-resource countries. It's so widespread in the United States that nearly all 65 children become infected with the virus before their second birthdays. Thus, the present disclosure provides RNA (e.g.,

18

mRNA) vaccines for maternal immunization to improve mother-to-child transmission of protection against the virus.

In some embodiments, the subject is a young adult between the ages of about 20 years and about 50 years (e.g., about 20, 25, 30, 35, 40, 45 or 50 years old).

In some embodiments, the subject is an elderly subject about 60 years old, about 70 years old, or older (e.g., about 60, 65, 70, 75, 80, 85 or 90 years old).

In some embodiments, the subject is has a chronic pulmonary disease (e.g., chronic obstructive pulmonary disease (COPD) or asthma). Two forms of COPD include chronic bronchitis, which involves a long-term cough with mucus, and emphysema, which involves damage to the lungs over time. Thus, a subject administered a RNA (e.g., mRNA) vaccine may have chronic bronchitis or emphysema.

In some embodiments, the subject has been exposed to hMPV, PIV3, RSV, MeV, BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1), or 20 any combination of two or more of the foregoing viruses; the subject is infected with hMPV, PIV3, RSV, MeV, BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1), or any combination of two or more of the foregoing viruses; or subject is at risk of infection by hMPV, PIV3, RSV, MeV, BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1), or any combination of two or more of the foregoing viruses.

In some embodiments, the subject is immunocompromised (has an impaired immune system, e.g., has an immune disorder or autoimmune disorder).

In some embodiments the nucleic acid vaccines described herein are chemically modified. In other embodiments the

Yet other aspects provide compositions for and methods of vaccinating a subject comprising administering to the subject a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a first respiratory virus antigenic polypeptide, wherein the RNA polynucleotide does not include a stabilization element, and wherein an adjuvant is not coformulated or co-administered with the vaccine.

In other aspects the invention is a composition for or method of vaccinating a subject comprising administering to the subject a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a first antigenic polypeptide wherein a dosage of between 10 μg/kg and 400 μg/kg of the nucleic acid vaccine is administered to the subject. In some embodiments the dosage of the RNA polynucleotide is 1-5 µg, 5-10 µg, 10-15  $\mu g$ , 15-20  $\mu g$ , 10-25  $\mu g$ , 20-25  $\mu g$ , 20-50  $\mu g$ , 30-50  $\mu g$ , 40-50 μд, 40-60 μд, 60-80 μд, 60-100 μд, 50-100 μд, 80-120 μд, 40-120 μg, 40-150 μg, 50-150 μg, 50-200 μg, 80-200 μg, 100-200 μg, 120-250 μg, 150-250 μg, 180-280 μg, 200-300 μд, 50-300 μд, 80-300 μд, 100-300 μд, 40-300 μд, 50-350 μд, 100-350 μд, 200-350 μд, 300-350 μд, 320-400 μд, 40-380 μg, 40-100 μg, 100-400 μg, 200-400 μg, or 300-400 μg per dose. In some embodiments, the nucleic acid vaccine is administered to the subject by intradermal or intramuscular injection. In some embodiments, the nucleic acid vaccine is administered to the subject on day zero. In some embodiments, a second dose of the nucleic acid vaccine is administered to the subject on day twenty one.

In some embodiments, a dosage of 25 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage

of 100 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage of 50 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage 5 of 75 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage of 150 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage 10 of 400 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, a dosage of 200 micrograms of the RNA polynucleotide is included in the nucleic acid vaccine administered to the subject. In some embodiments, the RNA polynucleotide accumulates at a 100 fold higher level in the local lymph node in comparison with the distal lymph node. In other embodiments the nucleic acid vaccine is chemically modified and in other embodiments the nucleic acid vaccine is not chemically modified.

Aspects of the invention provide a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a first antigenic polypeptide, wherein the RNA polynucleotide does not include a stabilization element, and a pharmaceutically acceptable carrier 25 or excipient, wherein an adjuvant is not included in the vaccine. In some embodiments, the stabilization element is a histone stem-loop. In some embodiments, the stabilization element is a nucleic acid sequence having increased GC content relative to wild type sequence.

Aspects of the invention provide nucleic acid vaccines comprising one or more RNA polynucleotides having an open reading frame encoding a first antigenic polypeptide, wherein the RNA polynucleotide is present in the formulation for in vivo administration to a host, which confers an 35 antibody titer superior to the criterion for seroprotection for the first antigen for an acceptable percentage of human subjects. In some embodiments, the antibody titer produced by the mRNA vaccines of the invention is a neutralizing antibody titer. In some embodiments the neutralizing anti- 40 body titer is greater than a protein vaccine. In other embodiments the neutralizing antibody titer produced by the mRNA vaccines of the invention is greater than an adjuvanted protein vaccine. In yet other embodiments the neutralizing antibody titer produced by the mRNA vaccines of the 45 invention is 1,000-10,000, 1,200-10,000, 1,400-10,000, 1,500-10,000, 1,000-5,000, 1,000-4,000, 1,800-10,000, 2000-10,000, 2,000-5,000, 2,000-3,000, 2,000-4,000, 3,000-5,000, 3,000-4,000, or 2,000-2,500. A neutralization titer is typically expressed as the highest serum dilution required to 50 achieve a 50% reduction in the number of plaques.

Also provided are nucleic acid vaccines comprising one or more RNA polynucleotides having an open reading frame encoding a first antigenic polypeptide, wherein the RNA polynucleotide is present in a formulation for in vivo administration to a host for eliciting a longer lasting high antibody titer than an antibody titer elicited by an mRNA vaccine having a stabilizing element or formulated with an adjuvant and encoding the first antigenic polypeptide. In some embodiments, the RNA polynucleotide is formulated to 60 produce a neutralizing antibodies within one week of a single administration. In some embodiments, the adjuvant is selected from a cationic peptide and an immunostimulatory nucleic acid. In some embodiments, the cationic peptide is protamine.

Aspects provide nucleic acid vaccines comprising one or more RNA polynucleotides having an open reading frame comprising at least one chemical modification or optionally no nucleotide modification, the open reading frame encoding a first antigenic polypeptide, wherein the RNA polynucleotide is present in the formulation for in vivo administration to a host such that the level of antigen expression in the host significantly exceeds a level of antigen expression produced by an mRNA vaccine having a stabilizing element or formulated with an adjuvant and encoding the first antigenic polypeptide.

20

Other aspects provide nucleic acid vaccines comprising one or more RNA polynucleotides having an open reading frame comprising at least one chemical modification or optionally no nucleotide modification, the open reading frame encoding a first antigenic polypeptide, wherein the vaccine has at least 10 fold less RNA polynucleotide than is required for an unmodified mRNA vaccine to produce an equivalent antibody titer. In some embodiments, the RNA polynucleotide is present in a dosage of 25-100 micrograms.

Aspects of the invention also provide a unit of use vaccine, comprising between 10 ug and 400 ug of one or more RNA polynucleotides having an open reading frame comprising at least one chemical modification or optionally no nucleotide modification, the open reading frame encoding a first antigenic polypeptide, and a pharmaceutically acceptable carrier or excipient, formulated for delivery to a human subject. In some embodiments, the vaccine further comprises a cationic lipid nanoparticle.

Aspects of the invention provide methods of creating, maintaining or restoring antigenic memory to a respiratory virus strain in an individual or population of individuals comprising administering to said individual or population an antigenic memory booster nucleic acid vaccine comprising (a) at least one RNA polynucleotide, said polynucleotide comprising at least one chemical modification or optionally no nucleotide modification and two or more codon-optimized open reading frames, said open reading frames encoding a set of reference antigenic polypeptides, and (b) optionally a pharmaceutically acceptable carrier or excipient. In some embodiments, the vaccine is administered to the individual via a route selected from the group consisting of intramuscular administration, intradermal administration and subcutaneous administration. In some embodiments, the administering step comprises contacting a muscle tissue of the subject with a device suitable for injection of the composition. In some embodiments, the administering step comprises contacting a muscle tissue of the subject with a device suitable for injection of the composition in combination with electroporation.

Aspects of the invention provide methods of vaccinating a subject comprising administering to the subject a single dosage of between 25 ug/kg and 400 ug/kg of a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a first antigenic polypeptide in an effective amount to vaccinate the subject.

Other aspects provide nucleic acid vaccines comprising one or more RNA polynucleotides having an open reading frame comprising at least one chemical modification, the open reading frame encoding a first antigenic polypeptide, wherein the vaccine has at least 10 fold less RNA polynucleotide than is required for an unmodified mRNA vaccine to produce an equivalent antibody titer. In some embodiments, the RNA polynucleotide is present in a dosage of 25-100 micrograms.

Other aspects provide nucleic acid vaccines comprising an LNP formulated RNA polynucleotide having an open reading frame comprising no nucleotide modifications (unmodified), the open reading frame encoding a first antigenic

polypeptide, wherein the vaccine has at least 10 fold less RNA polynucleotide than is required for an unmodified mRNA vaccine not formulated in a LNP to produce an equivalent antibody titer. In some embodiments, the RNA polynucleotide is present in a dosage of 25-100 micrograms. 5

The data presented in the Examples demonstrate significant enhanced immune responses using the formulations of the invention. Both chemically modified and unmodified RNA vaccines are useful according to the invention. Surprisingly, in contrast to prior art reports that it was preferable 10 to use chemically unmodified mRNA formulated in a carrier for the production of vaccines, it is described herein that chemically modified mRNA-LNP vaccines required a much lower effective mRNA dose than unmodified mRNA, i.e., tenfold less than unmodified mRNA when formulated in 15 carriers other than LNP. Both the chemically modified and unmodified RNA vaccines of the invention produce better immune responses than mRNA vaccines formulated in a different lipid carrier.

In other aspects the invention encompasses a method of 20 treating an elderly subject age 60 years or older comprising administering to the subject a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a respiratory virus antigenic polypeptide in an effective amount to vaccinate the subject.

In other aspects the invention encompasses a method of treating a young subject age 17 years or younger comprising administering to the subject a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a respiratory virus antigenic polypeptide in an effective amount to vaccinate the subject.

In other aspects the invention encompasses a method of treating an adult subject comprising administering to the subject a nucleic acid vaccine comprising one or more RNA polynucleotides having an open reading frame encoding a 35 respiratory virus antigenic polypeptide in an effective amount to vaccinate the subject.

In some aspects the invention is a method of vaccinating a subject with a combination vaccine including at least two nucleic acid sequences encoding respiratory antigens 40 wherein the dosage for the vaccine is a combined therapeutic dosage wherein the dosage of each individual nucleic acid encoding an antigen is a sub therapeutic dosage. In some embodiments, the combined dosage is 25 micrograms of the RNA polynucleotide in the nucleic acid vaccine adminis- 45 tered to the subject. In some embodiments, the combined dosage is 100 micrograms of the RNA polynucleotide in the nucleic acid vaccine administered to the subject. In some embodiments the combined dosage is 50 micrograms of the RNA polynucleotide in the nucleic acid vaccine adminis- 50 tered to the subject. In some embodiments, the combined dosage is 75 micrograms of the RNA polynucleotide in the nucleic acid vaccine administered to the subject. In some embodiments, the combined dosage is 150 micrograms of the RNA polynucleotide in the nucleic acid vaccine admin- 55 istered to the subject. In some embodiments, the combined dosage is 400 micrograms of the RNA polynucleotide in the nucleic acid vaccine administered to the subject. In some embodiments, the sub therapeutic dosage of each individual nucleic acid encoding an antigen is 1, 2, 3, 4, 5, 6, 7, 8, 9, 60 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 micrograms. In other embodiments the nucleic acid vaccine is chemically modified and in other embodiments the nucleic acid vaccine is not chemically modified.

The RNA polynucleotide is one of SEQ ID NO: 1-4, 9-12, 65 20-23, 35-46, 57-61, and 64-80 and includes at least one chemical modification. In other embodiments the RNA

polynucleotide is one of SEQ ID NO: 1-4, 9-12, 20-23, 35-46, 57-61, and 64-80 and does not include any nucleotide modifications, or is unmodified. In yet other embodiments the at least one RNA polynucleotide encodes an antigenic protein of any of SEQ ID NO: 5-8, 12-13, 24-34, and 47-50 and includes at least one chemical modification. In other embodiments the RNA polynucleotide encodes an antigenic protein of any of SEQ ID NO: 5-8, 12-13, 24-34, and 47-50 and does not include any nucleotide modifications, or is

22

In preferred aspects, vaccines of the invention (e.g., LNP-encapsulated mRNA vaccines) produce prophylactically- and/or therapeutically-efficacious levels, concentrations and/or titers of antigen-specific antibodies in the blood or serum of a vaccinated subject. As defined herein, the term antibody titer refers to the amount of antigen-specific antibody produces in s subject, e.g., a human subject. In exemplary embodiments, antibody titer is expressed as the inverse of the greatest dilution (in a serial dilution) that still gives a positive result. In exemplary embodiments, antibody titer is determined or measured by enzyme-linked immunosorbent assay (ELISA). In exemplary embodiments, antibody titer is determined or measured by neutralization assay, 25 e.g., by microneutralization assay. In certain aspects, antibody titer measurement is expressed as a ratio, such as 1:40, 1:100, etc.

In exemplary embodiments of the invention, an efficacious vaccine produces an antibody titer of greater than 1:40, greater that 1:100, greater than 1:400, greater than 1:1000, greater than 1:2000, greater than 1:3000, greater than 1:4000, greater than 1:500, greater than 1:6000, greater than 1:7500, greater than 1:10000. In exemplary embodiments, the antibody titer is produced or reached by 10 days following vaccination, by 20 days following vaccination, by 30 days following vaccination, or by 50 or more days following vaccination. In exemplary embodiments, the titer is produced or reached following a single dose of vaccine administered to the subject. In other embodiments, the titer is produced or reached following multiple doses, e.g., following a first and a second dose (e.g., a booster dose.)

In exemplary aspects of the invention, antigen-specific antibodies are measured in units of µg/ml or are measured in units of IU/L (International Units per liter) or mIU/ml (milli International Units per ml). In exemplary embodiments of the invention, an efficacious vaccine produces >0.5 μg/ml,  $>0.1 \mu g/ml$ ,  $>0.2 \mu g/ml$ ,  $>0.35 \mu g/ml$ ,  $>0.5 \mu g/ml$ ,  $>1 \mu g/ml$ ,  $>2 \mu g/ml$ ,  $>5 \mu g/ml$  or  $>10 \mu g/ml$ . In exemplary embodiments of the invention, an efficacious vaccine produces >10 mIU/ml, >20 mIU/ml, >50 mIU/ml, >100 mIU/ml, >200 mIU/ml, >500 mIU/ml or >1000 mIU/ml. In exemplary embodiments, the antibody level or concentration is produced or reached by 10 days following vaccination, by 20 days following vaccination, by 30 days following vaccination, by 40 days following vaccination, or by 50 or more days following vaccination. In exemplary embodiments, the level or concentration is produced or reached following a single dose of vaccine administered to the subject. In other embodiments, the level or concentration is produced or reached following multiple doses, e.g., following a first and a second dose (e.g., a booster dose.) In exemplary embodiments, antibody level or concentration is determined or measured by enzyme-linked immunosorbent assay (ELISA). In exemplary embodiments, antibody level or concentration is determined or measured by neutralization assay, e.g., by microneutralization assay.

canavalin A was included. The cytokines tested include IL-10 (FIG. 8A), TNF-α (FIG. 8B), IL4 (FIG. 8C), IL-5 (FIG. 8D) and IL-6 (FIG. 8E).

24

The details of various embodiments of the disclosure are set forth in the description below. Other features, objects, and advantages of the disclosure will be apparent from the description and from the claims.

## BRIEF DESCRIPTION OF THE DRAWINGS

The foregoing and other objects, features and advantages will be apparent from the following description of particular embodiments of the disclosure, as illustrated in the accompanying drawings in which like reference characters refer to the same parts throughout the different views. The drawings are not necessarily to scale, emphasis instead being placed upon illustrating the principles of various embodiments of the disclosure.

FIG. 1 shows a schematic of one example of a RNA (e.g. mRNA) vaccine construct of the present disclosure. The construct depicts a human *Metapneumovirus* and human respiratory syncytial virus full length fusion protein obtained from wild-type strains (*The Journal of General Virology.* 20 2008; 89(Pt 12): 3113-3118, incorporated herein by reference).

FIGS. 2A-2C are graphs showing the levels of anti-hMPV fusion protein-specific antibodies in the serum of mice immunized with hMPV mRNA vaccines on day 0 (FIG. 2A), 25 day 14 (FIG. 2B) and day 35 (FIG. 2C) post immunization. The mice were immunized with a single dose (2  $\mu$ g or 10  $\mu$ g) on day 0 and were given a boost dose (2  $\mu$ g or 10  $\mu$ g) on day 21, hMPV fusion protein-specific antibodies were detected at up to 1:10000 dilution of serum on day 35 for both doses. 30

FIGS. 3A-3C are graphs showing the result of IgG isotyping in the serum of mice immunized with hMPV mRNA vaccines. The levels of hMPV fusion protein-specific IgG2a (FIG. 3A) and IgG1 (FIG. 3B) antibodies in the serum are measured by ELISA. FIG. 3C shows that hMPV fusion 35 protein mRNA vaccine induced a mixed Th1/Th2 cytokine response with a Th1 bias.

FIG. **4** is a graph showing in vitro neutralization of a hMPV B2 strain (TN/91-316) using the sera of mice immunized with a mRNA vaccine encoding hMPV fusion protein. 40 Mouse serum obtained from mice receiving a 10 µg or a 2 µg dose contained hMPV-neutralizing antibodies.

FIGS. 5A-5C are graphs showing a Th1 cytokine response induced by a hMPV fusion peptide pool (15-mers-50 (overlap)) in splenocytes isolated from mice immunized with the 45 hMPV mRNA vaccines. Virus-free media was used as a negative control and Concanavalin A (ConA, a positive control for splenocyte stimulation) was included. The cytokines tested included IFN- $\gamma$  (FIG. 5A), IL-2 (FIG. 5B) and IL12 (FIG. 5C).

FIGS. 6A-6E are graphs showing the Th2 cytokine response induced by a hMPV fusion peptide pool (15-mers-50) in splenocytes isolated from mice immunized with the hMPV mRNA vaccines. Virus-free media was used as a negative control and Concanavalin A was also included. The 55 cytokines tested included IL-10 (FIG. 6A), TNF-α (FIG. 6B), IL4 (FIG. 6C), IL-5 (FIG. 6D) and IL-6 (FIG. 6E).

FIGS. 7A-7C are graphs showing the Th1 response induced by inactivated hMPV virus in splenocytes isolated from mice immunized with hMPV mRNA vaccines. Virusfree media was used as a negative control and Concanavalin A was included. The cytokines tested included IFN-γ (FIG. 7A), IL-2 (FIG. 7B) and IL12 (FIG. 7C).

FIGS. 8A-8E are graphs showing the Th2 response induced by inactivated hMPV virus in splenocytes isolated from mice immunized with the hMPV mRNA vaccines. Virus-free media was used as a negative control and Con-

FIGS. 9A-9B are graphs showing the results of cotton rat challenge experiments. Two different doses of the hMPV mRNA vaccines were used (2 μg or 10 μg doses) to immunize the cotton rats before challenge. The hMPV mRNA vaccines reduced the viral titer in the lung and nose of the cotton rat, with the 10 μg dose being more effective in reducing viral titer. Use of a 10 μg dose resulted in 100% protection in the lung and a 2 log reduction in nose viral titer. Use of a 2 μg dose resulted in a 1 log reduction in lung vital titer and no reduction in nose viral titer. The vaccine was administered on Day 0, and a boost was administered on Day 21.

FIG. 10 is a graph showing the lung histopathology of cotton rats that received hMPV mRNA vaccines. Pathology associated with vaccine-enhanced disease was not observed in immunized groups.

FIG. 11 is a graph showing hMPV neutralization antibody titers in cotton rats that received hMPV mRNA vaccines (2  $\mu$ g or 10  $\mu$ g doses) on days 35 and 42 post immunization.

FIG. 12 is a graph showing the lung and nose viral load in cotton rats challenged with a hMPV/A2 strain after immunization with the indicated mRNA vaccines (hMPV mRNA vaccine or hMPV/PIV mRNA combination vaccine). Vaccinated cotton rats showed reduced lung and nose viral loads after challenge, compared to control.

FIG. 13 is a graph showing the lung and nose viral load in cotton rats challenged with PIV3 strain after immunization with indicated mRNA vaccines (PIV mRNA vaccine or hMPV/PIV combination vaccine). Vaccinated cotton rats showed reduced lung and nose viral loads after challenge, compared to control.

FIG. 14 is a graph showing hMPV neutralizing antibody titers in cotton rats that received different dosages of hMPV mRNA vaccines or hMPV/PIV combination mRNA vaccines on day 42 post immunization. The dosages of the vaccine are indicated in Table 9.

FIG. 15 is a graph showing PIV3 neutralizing antibody titers in cotton rats that received different dosages of PIV mRNA vaccines or hMPV/PIV combination mRNA vaccines on day 42 post immunization. The dosages of the vaccine are indicated in Table 9.

FIG. 16 is a graph showing the lung histopathology score of cotton rats immunized with hMPV mRNA vaccines, PIV mRNA vaccines or hMPV/PIV combination mRNA vaccines as indicated in Table 9. Low occurrence of alevolitis and interstitial pneumonia was observed, indicating no antibody-dependent enhancement (ADE) of hMPV associated diseases.

FIG. 17 is a graph showing the reciprocal MERS-CoV neutralizing antibody titers in mice immunized with *Beta-coronavirus* mRNA vaccine encoding the MERS-CoV full-length Spike protein, on days 0, 21, 42, and 56 post immunization.

FIG. 18 is a graph showing the reciprocal MERS-CoV neutralizing antibody titers in mice immunized with *Beta-coronavirus* mRNA vaccine encoding either the MERS-CoV full-length Spike protein, or the S2 subunit of the Spike protein. The full length spike protein induced a stronger immune response compared to the S2 subunit alone.

FIGS. 19A-19C are graphs showing the viral load in the nose and throat, the bronchoalveolar lavage (BAL), or the lungs of New Zealand white rabbits 4 days post challenge with MERS-CoV. The New Zealand white rabbits were immunized with one 20 µg-dose (on day 0) or two 20

µg-doses (on day 0 and 21) of MERS-CoV mRNA vaccine encoding the full-length Spike protein before challenge. FIG. 19A shows that two doses of MERS-CoV mRNA vaccine resulted in a 3 log reduction of viral load in the nose and led to complete protection in the throat of the New Zealand white rabbits. FIG. 19B shows that two doses of MERS-CoV mRNA vaccine resulted in a 4 log reduction of viral load in the BAL of the New Zealand white rabbits. FIG. 19C show one dose of MERS-CoV mRNA vaccine resulted in a 2 log reduction of viral load, while two doses of MERS-CoV mRNA vaccine resulted in an over 4 log reduction of viral load in the lungs of the New Zealand white rabbits

FIGS. **20**A-**20**B are images and graphs showing viral load or replicating virus detected by PCR in the lungs of New Zealand white rabbits 4 days post challenge with MERS-CoV. The New Zealand white rabbits were immunized with a single 20 μg dose (on day 0, Group 1a) of MERS-CoV mRNA vaccine encoding the full-length Spike protein, two 20 μg doses (on day 0 and 21, Group 1b) of MERS-CoV mRNA vaccine encoding the full-length Spike protein, or placebo (Group 2) before challenge. FIG. **20**A shows that two doses of 20 μg a MERS-CoV mRNA vaccine reduced over 99% (2 log) of viruses in the lungs of New Zealand white rabbits. FIG. **20**B shows that the group of New Zealand white rabbits that received 2 doses of 20 μg MERS-CoV mRNA vaccine did not have any detectable replicating MERS-CoV virus in their lungs.

FIG. 21 is a graph showing the MERS-CoV neutralizing 30 antibody titers in New Zealand white rabbits immunized with MERS-CoV mRNA vaccine encoding the full-length Spike protein. Immunization of the in New Zealand white rabbits were carried out as described in FIGS. 21A-21C. The results show that two doses of 20  $\mu$ g MERS-CoV mRNA 35 vaccine induced a significant amount of neutralizing antibodies against MERS-CoV (EC50 between 500-1000). The MERS-CoV mRNA vaccine induced antibody titer is 3-5 fold better than any other vaccines tested in the same model.

## DETAILED DESCRIPTION

The present disclosure provides, in some embodiments, vaccines that comprise RNA (e.g., mRNA) polynucleotides encoding a human Metapneumovirus (hMPV) antigenic 45 polypeptide, a parainfluenza virus type 3 (PIV3) antigenic polypeptide, a respiratory syncytial virus (RSV) antigenic polypeptide, a measles virus (MeV) antigenic polypeptide, or a Betacoronavirus antigenic polypeptide (e.g., Middle East respiratory syndrome coronavirus (MERS-CoV), 50 SARS-CoV, human coronavirus (HCoV)-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH (New Haven) and HCoV-HKU1) (see, e.g., Esper F. et al. Emerging Infectious Diseases, 12(5), 2006; and Pyrc K. et al. Journal of Virology, 81(7):3051-57, 2007, the contents of each of 55 which is here incorporated by reference in their entirety). The present disclosure also provides, in some embodiments, combination vaccines that comprise at least one RNA (e.g., mRNA) polynucleotide encoding at least two antigenic polypeptides selected from hMPV antigenic polypeptides, 60 PIV3 antigenic polypeptides, RSV antigenic polypeptides, MeV antigenic polypeptides and BetaCoV antigenic polypeptides. Also provided herein are methods of administering the RNA (e.g., mRNA) vaccines, methods of producing the RNA (e.g., mRNA) vaccines, compositions (e.g., pharma- 65 ceutical compositions) comprising the RNA (e.g., mRNA) vaccines, and nucleic acids (e.g., DNA) encoding the RNA

26

(e.g., mRNA) vaccines. In some embodiments, a RNA (e.g., mRNA) vaccine comprises an adjuvant, such as a flagellin adjuvant, as provided herein.

The RNA (e.g., mRNA) vaccines (e.g., hMPV, PIV3, RSV, MeV, BetaCoV RNA vaccines and combinations thereof), in some embodiments, may be used to induce a balanced immune response, comprising both cellular and humoral immunity, without many of the risks associated with DNA vaccination.

The entire contents of International Application No. PCT/US2015/02740 is incorporated herein by reference. Human *Metapneumovirus* (hMPV)

hMPV shares substantial homology with respiratory syncytial virus (RSV) in its surface glycoproteins. hMPV fusion protein (F) is related to other paramyxovirus fusion proteins and appears to have homologous regions that may have similar functions. The hMPV fusion protein amino acid sequence contains features characteristic of other paramyxovirus F proteins, including a putative cleavage site and potential N-linked glycosylation sites. Paramyxovirus fusion proteins are synthesized as inactive precursors (F0) that are cleaved by host cell proteases into the biologically fusion-active F1 and F2 domains (see, e.g., Cseke G. et al. Journal of Virology 2007; 81(2):698-707, incorporated herein by reference). hMPV has one putative cleavage site, in contrast to the two sites established for RSV F, and only shares 34% amino acid sequence identity with RSV F. F2 is extracellular and disulfide linked to F1. Fusion proteins are type I glycoproteins existing as trimers, with two 4-3 heptad repeat domains at the N- and C-terminal regions of the protein (HR1 and HR2), which form coiled-coil alphahelices. These coiled coils become apposed in an antiparallel fashion when the protein undergoes a conformational change into the fusogenic state. There is a hydrophobic fusion peptide N proximal to the N-terminal heptad repeat, which is thought to insert into the target cell membrane. while the association of the heptad repeats brings the trans-40 membrane domain into close proximity, inducing membrane fusion (see, e.g., Baker, K A et al. Mol. Cell 1999; 3:309-319). This mechanism has been proposed for a number of different viruses, including RSV, influenza virus, and human immunodeficiency virus. Fusion proteins are major antigenic determinants for all known paramyxoviruses and for other viruses that possess similar fusion proteins such as human immunodeficiency virus, influenza virus, and Ebola

In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV fusion protein (F). In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding a F1 or F2 subunit of a hMPV F protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV glycoprotein (G). In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV matrix protein (M). In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV phosphoprotein (P). In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV nucleoprotein (N). In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding hMPV SH protein (SH).

In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein, M protein, P protein, N protein and SH protein.

In some embodiments, a hMPV vaccine of the present 5 disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and G protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and M protein. In some embodiments, a hMPV vaccine of the 10 present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and P protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and N protein. In some embodiments, a hMPV 15 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and SH protein.

In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and M protein. In some embodiments, a 20 hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and P protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and N protein. In some 25 embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and SH protein.

In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 30 encoding F protein, G protein and M protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and P protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA 35 (e.g., mRNA) polynucleotide encoding F protein, G protein and N protein. In some embodiments, a hMPV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and SH protein.

A hMPV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one hMPV antigenic polypeptide identified by any one of SEQ ID NO: 5-8 (Table 3; see also amino acid sequences of Table 4).

A hMPV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide encoded by a nucleic acid (e.g., DNA) identified by any one of SEQ ID NO: 1-4 (Table 2).

The present disclosure is not limited by a particular strain of hMPV. The strain of hMPV used in a vaccine may be any strain of hMPV. Non-limiting examples of strains of hMPV for use as provide herein include the CAN98-75 (CAN75) and the CAN97-83 (CAN83) hMPV strains (Skiadopoulos M H et al. *J Virol.* 20014; 78(13)6927-37, incorporated 55 herein by reference), a hMPV A1, A2, B1 or B2 strain (see, e.g., de Graaf M et al. *The Journal of General Virology* 2008; 89:975-83; Peret T C T et al. *The Journal of Infectious Disease* 2002; 185:1660-63, incorporated herein by reference), a hMPV isolate TN/92-4 (e.g., SEQ ID NO: 1 and 5), 60 a hMPV isolate NL/1/99 (e.g., SEQ ID NO: 2 and 6), or a hMPV isolate PER/CFI0497/2010/B (e.g., SEQ ID NO: 3 and 7).

In some embodiments, at least one hMPV antigenic polypeptide is obtained from a hMPV A1, A2, B1 or B2 65 strain (see, e.g., de Graaf M et al. *The Journal of General Virology* 2008; 89:975-83; Peret T C T et al. *The Journal of* 

28

Infectious Disease 2002; 185:1660-63, incorporated herein by reference). In some embodiments, at least one antigenic polypeptide is obtained from the CAN98-75 (CAN75) hMPV strain. In some embodiments, at least one antigenic polypeptide is obtained from the CAN97-83 (CAN83) hMPV strain. In some embodiments, at least one antigenic polypeptide is obtained from hMPV isolate TN/92-4 (e.g., SEQ ID NO: 1 and 5). In some embodiments, at least one antigenic polypeptide is obtained from hMPV isolate NL/1/99 (e.g., SEQ ID NO: 2 and 6). In some embodiments, at least one antigenic polypeptide is obtained from hMPV isolate PER/CFI0497/2010/B (e.g., SEQ ID NO: 3 and 7).

In some embodiments, hMPV vaccines comprise RNA (e.g., mRNA) polynucleotides encoding a hMPV antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with hMPV F protein and having F protein activity.

A protein is considered to have F protein activity if, for example, the protein acts to fuse the viral envelope and host cell plasma membrane, mediates viral entry into a host cell via an interaction with arginine-glycine-aspartate RGD-binding integrins, or a combination thereof (see, e.g., Cox R G et al. *J Virol.* 2012; 88(22):12148-60, incorporated herein by reference).

In some embodiments, hMPV vaccines comprise RNA (e.g., mRNA) polynucleotides encoding hMPV antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with hMPV G protein and having G protein activity.

A protein is considered to have G protein activity if, for example, the protein acts to modulate (e.g., inhibit) hMPV-induced cellular (immune) responses (see, e.g., Bao X et al. *PLoS Pathog.* 2008; 4(5):e1000077, incorporated herein by reference).

Human parainfluenza virus type 3 (PIV3)

Parainfluenza viruses belong to the family Paramyxoviridae. These are enveloped viruses with a negative-sense single-stranded RNA genome. Parainfluenza viruses belong to the subfamily Paramyxoviridae, which is subdivided into 40 three genera: Respirovirus (PIV-1, PIV-3, and Sendai virus (SeV)), Rubulavirus (PIV-2, PIV-4 and mumps virus) and Morbillivirus (measles virus, rinderpest virus and canine distemper virus (CDV)). Their genome, a ~15 500 nucleotide-long negative-sense RNA molecule, encodes two envelope glycoproteins, the hemagglutinin-neuraminidase (HN), the fusion protein (F or F0), which is cleaved into F1 and F2 subunits, a matrix protein (M), a nucleocapsid protein (N) and several nonstructural proteins including the viral replicase (L). All parainfluenza viruses, except for PIV-1, express a non-structural V protein that blocks IFN signaling in the infected cell and acts therefore as a virulence factor (see, e.g., Nishio M et al. J Virol. 2008; 82(13):6130-38).

PIV3 hemagglutinin-neuraminidase (HN), a structural protein, is found on the viral envelope, where it is necessary for attachment and cell entry. It recognizes and binds to sialic acid-containing receptors on the host cell's surface. As a neuroaminidase, HN removes sialic acid from virus particles, preventing self-aggregation of the virus, and promoting the efficient spread of the virus. Furthermore, HN promotes the activity of the fusion (F or F0) protein, contributing to the penetration of the host cell's surface.

PIV3 fusion protein (PIV3 F) is located on the viral envelope, where it facilitates the viral fusion and cell entry. The F protein is initially inactive, but proteolytic cleavage leads to its active forms, F1 and F2, which are linked by disulfide bonds. This occurs when the HN protein binds its receptor on the host cell's surface. During early phases of

infection, the F glycoprotein mediates penetration of the host cell by fusion of the viral envelope to the plasma membrane. In later stages of the infection, the F protein facilitates the fusion of the infected cells with neighboring uninfected cells, which leads to the formation of a syncytium 5 and spread of the infection.

PIV3 matrix protein (M) is found within the viral envelope and assists with viral assembly. It interacts with the nucleocapsid and envelope glycoproteins, where it facilitates the budding of progeny viruses through its interactions with specific sites on the cytoplasmic tail of the viral glycoproteins and nucleocapsid. It also plays a role in transporting viral components to the budding site.

PIV3 phosphoprotein (P) and PIV3 large polymerase protein (L) are found in the nucleocapsid where they form 15 part of the RNA polymerase complex. The L protein, a viral RNA-dependent RNA polymerase, facilitates genomic transcription, while the host cell's ribosomes translate the viral mRNA into viral proteins.

PIV3 V is a non-structural protein that blocks IFN signaling in the infected cell, therefore acting as a virulence factor.

PIV3 nucleoprotein (N) encapsidates the genome in a ratio of 1 N per 6 ribonucleotides, protecting it from nucleases. The nucleocapsid (NC) has a helical structure. 25 The encapsidated genomic RNA is termed the NC and serves as template for transcription and replication. During replication, encapsidation by PIV3 N is coupled to RNA synthesis and all replicative products are resistant to nucleases. PIV3 N homo-multimerizes to form the nucleocapsid 30 and binds to viral genomic RNA. PIV3 N binds the P protein and thereby positions the polymerase on the template.

In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding PIV3 fusion protein (F). In some embodiments, a 35 PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding a F1 or F2 subunit of a PIV3 F protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding PIV3 hemagglutinin-neuramini- 40 dase (HN) (see, e.g., van Wyke Coelingh K L et al. J Virol. 1987; 61(5):1473-77, incorporated herein by reference). In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding PIV3 matrix protein (M). In some embodiments, a PIV3 45 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding PIV3 phosphoprotein (P). In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding PIV3 nucleoprotein (N).

In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, HN protein, M protein, P protein, and N protein.

In some embodiments, a PIV3 vaccine of the present 55 disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and HN protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and M protein. In some embodiments, a PIV3 vaccine of the 60 present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and P protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and N protein.

In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide

30

encoding HN protein and M protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HN protein and P protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HN protein and N protein.

In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, HN protein and M protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, HN protein and P protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, HN protein and N protein.

A PIV3 vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one PIV3 antigenic polypeptide identified by any one of SEQ ID NO: 12-13 (Table 6; see also amino acid sequences of Table 7).

A PIV3 vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide encoded by a nucleic acid (e.g., DNA) identified by any one of SEQ ID NO: 9-12 (Table 5; see also nucleic acid sequences of Table 7).

The present disclosure is not limited by a particular strain of PIV3. The strain of PIV3 used in a vaccine may be any strain of PIV3. A non-limiting example of a strain of PIV3 for use as provide herein includes HPIV3/Homo sapiens/PER/FLA4815/2008.

In some embodiments, PIV3 vaccines comprise RNA (e.g., mRNA) polynucleotides encoding a PIV3 antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with PIV3 F protein and having F protein activity.

In some embodiments, PIV3 vaccines comprise RNA (e.g., mRNA) polynucleotides encoding PIV3 antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with PIV3 hemagglutinin-neuraminidase (HN) and having hemagglutinin-neuraminidase activity.

A protein is considered to have hemagglutinin-neuraminidase activity if, for example, it is capable of both receptor binding and receptor cleaving. Such proteins are major surface glycoproteins that have functional sites for cell attachment and for neuraminidase activity. They are able to cause red blood cells to agglutinate and to cleave the glycosidic linkages of neuraminic acids, so they have the potential to both bind a potential host cell and then release the cell if necessary, for example, to prevent self-aggregation of the virus.

In some embodiments, PIV3 vaccines comprise RNA (e.g., mRNA) polynucleotides encoding PIV3 antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with PIV3 HN, F (e.g., F, F1 or F2), M, N, L or V and having HN, F (e.g., F, F1 or F2), M, N, L or V activity, respectively. Respiratory Syncytial Virus (RSV)

RSV is a negative-sense, single-stranded RNA virus of the genus *Pneumovirinae*. The virus is present in at least two antigenic subgroups, known as Group A and Group B, primarily resulting from differences in the surface G glycoproteins. Two RSV surface glycoproteins—G and F—mediate attachment with and attachment to cells of the respiratory epithelium. F surface glycoproteins mediate coalescence of neighboring cells. This results in the formation of syncytial cells. RSV is the most common cause of bronchiolitis. Most infected adults develop mild cold-like

symptoms such as congestion, low-grade fever, and wheezing. Infants and small children may suffer more severe symptoms such as bronchiolitis and pneumonia. The disease may be transmitted among humans via contact with respiratory secretions.

The genome of RSV encodes at least three surface glycoproteins, including F, G, and SH, four nucleocapsid proteins, including L, P, N, and M2, and one matrix protein, M. Glycoprotein F directs viral penetration by fusion between the virion and the host membrane. Glycoprotein G is a type II transmembrane glycoprotein and is the major attachment protein. SH is a short integral membrane protein. Matrix protein M is found in the inner layer of the lipid bilayer and assists virion formation. Nucleocapsid proteins L, P, N, and M2 modulate replication and transcription of the RSV 15 genome. It is thought that glycoprotein G tethers and stabilizes the virus particle at the surface of bronchial epithelial cells, while glycoprotein F interacts with cellular glycosaminoglycans to mediate fusion and delivery of the RSV virion contents into the host cell (Krzyzaniak M A et al. PLoS 20 of RSV. The strain of RSV used in a vaccine may be any Pathog 2013; 9(4)).

In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) 25 polynucleotide encoding G protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding L protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding P protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding N protein. In some embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding M2 protein. In some 35 embodiments, a PIV3 vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding M

In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 40 encoding F protein, G protein, L protein, P protein, N protein, M2 protein and M protein.

In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and G protein. In some embodiments, a 45 RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and L protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and P protein. In some embodiments, a 50 RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and N protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and M2 protein. In some embodiments, 55 a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and M protein.

In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 60 encoding G protein and L protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and P protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 65 encoding G protein and N protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA

32

(e.g., mRNA) polynucleotide encoding G protein and M2 protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding G protein and M protein.

In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and L protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and P protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and N protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and M2 protein. In some embodiments, a RSV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein, G protein and M protein.

The present disclosure is not limited by a particular strain

In some embodiments, RSV vaccines comprise RNA (e.g., mRNA) polynucleotides encoding a RSV antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with RSV F protein and having F protein activity.

In some embodiments, RSV vaccines comprise RNA (e.g., mRNA) polynucleotides encoding RSV antigenic polypeptides having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with RSV G protein and having G protein activity.

A protein is considered to have G protein activity if, for example, the protein acts to modulate (e.g., inhibit) hMPVinduced cellular (immune) responses (see, e.g., Bao X et al. PLoS Pathog. 2008; 4(5):e1000077, incorporated herein by reference).

Measles Virus (MeV)

Molecular epidemiologic investigations and virologic surveillance contribute notably to the control and prevention of measles. Nearly half of measles-related deaths worldwide occur in India, yet virologic surveillance data are incomplete for many regions of the country. Previous studies have documented the presence of measles virus genotypes D4, D7, and D8 in India, and genotypes D5, D9, D11, H1, and G3 have been detected in neighboring countries. Recently, MeV genotype B3 was detected in India (Kuttiatt V S et al. Emerg Infect Dis. 2014; 20(10): 1764-66).

The glycoprotein complex of paramyxoviruses mediates receptor binding and membrane fusion. In particular, the MeV fusion (F) protein executes membrane fusion, after receptor binding by the hemagglutinin (HA) protein (Muhlebach M D et al. Journal of Virology 2008; 82(22):11437-45). The MeV P gene codes for three proteins: P, an essential polymerase cofactor, and V and C, which have multiple functions but are not strictly required for viral propagation in cultured cells. V shares the amino-terminal domain with P but has a zinc-binding carboxyl-terminal domain, whereas C is translated from an overlapping reading frame. The MeV C protein is an infectivity factor. During replication, the P protein binds incoming monomeric nucleocapsid (N) proteins with its amino-terminal domain and positions them for assembly into the nascent ribonucleocapsid. The P protein amino-terminal domain is natively unfolded (Deveaux P et al. Journal of Virology 2004; 78(21):11632-40).

In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein. In some embodiments, a MeV vaccine

of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding P protein. In some embodiments, a MeV vaccine of the present disclosure 5 comprises a RNA (e.g., mRNA) polynucleotide encoding V protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding C protein.

In some embodiments, a MeV vaccine of the present 10 disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein, F protein, P protein, V protein and C protein.

In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 15 encoding HA protein and F protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein and P protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide 20 encoding HA protein and V protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein and C protein.

some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and P protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and V protein. In some embodiments, a MeV vaccine of the present disclosure 30 comprises a RNA (e.g., mRNA) polynucleotide encoding F protein and C protein.

In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein, F protein and P protein. In some 35 embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein, F protein and V protein. In some embodiments, a MeV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding HA protein, F protein and C protein.

In some embodiments, MeV vaccines comprise RNA (e.g., mRNA) encoding a MeV antigenic polypeptide having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with MeV HA protein and having MeV 45 HA protein activity.

In some embodiments, MeV vaccines comprise RNA (e.g., mRNA) encoding a MeV antigenic polypeptide having at least 95%, at least 96%, at least 97%, at least 98% or at least 99% identity with MeV F protein and having MeV F 50 protein activity.

A protein is considered to have HA protein activity if the protein mediates receptor binding and/or membrane fusion. MeV F protein executes membrane fusion, after receptor binding by the MeV HA protein.

A MeV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one MeV antigenic polypeptide identified by any one of SEQ ID NO: 47-50 (Table 14; see also amino acid sequences of Table 15).

A MeV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide identified by any one of SEQ ID NO: 37, 40, 43, 46 (Table 13).

A MeV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide encoded by a nucleic 65 acid (e.g., DNA) identified by any one of SEQ ID NO: 35, 36, 38, 39, 41, 42, 44 and 45 (Table 13).

34

The present disclosure is not limited by a particular strain of MeV. The strain of MeV used in a vaccine may be any strain of MeV. Non-limiting examples of strains of MeV for use as provide herein include B3/B3.1, C2, D4, D6, D7, D8, G3, H1, Moraten, Rubeovax, MVi/New Jersey.USA/45.05, MVi/Texas.USA/4.07, AIK-C, MVi/New York.USA/26.09/3, MVi/California.USA/16.03, MVi/Virginia.USA/15.09, MVi/California.USA/8.04, and MVi/Pennsylvania.USA/20.09.

MeV proteins may be from MeV genotype D4, D5, D7, D8, D9, D11, H1, G3 or B3. In some embodiments, a MeV HA protein or a MeV F protein is from MeV genotype D8. In some embodiments, a MeV HA protein or a MeV F protein is from MeV genotype B3.

Betacoronaviruses (BetaCoV)

MERS-Co V. MERS-CoV is a positive-sense, singlestranded RNA virus of the genus Betacoronavirus. The genomes are phylogenetically classified into two clades, clade A and clade B. It has a strong tropism for non-ciliated bronchial epithelial cells, evades the innate immune response and antagonizes interferon (IFN) production in infected cells. Dipeptyl peptidase 4 (DDP4, also known as CD26) has been identified as a functional cellular receptor for MERS-CoV. Its enzymatic activity is not required for infection, although its amino acid sequence is highly conserved across species and is expressed in the human bronchial epithelium and kidneys. Most infected individuals develop severe acute respiratory illnesses, including fever, cough, and shortness of breath, and the virus can be fatal. The disease may be transmitted among humans, generally among those in close contact.

The genome of MERS-CoV encodes at least four unique accessory proteins, such as 3, 4a, 4b and 5, two replicase proteins (open reading frame 1a and 1b), and four major structural proteins, including spike (S), envelope (E), nucleocapsid (N), and membrane (M) proteins (Almazan F et al. MBio 2013; 4(5):e00650-13). The accessory proteins play nonessential roles in MERS-CoV replication, but they are likely structural proteins or interferon antagonists, modulating in vivo replication efficiency and/or pathogenesis, as in the case of SARS-CoV (Almazan F et al. MBio 2013; 4(5):e00650-13; Totura A L et al. Curr Opin Virol 2012; 2(3):264-75; Scobey T et al. Proc Natl Acad Sci USA 2013; 110(40):16157-62). The other proteins of MERS-CoV maintain different functions in virus replication. The E protein, for example, involves in virulence, and deleting the E-coding gene results in replication-competent and propagation-defective viruses or attenuated viruses (Almazan F et al. MBio 2013; 4(5):e00650-13). The S protein is particularly essential in mediating virus binding to cells expressing receptor dipeptidyl peptidase-4 (DPP4) through receptorbinding domain (RBD) in the S1 subunit, whereas the S2 subunit subsequently mediates virus entry via fusion of the virus and target cell membranes (Li F. J Virol 2015; 89(4): 55 1954-64; Raj V S et al. Nature 2013; 495(7440):251-4).

In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding the S1 subunit of the S protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding the S2 subunit of the S protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding E protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a

RNA (e.g., mRNA) polynucleotide encoding N protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding M protein.

In some embodiments, a MERS-CoV vaccine of the 5 present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), E protein, N protein and M protein.

In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and E protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and N protein. In some embodiments, a MERS-CoV vaccine of the 15 present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and M protein.

In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) poly-20 nucleotide encoding S protein (S, S1 and/or S2), E protein and M protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), E protein and N protein. In some embodiments, a 25 MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), M protein and N protein. In some embodiments, a MERS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding E 30 protein, M protein and N protein.

A MERS-CoV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one MERS-CoV antigenic polypeptide identified by any one of SEQ ID NO: 24-38 or 35 33 (Table 11; see also amino acid sequences of Table 12).

A MERS-CoV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide encoded by a nucleic acid (e.g., DNA) identified by any one of SEQ ID NO: 20-23 (Table 10).

The present disclosure is not limited by a particular strain of MERS-CoV. The strain of MERS-CoV used in a vaccine may be any strain of MERS-CoV. Non-limiting examples of strains of MERS-CoV for use as provide herein include Riyadh\_14\_2013, and 2cEMC/2012, Hasa\_1\_2013.

SARS-CoV. The genome of SARS-CoV includes of a single, positive-strand RNA that is approximately 29,700 nucleotides long. The overall genome organization of SARS-CoV is similar to that of other coronaviruses. The reference genome includes 13 genes, which encode at least 50 14 proteins. Two large overlapping reading frames (ORFs) encompass 71% of the genome. The remainder has 12 potential ORFs, including genes for structural proteins S (spike), E (small envelope), M (membrane), and N (nucleocapsid). Other potential ORFs code for unique putative 55 SARS-CoV-specific polypeptides that lack obvious sequence similarity to known proteins. A detailed analysis of the SARS-CoV genome has been published in *J Mol Biol* 2003; 331: 991-1004.

In some embodiments, a SARS-CoV vaccine of the 60 present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), E protein, N protein and M protein.

In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and E protein. In some embodiments, a SARS-CoV vaccine of the

36

present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and N protein. In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2) and M protein.

In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), E protein and M protein. In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), E protein and N protein. In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding S protein (S, S1 and/or S2), M protein and N protein. In some embodiments, a SARS-CoV vaccine of the present disclosure comprises a RNA (e.g., mRNA) polynucleotide encoding E protein, M protein and N protein.

A SARS-CoV vaccine may comprise, for example, at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one SARS-CoV antigenic polypeptide identified by any one of SEQ ID NO: 29, 32 or 34 (Table 11; see also amino acid sequences of Table 12).

The present disclosure is not limited by a particular strain of SARS-CoV. The strain of SARS-CoV used in a vaccine may be any strain of SARS-CoV.

HCoV-OC43. Human coronavirus OC43 is an enveloped, positive-sense, single-stranded RNA virus in the species Betacoronavirus-1 (genus Betacoronavirus, subfamily Coronavirinae, family Coronaviridae, order Nidovirales). Four HCoV-OC43 genotypes (A to D), have been identified with genotype D most likely arising from recombination. The complete genome sequencing of two genotype C and D strains and bootscan analysis shows recombination events between genotypes B and C in the generation of genotype D. Of 29 strains identified, none belong to the more ancient genotype A. Along with HCoV-229E, a species in the Alphacoronavirus genus, HCoV-OC43 are among the known viruses that cause the common cold. Both viruses can cause severe lower respiratory tract infections, including pneumonia in infants, the elderly, and immunocompromised individuals such as those undergoing chemotherapy and those with HIV-AIDS.

HCoV-HKU1. Human coronavirus HKU1 (HCoV-HKU1) is a positive-sense, single-stranded RNA virus with the HE gene, which distinguishes it as a group 2, or Betacoronavirus. It was discovered in January 2005 in two patients in Hong Kong. The genome of HCoV-HKU1 is a 29,926-nucleotide, polyadenylated RNA. The GC content is 32%, the lowest among all known coronaviruses. The genome organization is the same as that of other group II coronaviruses, with the characteristic gene order 1a, 1b, HE, S, E, M, and N. Furthermore, accessory protein genes are present between the S and E genes (ORF4) and at the position of the N gene (ORF8). The TRS is presumably located within the AAUCUAAAC sequence, which precedes each ORF except E. As in sialodacryoadenitis virus and mouse hepatitis virus (MHV), translation of the E protein possibly occurs via an internal ribosomal entry site. The 3' untranslated region contains a predicted stem-loop structure immediately downstream of the N ORF (nucleotide position 29647 to 29711). Further downstream, a pseudoknot structure is present at nucleotide position 29708 to 29760. Both RNA structures are conserved in group II coronaviruses and are critical for virus replication.

HCoV-NL63. The RNA genome of human coronavirus NL63 (HCoV-NL63) is 27,553 nucleotides, with a poly(A) tail (FIG. 1). With a GC content of 34%, HCoV-NL63 has one of the lowest GC contents of the coronaviruses, for which GC content ranges from 32 to 42%. Untranslated 5 regions of 286 and 287 nucleotides are present at the 5' and 3' termini, respectively. Genes predicted to encode the S, E, M, and N proteins are found in the 3' part of the HCoV-NL63 genome. The HE gene, which is present in some group II coronaviruses, is absent, and there is only a single, mono- 10 cistronic accessory protein ORF (ORF3) located between the S and E genes. Subgenomic mRNAs are generated for all ORFs (S, ORF3, E, M, and N), and the core sequence of the TRS of HCoV-NL63 is defined as AACUAAA. This sequence is situated upstream of every ORF except for the 15 E ORF, which contains the suboptimal core sequence AAC-UAUA. Interestingly, a 13-nucleotide sequence with perfect homology to the leader sequence is situated upstream of the suboptimal E TRS. Annealing of this 13-nucleotide sequence to the leader sequence may act as a compensatory 20 mechanism for the disturbed leader-TRS/body-TRS interac-

HCoV-229E. Human coronavirus 229E (HCoV-229E) is a single-stranded, positive-sense, RNA virus species in the Alphacoronavirus genus of the subfamily Coronavirinae, in 25 the family Coronaviridae, of the order Nidovirales. Along with Human coronavirus OC43, it is responsible for the common cold. HCoV-NL63 and HCoV-229E are two of the four human coronaviruses that circulate worldwide. These two viruses are unique in their relationship towards each 30 other. Phylogenetically, the viruses are more closely related to each other than to any other human coronavirus, yet they only share 65% sequence identity. Moreover, the viruses use different receptors to enter their target cell. HCoV-NL63 is associated with croup in children, whereas all signs suggest 35 that the virus probably causes the common cold in healthy adults. HCoV-229E is a proven common cold virus in healthy adults, so it is probable that both viruses induce comparable symptoms in adults, even though their mode of infection differs (HCoV-NL63 and HCoV-229E are two of 40 the four human coronaviruses that circulate worldwide. These two viruses are unique in their relationship towards each other. Phylogenetically, the viruses are more closely related to each other than to any other human coronavirus, yet they only share 65% sequence identity. Moreover, the 45 viruses use different receptors to enter their target cell. HCoV-NL63 is associated with croup in children, whereas all signs suggest that the virus probably causes the common cold in healthy adults. HCoV-229E is a proven common cold virus in healthy adults, so it is probable that both viruses 50 induce comparable symptoms in adults, even though their mode of infection differs (Dijkman R. et al. J Formos Med Assoc. 2009 April; 108(4):270-9, the contents of which is incorporated herein by reference in their entirety). Combination Vaccines

Embodiments of the present disclosure also provide combination RNA (e.g., mRNA) vaccines. A "combination RNA (e.g., mRNA) vaccine" of the present disclosure refers to a vaccine comprising at least one (e.g., at least 2, 3, 4, or 5) RNA (e.g., mRNA) polynucleotide having an open reading 60 frame encoding a combination of any two or more (or all of) antigenic polypeptides selected from hMPV antigenic polypeptides, PIV3 antigenic polypeptides, RSV antigenic polypeptides, MeV antigenic polypeptides, and BetaCoV antigenic polypeptides (e.g., selected from MERS-CoV, SARS-65 CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

38

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide, a MeV antigenic polypeptide, a MeV antigenic polypeptide, and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide and a PIV3 antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide and a RSV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide and a BetaCoV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a PIV3 antigenic polypeptide and a RSV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a PIV3 antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a PIV3 antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a RSV antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a RSV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a MeV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, 55 HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide, a RSV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide, a MeV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, 5 SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

39

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a RSV antigenic 10 polypeptide, a MeV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) 15 vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a PIV3 antigenic polypeptide, a RSV antigenic polypeptide, a MeV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, 20 HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide and a RSV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) 30 vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a PIV3 antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and 35 HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a RSV antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a RSV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, 45 HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a hMPV antigenic polypeptide, a MeV antigenic 50 polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) 55 vaccine comprises a RNA (e.g., mRNA) polynucleotide encoding a PIV3 antigenic polypeptide, a RSV antigenic polypeptide and a MeV antigenic polypeptide.

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide 60 encoding a PIV3 antigenic polypeptide, a RSV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

In some embodiments, a combination RNA (e.g., mRNA) vaccine comprises a RNA (e.g., mRNA) polynucleotide

40

encoding a RSV antigenic polypeptide, a MeV antigenic polypeptide and a BetaCoV antigenic polypeptide (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1).

Other combination respiratory virus RNA (e.g., mRNA) vaccines are encompassed by the present disclosure.

It has been discovered that the mRNA vaccines described herein are superior to current vaccines in several ways. First, the lipid nanoparticle (LNP) delivery is superior to other formulations including a protamine base approach described in the literature and no additional adjuvants are to be necessary. The use of LNPs enables the effective delivery of chemically modified or unmodified mRNA vaccines. Additionally it has been demonstrated herein that both modified and unmodified LNP formulated mRNA vaccines were superior to conventional vaccines by a significant degree. In some embodiments the mRNA vaccines of the invention are superior to conventional vaccines by a factor of at least 10 fold, 20 fold, 40 fold, 50 fold, 100 fold, 500 fold or 1,000 fold

Although attempts have been made to produce functional RNA vaccines, including mRNA vaccines and self-replicating RNA vaccines, the therapeutic efficacy of these RNA vaccines have not yet been fully established. Quite surprisingly, the inventors have discovered, according to aspects of the invention a class of formulations for delivering mRNA vaccines in vivo that results in significantly enhanced, and in many respects synergistic, immune responses including enhanced antigen generation and functional antibody production with neutralization capability. These results can be achieved even when significantly lower doses of the mRNA are administered in comparison with mRNA doses used in other classes of lipid based formulations. The formulations of the invention have demonstrated significant unexpected in vivo immune responses sufficient to establish the efficacy of functional mRNA vaccines as prophylactic and therapeutic agents. Additionally, self-replicating RNA vaccines rely on viral replication pathways to deliver enough RNA to a cell 40 to produce an immunogenic response. The formulations of the invention do not require viral replication to produce enough protein to result in a strong immune response. Thus, the mRNA of the invention are not self-replicating RNA and do not include components necessary for viral replication.

The invention involves, in some aspects, the surprising finding that lipid nanoparticle (LNP) formulations significantly enhance the effectiveness of mRNA vaccines, including chemically modified and unmodified mRNA vaccines. The efficacy of mRNA vaccines formulated in LNP was examined in vivo using several distinct antigens. The results presented herein demonstrate the unexpected superior efficacy of the mRNA vaccines formulated in LNP over other commercially available vaccines.

In addition to providing an enhanced immune response, the formulations of the invention generate a more rapid immune response with fewer doses of antigen than other vaccines tested. The mRNA-LNP formulations of the invention also produce quantitatively and qualitatively better immune responses than vaccines formulated in a different carriers.

The data described herein demonstrate that the formulations of the invention produced significant unexpected improvements over existing antigen vaccines. Additionally, the mRNA-LNP formulations of the invention are superior to other vaccines even when the dose of mRNA is lower than other vaccines. Mice immunized with either 10  $\mu g$  or 2  $\mu g$  doses of an hMPV fusion protein mRNA LNP vaccine or a

PIV3 mRNA LNP vaccine produced neutralizing antibodies which for instance, successfully neutralized the hMPV B2 virus. A 10 μg dose of mRNA vaccine protected 100% of mice from lethal challenge and drastically reduced the viral titer after challenge (~2 log reduction).

41

Two 20  $\mu g$  doses of MERS-CoV mRNA LNP vaccine significantly reduced viral load and induced significant amount of neutralizing antibodies against MERS-CoV (EC $_{50}$  between 500-1000). The MERS-CoV mRNA vaccine induced antibody titer was 3-5 fold better than any other 10 vaccines tested in the same model.

The LNP used in the studies described herein has been used previously to deliver siRNA in various animal models as well as in humans. In view of the observations made in association with the siRNA delivery of LNP formulations, 15 the fact that LNP is useful in vaccines is quite surprising. It has been observed that therapeutic delivery of siRNA formulated in LNP causes an undesirable inflammatory response associated with a transient IgM response, typically leading to a reduction in antigen production and a compromised immune response. In contrast to the findings observed with siRNA, the LNP-mRNA formulations of the invention are demonstrated herein to generate enhanced IgG levels, sufficient for prophylactic and therapeutic methods rather than transient IgM responses.

Nucleic Acids/Polynucleotides

Respiratory virus vaccines, as provided herein, comprise at least one (one or more) ribonucleic acid (RNA) (e.g., mRNA) polynucleotide having an open reading frame encoding at least one antigenic polypeptide selected from 30 hMPV, PIV3, RSV, MeV and BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1) antigenic polypeptides. The term "nucleic acid" includes any compound and/or substance that comprises a polymer of 35 nucleotides (nucleotide monomer). These polymers are referred to as polynucleotides. Thus, the terms "nucleic acid" and "polynucleotide" are used interchangeably.

Nucleic acids may be or may include, for example, ribonucleic acids (RNAs), deoxyribonucleic acids (DNAs), 40 threose nucleic acids (TNAs), glycol nucleic acids (GNAs), peptide nucleic acids (PNAs), locked nucleic acids (LNAs, including LNA having a  $\beta$ -D-ribo configuration,  $\alpha$ -LNA having an  $\alpha$ -L-ribo configuration (a diastereomer of LNA), 2'-amino-LNA having a 2'-amino functionalization, and 45 2'-amino- $\alpha$ -LNA having a 2'-amino functionalization), ethylene nucleic acids (ENA), cyclohexenyl nucleic acids (CeNA) or chimeras or combinations thereof.

In some embodiments, polynucleotides of the present disclosure function as messenger RNA (mRNA). "Messen- 50 ger RNA" (mRNA) refers to any polynucleotide that encodes a (at least one) polypeptide (a naturally-occurring, non-naturally-occurring, or modified polymer of amino acids) and can be translated to produce the encoded polypeptide in vitro, in vivo, in situ or ex vivo. The skilled artisan 55 will appreciate that, except where otherwise noted, polynucleotide sequences set forth in the instant application will recite "T"s in a representative DNA sequence but where the sequence represents RNA (e.g., mRNA), the "T"s would be substituted for "U"s. Thus, any of the RNA polynucleotides 60 encoded by a DNA identified by a particular sequence identification number may also comprise the corresponding RNA (e.g., mRNA) sequence encoded by the DNA, where each "T" of the DNA sequence is substituted with "U."

The basic components of an mRNA molecule typically 65 include at least one coding region, a 5' untranslated region (UTR), a 3' UTR, a 5' cap and a poly-A tail. Polynucleotides

42

of the present disclosure may function as mRNA but can be distinguished from wild-type mRNA in their functional and/or structural design features, which serve to overcome existing problems of effective polypeptide expression using nucleic-acid based therapeutics.

In some embodiments, a RNA polynucleotide of an RNA (e.g., mRNA) vaccine encodes 2-10, 2-9, 2-8, 2-7, 2-6, 2-5, 2-4, 2-3, 3-10, 3-9, 3-8, 3-7, 3-6, 3-5, 3-4, 4-10, 4-9, 4-8, 4-7, 4-6, 4-5, 5-10, 5-9, 5-8, 5-7, 5-6, 6-10, 6-9, 6-8, 6-7, 7-10, 7-9, 7-8, 8-10, 8-9 or 9-10 antigenic polypeptides. In some embodiments, a RNA (e.g., mRNA) polynucleotide of a respiratory virus vaccine encodes at least 10, 20, 30, 40, 50, 60, 70, 80, 90 or 100 antigenic polypeptides. In some embodiments, a RNA (e.g., mRNA) polynucleotide of a respiratory virus vaccine encodes at least 100 or at least 200 antigenic polypeptides. In some embodiments, a RNA polynucleotide of an respiratory virus vaccine encodes 1-10, 5-15, 10-20, 15-25, 20-30, 25-35, 30-40, 35-45, 40-50, 1-50, 1-100, 2-50 or 2-100 antigenic polypeptides.

Polynucleotides of the present disclosure, in some embodiments, are codon optimized. Codon optimization methods are known in the art and may be used as provided herein. Codon optimization, in some embodiments, may be used to match codon frequencies in target and host organ-25 isms to ensure proper folding; bias GC content to increase mRNA stability or reduce secondary structures; minimize tandem repeat codons or base runs that may impair gene construction or expression; customize transcriptional and translational control regions; insert or remove protein trafficking sequences; remove/add post translation modification sites in encoded protein (e.g. glycosylation sites); add, remove or shuffle protein domains; insert or delete restriction sites; modify ribosome binding sites and mRNA degradation sites; adjust translational rates to allow the various domains of the protein to fold properly; or to reduce or eliminate problem secondary structures within the polynucleotide. Codon optimization tools, algorithms and services are known in the art—non-limiting examples include services from GeneArt (Life Technologies), DNA2.0 (Menlo Park Calif.) and/or proprietary methods. In some embodiments, the open reading frame (ORF) sequence is optimized using optimization algorithms.

In some embodiments, a codon optimized sequence shares less than 95% sequence identity, less than 90% sequence identity, less than 85% sequence identity, less than 80% sequence identity, or less than 75% sequence identity to a naturally-occurring or wild-type sequence (e.g., a naturally-occurring or wild-type mRNA sequence encoding a polypeptide or protein of interest (e.g., an antigenic protein or antigenic polypeptide)).

In some embodiments, a codon-optimized sequence shares between 65% and 85% (e.g., between about 67% and about 85%, or between about 67% and about 80%) sequence identity to a naturally-occurring sequence or a wild-type sequence (e.g., a naturally-occurring or wild-type mRNA sequence encoding a polypeptide or protein of interest (e.g., an antigenic protein or polypeptide)). In some embodiments, a codon-optimized sequence shares between 65% and 75%, or about 80% sequence identity to a naturally-occurring sequence or wild-type sequence (e.g., a naturally-occurring or wild-type mRNA sequence encoding a polypeptide or protein of interest (e.g., an antigenic protein or polypeptide)).

In some embodiments a codon-optimized RNA (e.g., mRNA) may, for instance, be one in which the levels of G/C are enhanced. The G/C-content of nucleic acid molecules may influence the stability of the RNA. RNA having an

increased amount of guanine (G) and/or cytosine (C) residues may be functionally more stable than nucleic acids containing a large amount of adenine (A) and thymine (T) or uracil (U) nucleotides. WO02/098443 discloses a pharmaceutical composition containing an mRNA stabilized by sequence modifications in the translated region. Due to the degeneracy of the genetic code, the modifications work by substituting existing codons for those that promote greater RNA stability without changing the resulting amino acid. The approach is limited to coding regions of the RNA. Antigens/Antigenic Polypeptides

In some embodiments, an antigenic polypeptide (e.g., a hMPV, PIV3, RSV, MeV or BetaCoV antigenic polypeptide) is longer than 25 amino acids and shorter than 50 amino 15 acids. Polypeptides include gene products, naturally occurring polypeptides, synthetic polypeptides, homologs, orthologs, paralogs, fragments and other equivalents, variants, and analogs of the foregoing. A polypeptide may be a single molecule or may be a multi-molecular complex such 20 as a dimer, trimer or tetramer. Polypeptides may also comprise single chain polypeptides or multichain polypeptides, such as antibodies or insulin, and may be associated or linked to each other. Most commonly, disulfide linkages are found in multichain polypeptides. The term "polypeptide" 25 may also apply to amino acid polymers in which at least one amino acid residue is an artificial chemical analogue of a corresponding naturally-occurring amino acid.

A "polypeptide variant" is a molecule that differs in its amino acid sequence relative to a native sequence or a 30 reference sequence. Amino acid sequence variants may possess substitutions, deletions, insertions, or a combination of any two or three of the foregoing, at certain positions within the amino acid sequence, as compared to a native sequence or a reference sequence. Ordinarily, variants possess at least 50% identity to a native sequence or a reference sequence. In some embodiments, variants share at least 80% identity or at least 90% identity with a native sequence or a reference sequence.

In some embodiments "variant mimics" are provided. A 40 "variant mimic" contains at least one amino acid that would mimic an activated sequence. For example, glutamate may serve as a mimic for phosphoro-threonine and/or phosphoro-serine. Alternatively, variant mimics may result in deactivation or in an inactivated product containing the mimic. For 45 example, phenylalanine may act as an inactivating substitution for tyrosine, or alanine may act as an inactivating substitution for serine.

"Orthologs" refers to genes in different species that evolved from a common ancestral gene by speciation. Nor- 50 mally, orthologs retain the same function in the course of evolution. Identification of orthologs is important for reliable prediction of gene function in newly sequenced genomes.

"Analogs" is meant to include polypeptide variants that 55 differ by one or more amino acid alterations, for example, substitutions, additions or deletions of amino acid residues that still maintain one or more of the properties of the parent or starting polypeptide.

The present disclosure provides several types of compositions that are polynucleotide or polypeptide based, including variants and derivatives. These include, for example, substitutional, insertional, deletion and covalent variants and derivatives. The term "derivative" is synonymous with the term "variant" and generally refers to a molecule that has 65 been modified and/or changed in any way relative to a reference molecule or a starting molecule.

44

As such, polynucleotides encoding peptides or polypeptides containing substitutions, insertions and/or additions, deletions and covalent modifications with respect to reference sequences, in particular the polypeptide sequences disclosed herein, are included within the scope of this disclosure. For example, sequence tags or amino acids, such as one or more lysines, can be added to peptide sequences (e.g., at the N-terminal or C-terminal ends). Sequence tags can be used for peptide detection, purification or localization. Lysines can be used to increase peptide solubility or to allow for biotinylation. Alternatively, amino acid residues located at the carboxy and amino terminal regions of the amino acid sequence of a peptide or protein may optionally be deleted providing for truncated sequences. Certain amino acids (e.g., C-terminal residues or N-terminal residues) alternatively may be deleted depending on the use of the sequence, as for example, expression of the sequence as part of a larger sequence that is soluble, or linked to a solid support.

"Substitutional variants" when referring to polypeptides are those that have at least one amino acid residue in a native or starting sequence removed and a different amino acid inserted in its place at the same position. Substitutions may be single, where only one amino acid in the molecule has been substituted, or they may be multiple, where two or more (e.g., 3, 4 or 5) amino acids have been substituted in the same molecule.

As used herein the term "conservative amino acid substitution" refers to the substitution of an amino acid that is normally present in the sequence with a different amino acid of similar size, charge, or polarity. Examples of conservative substitutions include the substitution of a non-polar (hydrophobic) residue such as isoleucine, valine and leucine for another non-polar residue. Likewise, examples of conservative substitutions include the substitution of one polar (hydrophilic) residue for another such as between arginine and lysine, between glutamine and asparagine, and between glycine and serine. Additionally, the substitution of a basic residue such as lysine, arginine or histidine for another, or the substitution of one acidic residue such as aspartic acid or glutamic acid for another acidic residue are additional examples of conservative substitutions. Examples of nonconservative substitutions include the substitution of a nonpolar (hydrophobic) amino acid residue such as isoleucine, valine, leucine, alanine, methionine for a polar (hydrophilic) residue such as cysteine, glutamine, glutamic acid or lysine and/or a polar residue for a non-polar residue.

"Features" when referring to polypeptide or polynucleotide are defined as distinct amino acid sequence-based or nucleotide-based components of a molecule respectively. Features of the polypeptides encoded by the polynucleotides include surface manifestations, local conformational shape, folds, loops, half-loops, domains, half-domains, sites, termini and any combination(s) thereof.

As used herein when referring to polypeptides the term "domain" refers to a motif of a polypeptide having one or more identifiable structural or functional characteristics or properties (e.g., binding capacity, serving as a site for protein-protein interactions).

As used herein when referring to polypeptides the terms "site" as it pertains to amino acid based embodiments is used synonymously with "amino acid residue" and "amino acid side chain." As used herein when referring to polynucleotides the terms "site" as it pertains to nucleotide based embodiments is used synonymously with "nucleotide." A site represents a position within a peptide or polypeptide or

polynucleotide that may be modified, manipulated, altered, derivatized or varied within the polypeptide-based or polynucleotide-based molecules.

As used herein the terms "termini" or "terminus" when referring to polypeptides or polynucleotides refers to an 5 extremity of a polypeptide or polynucleotide respectively. Such extremity is not limited only to the first or final site of the polypeptide or polynucleotide but may include additional amino acids or nucleotides in the terminal regions. Polypeptide-based molecules may be characterized as hav- 10 ing both an N-terminus (terminated by an amino acid with a free amino group (NH2)) and a C-terminus (terminated by an amino acid with a free carboxyl group (COOH)). Proteins are in some cases made up of multiple polypeptide chains brought together by disulfide bonds or by non-covalent 15 forces (multimers, oligomers). These proteins have multiple N- and C-termini. Alternatively, the termini of the polypeptides may be modified such that they begin or end, as the case may be, with a non-polypeptide based moiety such as an organic conjugate.

As recognized by those skilled in the art, protein fragments, functional protein domains, and homologous proteins are also considered to be within the scope of polypeptides of interest. For example, provided herein is any protein fragment (meaning a polypeptide sequence at least one amino 25 acid residue shorter than a reference polypeptide sequence but otherwise identical) of a reference protein having a length of 10, 20, 30, 40, 50, 60, 70, 80, 90, 100 or longer than 100 amino acids. In another example, any protein that includes a stretch of 20, 30, 40, 50, or 100 (contiguous) 30 amino acids that are 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 100% identical to any of the sequences described herein can be utilized in accordance with the disclosure. In some embodiments, a polypeptide includes 2, 3, 4, 5, 6, 7, 8, 9, 10, or more mutations as shown in any of the sequences pro- 35 vided herein or referenced herein. In another example, any protein that includes a stretch of 20, 30, 40, 50, or 100 amino acids that are greater than 80%, 90%, 95%, or 100% identical to any of the sequences described herein, wherein the protein has a stretch of 5, 10, 15, 20, 25, or 30 amino 40 acids that are less than 80%, 75%, 70%, 65% to 60% identical to any of the sequences described herein can be utilized in accordance with the disclosure.

Polypeptide or polynucleotide molecules of the present disclosure may share a certain degree of sequence similarity 45 or identity with the reference molecules (e.g., reference polypeptides or reference polynucleotides), for example, with art-described molecules (e.g., engineered or designed molecules or wild-type molecules). The term "identity," as known in the art, refers to a relationship between the 50 sequences of two or more polypeptides or polynucleotides, as determined by comparing the sequences. In the art, identity also means the degree of sequence relatedness between two sequences as determined by the number of matches between strings of two or more amino acid residues 55 or nucleic acid residues. Identity measures the percent of identical matches between the smaller of two or more sequences with gap alignments (if any) addressed by a particular mathematical model or computer program (e.g., "algorithms"). Identity of related peptides can be readily 60 calculated by known methods. "% identity" as it applies to polypeptide or polynucleotide sequences is defined as the percentage of residues (amino acid residues or nucleic acid residues) in the candidate amino acid or nucleic acid sequence that are identical with the residues in the amino 65 acid sequence or nucleic acid sequence of a second sequence after aligning the sequences and introducing gaps, if neces46

sary, to achieve the maximum percent identity. Methods and computer programs for the alignment are well known in the art. Identity depends on a calculation of percent identity but may differ in value due to gaps and penalties introduced in the calculation. Generally, variants of a particular polynucleotide or polypeptide have at least 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% but less than 100% sequence identity to that particular reference polynucleotide or polypeptide as determined by sequence alignment programs and parameters described herein and known to those skilled in the art. Such tools for alignment include those of the BLAST suite (Stephen F. Altschul, et al. (1997)." Gapped BLAST and PSI-BLAST: a new generation of protein database search programs," Nucleic Acids Res. 25:3389-3402). Another popular local alignment technique is based on the Smith-Waterman algorithm (Smith, T. F. & Waterman, M. S. (1981) "Identification of common molecular subsequences." J. Mol. Biol. 147:195-197). A general global alignment technique based on dynamic programming is the Needleman-Wunsch algorithm (Needleman, S. B. & Wunsch, C. D. (1970) "A general method applicable to the search for similarities in the amino acid sequences of two proteins." J. Mol. Biol. 48:443-453). More recently, a Fast Optimal Global Sequence Alignment Algorithm (FOGSAA) was developed that purportedly produces global alignment of nucleotide and protein sequences faster than other optimal global alignment methods, including the Needleman-Wunsch algorithm. Other tools are described herein, specifically in the definition of "identity" below.

As used herein, the term "homology" refers to the overall relatedness between polymeric molecules, e.g. between nucleic acid molecules (e.g. DNA molecules and/or RNA molecules) and/or between polypeptide molecules. Polymeric molecules (e.g. nucleic acid molecules (e.g. DNA molecules and/or RNA molecules) and/or polypeptide molecules) that share a threshold level of similarity or identity determined by alignment of matching residues are termed homologous. Homology is a qualitative term that describes a relationship between molecules and can be based upon the quantitative similarity or identity. Similarity or identity is a quantitative term that defines the degree of sequence match between two compared sequences. In some embodiments, polymeric molecules are considered to be "homologous" to one another if their sequences are at least 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, or 99% identical or similar. The term "homologous" necessarily refers to a comparison between at least two sequences (polynucleotide or polypeptide sequences). Two polynucleotide sequences are considered homologous if the polypeptides they encode are at least 50%, 60%, 70%, 80%, 90%, 95%, or even 99% for at least one stretch of at least 20 amino acids. In some embodiments, homologous polynucleotide sequences are characterized by the ability to encode a stretch of at least 4-5 uniquely specified amino acids. For polynucleotide sequences less than 60 nucleotides in length, homology is determined by the ability to encode a stretch of at least 4-5 uniquely specified amino acids. Two protein sequences are considered homologous if the proteins are at least 50%, 60%, 70%, 80%, or 90% identical for at least one stretch of at least 20 amino acids.

Homology implies that the compared sequences diverged in evolution from a common origin. The term "homolog" refers to a first amino acid sequence or nucleic acid sequence (e.g., gene (DNA or RNA) or protein sequence) that is related to a second amino acid sequence or nucleic acid sequence by descent from a common ancestral sequence.

(1984)), BLASTP, BLASTN, and FASTA Altschul, S. F. et al., *J. Molec. Biol.*, 215, 403 (1990)).

Multiprotein and Multicomponent Vaccines

The present disclosure encompasses respiratory virus vaccines comprising multiple RNA (e.g., mRNA) polynucleotides, each encoding a single antigenic polypeptide, as well as respiratory virus vaccines comprising a single RNA polynucleotide encoding more than one antigenic polypeptide (e.g., as a fusion polypeptide). Thus, a vaccine composition comprising a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a first antigenic polypeptide and a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a second antigenic polypeptide encompasses (a) vaccines that comprise a first RNA polynucleotide encoding a first antigenic polypeptide and a second RNA polynucleotide encoding a second antigenic polypeptide, and (b) vaccines that comprise a single RNA polynucleotide encoding a first and second antigenic polypeptide (e.g., as a fusion polypeptide). RNA (e.g., mRNA) vaccines of the present disclosure, in some embodiments, comprise 2-10 (e.g., 2, 3, 4, 5, 6, 7, 8, 9 or 10), or more, RNA polynucleotides having an open reading frame, each of which encodes a different antigenic polypeptide (or a single RNA polynucleotide encoding 2-10, or more, different antigenic polypeptides). The antigenic polypeptides may be selected from hMPV, PIV3, RSV, MEV and BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1) antigenic polypeptides.

In some embodiments, a respiratory virus vaccine comprises a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral capsid protein, a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral premembrane/membrane protein, and a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral envelope protein. In some embodiments, a respiratory virus vaccine comprises a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral fusion (F) protein and a RNA polynucleotide having an open reading frame encoding a viral major surface glycoprotein (G protein). In some embodiments, a vaccine comprises a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral F protein. In some embodiments, a vaccine comprises a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a viral G protein. In some embodiments, a vaccine comprises a RNA (e.g., mRNA) polynucleotide having an open reading frame encoding a HN protein.

In some embodiments, a multicomponent vaccine comprises at least one RNA (e.g., mRNA) polynucleotide encoding at least one antigenic polypeptide fused to a signal peptide (e.g., any one of SEQ ID NO: 15-19). The signal peptide may be fused at the N-terminus or the C-terminus of an antigenic polypeptide. An antigenic polypeptide fused to a signal peptide may be selected from hMPV, PIV3, RSV, MEV and BetaCoV (e.g., selected from MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and HCoV-HKU1) antigenic polypeptides. Signal Peptides

In some embodiments, antigenic polypeptides encoded by respiratory virus RNA (e.g., mRNA) polynucleotides comprise a signal peptide. Signal peptides, comprising the N-terminal 15-60 amino acids of proteins, are typically needed for the translocation across the membrane on the secretory pathway and, thus, universally control the entry of most proteins both in eukaryotes and prokaryotes to the secretory pathway. Signal peptides generally include three

The term "homolog" may apply to the relationship between genes and/or proteins separated by the event of speciation or to the relationship between genes and/or proteins separated by the event of genetic duplication. "Orthologs" are genes (or proteins) in different species that evolved from a common ancestral gene (or protein) by speciation. Typically, orthologs retain the same function in the course of evolution. "Paralogs" are genes (or proteins) related by duplication within a genome. Orthologs retain the same function in the course of evolution, whereas paralogs evolve new functions, even if these are related to the original one.

The term "identity" refers to the overall relatedness between polymeric molecules, for example, between polynucleotide molecules (e.g. DNA molecules and/or RNA molecules) and/or between polypeptide molecules. Calculation of the percent identity of two polynucleic acid sequences, for example, can be performed by aligning the two sequences for optimal comparison purposes (e.g., gaps can be introduced in one or both of a first and a second 20 nucleic acid sequences for optimal alignment and nonidentical sequences can be disregarded for comparison purposes). In certain embodiments, the length of a sequence aligned for comparison purposes is at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 80%, 25 at least 90%, at least 95%, or 100% of the length of the reference sequence. The nucleotides at corresponding nucleotide positions are then compared. When a position in the first sequence is occupied by the same nucleotide as the corresponding position in the second sequence, then the 30 molecules are identical at that position. The percent identity between the two sequences is a function of the number of identical positions shared by the sequences, taking into account the number of gaps, and the length of each gap, which needs to be introduced for optimal alignment of the 35 two sequences. The comparison of sequences and determination of percent identity between two sequences can be accomplished using a mathematical algorithm. For example, the percent identity between two nucleic acid sequences can be determined using methods such as those described in 40 Computational Molecular Biology, Lesk, A. M., ed., Oxford University Press, New York, 1988; Biocomputing: Informatics and Genome Projects, Smith, D. W., ed., Academic Press, New York, 1993; Sequence Analysis in Molecular Biology, von Heinje, G., Academic Press, 1987; Computer Analysis 45 of Sequence Data, Part I, Griffin, A. M., and Griffin, H. G., eds., Humana Press, New Jersey, 1994; and Sequence Analysis Primer, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991; each of which is incorporated herein by reference. For example, the percent identity 50 between two nucleic acid sequences can be determined using the algorithm of Meyers and Miller (CABIOS, 1989, 4:11-17), which has been incorporated into the ALIGN program (version 2.0) using a PAM120 weight residue table, a gap length penalty of 12 and a gap penalty of 4. The 55 percent identity between two nucleic acid sequences can, alternatively, be determined using the GAP program in the GCG software package using an NWSgapdna.CMP matrix. Methods commonly employed to determine percent identity between sequences include, but are not limited to those 60 disclosed in Carillo, H., and Lipman, D., SIAM J Applied Math., 48:1073 (1988); incorporated herein by reference. Techniques for determining identity are codified in publicly available computer programs. Exemplary computer software to determine homology between two sequences 65 include, but are not limited to, GCG program package, Devereux, J., et al., Nucleic Acids Research, 12(1), 387

48

regions: an N-terminal region of differing length, which usually comprises positively charged amino acids; a hydrophobic region; and a short carboxy-terminal peptide region. In eukaryotes, the signal peptide of a nascent precursor protein (pre-protein) directs the ribosome to the rough 5 endoplasmic reticulum (ER) membrane and initiates the transport of the growing peptide chain across it for processing. ER processing produces mature proteins, wherein the signal peptide is cleaved from precursor proteins, typically by a ER-resident signal peptidase of the host cell, or they 10 remain uncleaved and function as a membrane anchor. A signal peptide may also facilitate the targeting of the protein to the cell membrane. The signal peptide, however, is not responsible for the final destination of the mature protein. Secretory proteins devoid of additional address tags in their 15 sequence are by default secreted to the external environment. During recent years, a more advanced view of signal peptides has evolved, showing that the functions and immunodominance of certain signal peptides are much more versatile than previously anticipated.

Respiratory virus vaccines of the present disclosure may comprise, for example, RNA (e.g., mRNA) polynucleotides encoding an artificial signal peptide, wherein the signal peptide coding sequence is operably linked to and is in frame with the coding sequence of the antigenic polypeptide. Thus, 25 respiratory virus vaccines of the present disclosure, in some embodiments, produce an antigenic polypeptide comprising an antigenic polypeptide (e.g., hMPV, PIV3, RSV, MeV or BetaCoV) fused to a signal peptide. In some embodiments, a signal peptide is fused to the N-terminus of the antigenic 30 polypeptide. In some embodiments, a signal peptide is fused to the C-terminus of the antigenic polypeptide.

In some embodiments, the signal peptide fused to the antigenic polypeptide is an artificial signal peptide. In some embodiments, an artificial signal peptide fused to the anti- 35 genic polypeptide encoded by the RNA (e.g., mRNA) vaccine is obtained from an immunoglobulin protein, e.g., an IgE signal peptide or an IgG signal peptide. In some embodiments, a signal peptide fused to the antigenic polypeptide encoded by a RNA (e.g., mRNA) vaccine is an Ig 40 heavy chain epsilon-1 signal peptide (IgE HC SP) having the sequence of: MDWTWILFLVAAATRVHS (SEQ ID NO: 16). In some embodiments, a signal peptide fused to the antigenic polypeptide encoded by the (e.g., mRNA) RNA (e.g., mRNA) vaccine is an IgGk chain V-III region HAH 45 signal peptide (IgGk SP) having the sequence of MET-PAOLLFLLLWLPDTTG (SEO ID NO: 15). In some embodiments, the signal peptide is selected from: Japanese encephalitis PRM signal sequence (MLG-SNSGQRVVFTILLLLVAPAYS; SEQ ID NO: 17), VSVg 50 protein signal sequence (MKCLLYLAFLFIGVNCA; SEQ ID NO: 18) and Japanese encephalitis JEV signal sequence (MWLVSLAIVTACAGA; SEQ ID NO: 19)

In some embodiments, the antigenic polypeptide encoded by a RNA (e.g., mRNA) vaccine comprises an amino acid 55 sequence identified by any one of SEQ ID NO: 5-8, 12-13, 24-34, 47-50 or 54-56 (Tables 3, 6, 11, 14 or 17; see also amino acid sequences of Tables 4, 7, 12 or 15) fused to a signal peptide identified by any one of SEQ ID NO: 15-19 (Table 8). The examples disclosed herein are not meant to be 60 limiting and any signal peptide that is known in the art to facilitate targeting of a protein to ER for processing and/or targeting of a protein to the cell membrane may be used in accordance with the present disclosure.

A signal peptide may have a length of 15-60 amino acids. 65 For example, a signal peptide may have a length of 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,

50

33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, or 60 amino acids. In some embodiments, a signal peptide has a length of 20-60, 25-60, 30-60, 35-60, 40-60, 45-60, 50-60, 55-60, 15-55, 20-55, 25-55, 30-55, 35-55, 40-55, 45-55, 50-55, 15-50, 20-50, 25-50, 30-50, 35-50, 40-50, 45-50, 15-45, 20-45, 25-45, 30-45, 35-45, 40-45, 15-40, 20-40, 25-40, 30-40, 35-40, 15-35, 20-35, 25-35, 30-35, 15-30, 20-30, 25-30, 15-25, 20-25, or 15-20 amino acids.

A signal peptide is typically cleaved from the nascent polypeptide at the cleavage junction during ER processing. The mature antigenic polypeptide produce by a respiratory virus RNA (e.g., mRNA) vaccine of the present disclosure typically does not comprise a signal peptide.

Chemical Modifications

Respiratory virus vaccines of the present disclosure, in some embodiments, comprise at least RNA (e.g. mRNA) polynucleotide having an open reading frame encoding at least one antigenic polypeptide that comprises at least one chemical modification.

The terms "chemical modification" and "chemically modified" refer to modification with respect to adenosine (A), guanosine (G), uridine (U), thymidine (T) or cytidine (C) ribonucleosides or deoxyribnucleosides in at least one of their position, pattern, percent or population. Generally, these terms do not refer to the ribonucleotide modifications in naturally occurring 5'-terminal mRNA cap moieties. With respect to a polypeptide, the term "modification" refers to a modification relative to the canonical set 20 amino acids. Polypeptides, as provided herein, are also considered "modified" of they contain amino acid substitutions, insertions or a combination of substitutions and insertions.

Polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides), in some embodiments, comprise various (more than one) different modifications. In some embodiments, a particular region of a polynucleotide contains one, two or more (optionally different) nucleoside or nucleotide modifications. In some embodiments, a modified RNA polynucleotide (e.g., a modified mRNA polynucleotide), introduced to a cell or organism, exhibits reduced degradation in the cell or organism, respectively, relative to an unmodified polynucleotide. In some embodiments, a modified RNA polynucleotide (e.g., a modified mRNA polynucleotide), introduced into a cell or organism, may exhibit reduced immunogenicity in the cell or organism, respectively (e.g., a reduced innate response).

Modifications of polynucleotides include, without limitation, those described herein. Polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) may comprise modifications that are naturally-occurring, non-naturally-occurring or the polynucleotide may comprise a combination of naturally-occurring and non-naturally-occurring modifications. Polynucleotides may include any useful modification, for example, of a sugar, a nucleobase, or an internucleoside linkage (e.g., to a linking phosphate, to a phosphodiester linkage or to the phosphodiester backbone).

Polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides), in some embodiments, comprise non-natural modified nucleotides that are introduced during synthesis or post-synthesis of the polynucleotides to achieve desired functions or properties. The modifications may be present on an internucleotide linkages, purine or pyrimidine bases, or sugars. The modification may be introduced with chemical synthesis or with a polymerase enzyme at the terminal of a chain or anywhere else in the chain. Any of the regions of a polynucleotide may be chemically modified.

The present disclosure provides for modified nucleosides and nucleotides of a polynucleotide (e.g., RNA polynucleotides, such as mRNA polynucleotides). A "nucleoside" refers to a compound containing a sugar molecule (e.g., a pentose or ribose) or a derivative thereof in combination 5 with an organic base (e.g., a purine or pyrimidine) or a derivative thereof (also referred to herein as "nucleobase"). A nucleotide" refers to a nucleoside, including a phosphate group. Modified nucleotides may by synthesized by any useful method, such as, for example, chemically, enzymatically, or recombinantly, to include one or more modified or non-natural nucleosides. Polynucleotides may comprise a region or regions of linked nucleosides. Such regions may have variable backbone linkages. The linkages may be standard phosphdioester linkages, in which case the poly- 15 nucleotides would comprise regions of nucleotides.

Modified nucleotide base pairing encompasses not only the standard adenosine-thymine, adenosine-uracil, or guanosine-cytosine base pairs, but also base pairs formed between nucleotides and/or modified nucleotides comprising non-standard or modified bases, wherein the arrangement of hydrogen bond donors and hydrogen bond acceptors permits hydrogen bonding between a non-standard base and a standard base or between two complementary non-standard base structures. One example of such non-standard base pairing is the base pairing between the modified nucleotide inosine and adenine, cytosine or uracil. Any combination of base/sugar or linker may be incorporated into polynucleotides of the present disclosure.

Modifications of polynucleotides (e.g., RNA polynucle- 30 otides, such as mRNA polynucleotides) that are useful in the vaccines of the present disclosure include, but are not limited to the following: 2-methylthio-N6-(cis-hydroxyisopentenyl)adenosine; 2-methylthio-N6-methyladenosine; 2-methylthio-N6-threonyl carbamoyladenosine; N6-glyci- 35 nylcarbamovladenosine; N6-isopentenyladenosine; N6-methyladenosine; N6-threonylcarbamoyladeno sine; 1,2'-O-dimethyladenosine; 1-methyladenosine; 2'-O-methyladenosine; 2'-O-ribosyladenosine (phosphate); 2-methyladenosine; 2-methylthio-N6 isopentenyladenosine; 2-meth- 40 ylthio-N6-hydroxynorvalyl carbamoyladenosine; (phosphate); methyladenosine; 2'-O-ribosyladenosine Isopentenyladenosine; N6-(cis-hydroxyisopentenyl)adenosine; N6,2'-O-dimethyladenosine; N6,2'-O-dimethyladenosine; N6,N6,2'-O-trimethyladenosine; N6,N6-dimethylad- 45 N6-acetyladenosine; enosine. N6-hydroxynorvalylcarbamovladenosine: N6-methyl-N6threonylcarbamoyladenosine; 2-methyladenosine; 2-methylthio-N6-isopentenyladenosine; 7-deaza-adenosine; N1-methyl-adenosine; N6,N6 (dimethyl)adenine; N6-cis- 50 hydroxy-isopentenyl-adenosine; α-thio-adenosine; 2 (amino)adenine; 2 (aminopropyl)adenine; 2 (methylthio) N6 (isopentenyl)adenine; 2-(alkyl)adenine; 2-(aminoalkyl)adenine; 2-(aminopropyl)adenine; 2-(halo)adenine; 2-(halo) 2-(propyl)adenine; 2'-Amino-2'-deoxy-ATP; 55 2'-Azido-2'-deoxy-ATP; 2'-Deoxy-2'-a-aminoadenosine TP; 2'-Deoxy-2'-a-azidoadenosine TP; 6 (alkyl)adenine; 6 (methyl)adenine; 6-(alkyl)adenine; 6-(methyl)adenine; 7 (deaza)adenine; 8 (alkenyl)adenine; 8 (alkynyl)adenine; 8 (amino)adenine; 8 (thioalkyl)adenine; 8-(alkenyl)adenine; 60 8-(alkyl)adenine; 8-(alkynyl)adenine; 8-(amino)adenine; 8-(halo)adenine; 8-(hydroxyl)adenine; 8-(thioalkyl)adenine; 8-(thiol)adenine; 8-azido-adeno sine; aza adenine; deaza adenine; N6 (methyl)adenine; N6-(isopentyl)adenine; 7-deaza-8-aza-adenosine; 7-methyladenine; 1-Deazaade- 65 nosine TP; 2'Fluoro-N6-Bz-deoxyadenosine TP; 2'-OMe-2-Amino-ATP; 2'O-methyl-N6-Bz-deoxyadenosine TP; 2'-a52

Ethynyladenosine TP; 2-aminoadenine; 2-Aminoadenosine TP; 2-Amino-ATP; 2'-a-Trifluoromethyladenosine TP; 2-Azidoadenosine TP; 2'-b-Ethynyladenosine TP; 2-Bromoadenosine TP; 2'-b-Trifluoromethyladenosine TP; 2-Chloroadenosine TP; 2'-Deoxy-2',2'-difluoroadenosine TP; 2'-Deoxy-2'-a-mercaptoadenosine TP; 2'-Deoxy-2'-athiomethoxyadenosine TP; 2'-Deoxy-2'-b-aminoadenosine TP; 2'-Deoxy-2'-b-azidoadenosine TP; 2'-Deoxy-2'-b-bromoadenosine TP; 2'-Deoxy-2'-b-chloroadenosine TP; 2'-Deoxy-2'-b-fluoroadenosine TP: 2'-Deoxy-2'-b-iodoadenosine TP; 2'-Deoxy-2'-b-mercaptoadenosine TP; 2'-Deoxy-2'-bthiomethoxyadenosine TP; 2-Fluoroadenosine TP; 2-Iodoadenosine TP; 2-Mercaptoadenosine TP; 2-methoxy-adenine; 2-methylthio-adenine; 2-Trifluoromethyladenosine TP; 3-Deaza-3-bromoadenosine TP; 3-Deaza-3-chloroadenosine TP; 3-Deaza-3-fluoroadenosine TP; 3-Deaza-3-iodoadenosine TP; 3-Deazaadenosine TP; 4'-Azidoadenosine TP; 4'-Carbocyclic adenosine TP; 4'-Ethynyladenosine TP; 5'-Homo-adenosine TP; 8-Aza-ATP; 8-bromo-adenosine TP: 8-Trifluoromethyladenosine TP: 9-Deazaadenosine TP: 2-aminopurine; 7-deaza-2,6-diaminopurine; 7-deaza-8-aza-2,6-diaminopurine; 7-deaza-8-aza-2-aminopurine; 2,6-diaminopurine; 7-deaza-8-aza-adenine, 7-deaza-2-aminopurine; 2-thiocytidine; 3-methylcytidine; 5-formylcytidine; 5-hydroxymethylcytidine; 5-methylcytidine; N4-acetylcytidine; 2'-O-methylcytidine; 2'-O-methylcytidine; 5,2'-O-dimethylcytidine; 5-formyl-2'-O-methylcytidine; Lysidine; N4,2'-O-dimethylcytidine; N4-acetyl-2'-O-methylcytidine; N4-methylcytidine; N4,N4-Dimethyl-2'-OMe-Cytidine TP; 4-methylcytidine; 5-aza-cytidine; Pseudo-iso-cytidine; pyrrolo-cytidine; α-thio-cytidine; 2-(thio)cytosine; 2'-Amino-2'-deoxy-CTP; 2'-Azido-2'-deoxy-CTP; 2'-Deoxy-2'-a-aminocytidine TP; 2'-Deoxy-2'-a-azidocytidine TP; 3 (deaza) 5 (aza)cytosine; 3 (methyl)cytosine; 3-(alkyl)cytosine; 3-(deaza) 5 (aza)cytosine; 3-(methyl)cytidine; 4,2'-O-dimethylcytidine; 5 (halo)cytosine; 5 (methyl)cytosine; 5 (propynyl)cytosine; 5 (trifluoromethyl)cytosine; 5-(alkyl)cytosine; 5-(alkynyl)cytosine; 5-(halo)cytosine; 5-(propynyl) cytosine; 5-(trifluoromethyl)cytosine; 5-bromo-cytidine; 5-iodo-cytidine; 5-propynyl cytosine; 6-(azo)cytosine; 6-aza-cytidine; aza cytosine; deaza cytosine; N4 (acetyl) cytosine; 1-methyl-1-deaza-pseudoisocytidine; 1-methylpseudoisocytidine; 2-methoxy-5-methyl-cytidine; 2-methoxy-cytidine; 2-thio-5-methyl-cytidine; 4-methoxy-1-methyl-pseudoisocytidine; 4-methoxy-pseudoisocytidine; 4-thio-1-methyl-1-deaza-pseudoisocytidine; 4-thio-1methyl-pseudoisocytidine; 4-thio-pseudoisocytidine; 5-azazebularine; 5-methyl-zebularine; pyrrolo-pseudoisocytidine; Zebularine; (E)-5-(2-Bromo-vinyl)cytidine TP; 2,2'-anhydro-cytidine TP hydrochloride; 2'Fluor-N4-Bz-cytidine TP; 2'Fluoro-N4-Acetyl-cytidine TP; 2'-O-Methyl-N4-Acetyl-cytidine TP; 2'O-methyl-N4-Bz-cytidine TP; 2'-a-Ethynylcytidine TP; 2'-a-Trifluoromethylcytidine TP; 2'-b-Ethynylcytidine TP; 2'-b-Trifluoromethylcytidine 2'-Deoxy-2',2'-difluorocytidine TP; 2'-Deoxy-2'-a-mercaptocytidine TP; 2'-Deoxy-2'-a-thiomethoxycytidine TP; 2'-Deoxy-2'-b-aminocytidine TP; 2'-Deoxy-2'-b-azidocytidine TP; 2'-Deoxy-2'-b-bromocytidine TP; 2'-Deoxy-2'-bchlorocytidine TP; 2'-Deoxy-2'-b-fluorocytidine TP; 2'-Deoxy-2'-b-iodocytidine TP; 2'-Deoxy-2'-b-mercaptocytidine TP; 2'-Deoxy-2'-b-thiomethoxycytidine TP; 2'-O-Methyl-5-(1-propynyl)cytidine TP; 3'-Ethynylcytidine TP; 4'-Azidocytidine TP; 4'-Carbocyclic cytidine TP; 4'-Ethynylcytidine TP; 5-(1-Propynyl)ara-cytidine TP; 5-(2-Chloro-phenyl)-2thiocytidine TP; 5-(4-Amino-phenyl)-2-thiocytidine TP; 5-Aminoallyl-CTP; 5-Cyanocytidine TP; 5-Ethynylara-cytidine TP; 5-Ethynylcytidine TP; 5'-Homo-cytidine TP;

54

5-Methoxycytidine TP; 5-Trifluoromethyl-Cytidine TP; N4-Amino-cytidine TP; N4-Benzoyl-cytidine TP; Pseudoisocytidine; 7-methylguanosine; N2,2'-O-dimethylguanosine; N2-methylguanosine; Wyosine; 1,2'-O-dimethylguanosine; 1-methylguanosine; 2'-O-methylguanosine; 5 2'-O-ribosylguanosine (phosphate); 2'-O-methylguanosine; 2'-O-ribosylguanosine (phosphate); 7-aminomethyl-7-deazaguanosine; 7-cyano-7-deazaguanosine; Archaeosine; Methylwyo sine; N2,7-dimethylguanosine; N2,N2,2'-Otrimethylguanosine; N2.N2.7-trimethylguanosine; N2.N2- 10 dimethylguanosine; N2,7,2'-O-trimethylguanosine; 6-thioguanosine; 7-deaza-guanosine; 8-oxo-guanosine; N1-methyl-guanosine; α-thio-guanosine; 2 (propyl)guanine; 2-(alkyl)guanine; 2'-Amino-2'-deoxy-GTP; 2'-Azido-2'-deoxy-GTP; 2'-Deoxy-2'-a-aminoguanosine TP; 2'-Deoxy-2'- 15 a-azidoguanosine TP; 6 (methyl)guanine; 6-(alkyl)guanine; 6-(methyl)guanine; 6-methyl-guanosine; 7 (alkyl)guanine; 7 (deaza)guanine; 7 (methyl)guanine; 7-(alkyl)guanine; 7-(deaza)guanine; 7-(methyl)guanine; 8 (alkyl)guanine; 8 (alkynyl)guanine; 8 (halo)guanine; 8 (thioalkyl)guanine; 20 8-(alkenyl)guanine; 8-(alkyl)guanine; 8-(alkynyl)guanine; 8-(amino)guanine; 8-(halo)guanine; 8-(hydroxyl)guanine; 8-(thioalkyl)guanine; 8-(thiol)guanine; aza guanine; deaza guanine; N (methyl)guanine; N-(methyl)guanine; 1-methyl-6-thio-guanosine; 6-methoxy-guanosine; 6-thio-7-deaza-8- 25 aza-guanosine; 6-thio-7-deaza-guanosine; 6-thio-7-methyl-7-deaza-8-aza-guanosine; 7-methyl-8-oxoguanosine; guanosine; N2,N2-dimethyl-6-thio-guanosine; N2-methyl-6-thio-guanosine; 1-Me-GTP; 2'Fluoro-N2-isobutylguanosine TP; 2'O-methyl-N2-isobutyl-guanosine TP; 2'-a- 30 Ethynylguanosine TP; 2'-a-Trifluoromethylguanosine TP; 2'-b-Ethynylguano sine TP; 2'-b-Trifluoromethylguanosine TP; 2'-Deoxy-2',2'-difluoroguanosine TP; 2'-Deoxy-2'-amercaptoguanosine TP; 2'-Deoxy-2'-a-thiomethoxyguanosine TP; 2'-Deoxy-2'-b-aminoguanosine TP; 2'-Deoxy-2'-b- 35 azidoguanosine TP; 2'-Deoxy-2'-b-bromoguanosine TP; 2'-Deoxy-2'-b-chloroguanosine TP; 2'-Deoxy-2'-b-fluoroguanosine TP; 2'-Deoxy-2'-b-iodoguanosine TP; 2'-Deoxy-2'-b-mercaptoguanosine TP: 2'-Deoxy-2'-b-thiomethoxyguanosine TP; 4'-Azidoguanosine 4'-Carbocyclic guanosine TP; 4'-Ethynylguanosine TP; 5'-Homo-guanosine TP; 8-bromo-guanosine TP; 9-Deazaguanosine TP; N2-isobutyl-guanosine TP; 1-methylinosine; Inosine; 1,2'-O-dimethylinosine; 2'-O-methylinosine; 7-methylinosine; 2'-O-methylinosine; Epoxyqueuosine; 45 galactosyl-queuosine; Mannosylqueuosine; Queuosine; allyamino-thymidine; aza thymidine; deaza thymidine; deoxy-thymidine; 2'-O-methyluridine; 2-thiouridine; 3-methyluridine; 5-carboxymethyluridine; 5-hydroxyuridine; 5-methyluridine; 5-taurinomethyl-2-thiouridine; 5-tau- 50 rinomethyluridine; Dihydrouridine; Pseudouridine; (3-(3amino-3-carboxypropyl)uridine; 1-methyl-3-(3-amino-5carboxypropyl)pseudouridine; 1-methylpseduouridine; 1-methyl-pseudouridine; 2'-O-methyluridine; 2'-O-methylpseudouridine; 2'-O-methyluridine; 2-thio-2'-O-methyluri- 55 dine; 3-(3-amino-3-carboxypropyl)uridine; 3,2'-O-dimethyluridine; 3-Methyl-pseudo-Uridine TP; 4-thiouridine; 5-(carboxyhydroxymethyl)uridine; 5-(carboxyhydroxymethyl)uridine methyl ester; 5,2'-O-dimethyluridine; 5,6-dihydro-uridine; 5-aminomethyl-2-thiouridine; 5-carbamoyl- 60 methyl-2'-O-methyluridine; 5-carbamoylmethyluridine; 5-carboxyhydroxymethyluridine; 5-carboxyhydroxymethyluridine methyl ester; 5-carboxymethylaminomethyl-2'-O-5-carboxymethylaminomethyl-2-thiourimethyluridine; dine; 5-carboxymethylaminomethyl-2-thiouridine; 65 5-carboxymethylaminomethyluridine; 5-carboxymethylaminomethyluridine; 5-Carbamoylmethyluridine

5-methoxycarbonylmethyl-2'-O-methyluridine; 5-methoxycarbonylmethyl-2-thiouridine; 5-methoxycarbonylmethyluridine; 5-methoxyuridine; 5-methyl-2-thiouridine; 5-methylaminomethyl-2-selenouridine; 5-methylaminomethyl-2-5-methylaminomethyluridine; thiouridine; 5-Methyldihydrouridine; 5-Oxyacetic acid-Uridine TP; 5-Oxyacetic acid-methyl ester-Uridine TP; N1-methylpseudo-uridine; uridine 5-oxyacetic acid; uridine 5-oxyacetic acid methyl ester; 3-(3-Amino-3-carboxypropyl)-Uridine TP; 5-(iso-Pentenylaminomethyl)-2-thiouridine TP; 5-(iso-Pentenylaminomethyl)-2'-O-methyluridine 5-(iso-Pentenylaminomethyl)uridine TP; 5-propynyl uracil; α-thio-uridine; 1 (aminoalkylamino-carbonylethylenyl)-2 (thio)-pseudouracil; 1 (aminoalkylaminocarbonylethylenyl)-2,4-(dithio)pseudouracil; 1 (aminoalkylaminocarbonylethylenyl)-4 (thio)pseudouracil; (aminoalkylaminocarbonylethylenyl)-pseudouracil; 1 (aminocarbonylethylenyl)-2(thio)-pseudouracil; 1 (aminocarbonylethylenyl)-2,4-(dithio)pseudouracil; 1 (aminocarbonylethylenyl)-4 (thio)pseudouracil: (aminocarbonylethylenyl)-pseudouracil; substituted 2(thio)-pseudouracil; 1 substituted 2,4-(dithio)pseudouracil; 1 substituted 4 (thio)pseudouracil; 1 substituted pseudouracil: 1-(aminoalkylamino-carbonylethylenyl)-2-(thio)-1-Methyl-3-(3-amino-3-carboxypropyl) pseudouracil; pseudouridine TP; 1-Methyl-3-(3-amino-3-carboxypropyl) pseudo-UTP; 1-Methyl-pseudo-UTP; 2 (thio)pseudouracil; 2' deoxy uridine; 2' fluorouridine; 2-(thio)uracil; 2,4-(dithio) psuedouracil; 2' methyl, 2'amino, 2'azido, 2'fluro-guanosine; 2'-Amino-2'-deoxy-UTP; 2'-Azido-2'-deoxy-UTP; 2'-Azido-deoxyuridine TP; 2'-O-methylpseudouridine; 2' deoxy uridine; 2' fluorouridine; 2'-Deoxy-2'-a-aminouridine TP; 2'-Deoxy-2'-a-azidouridine TP; 2-methylpseudouridine; 3 (3 amino-3 carboxypropyl)uracil; 4 (thio)pseudouracil; 4-(thio)pseudouracil; 4-(thio)uracil; 4-thiouracil; 5 (1,3-diazole-1-alkyl)uracil; 5 (2-aminopropyl)uracil; 5 (aminoalkyl)uracil; 5 (dimethylaminoalkyl)uracil; 5 (guanidiniumalkyl)uracil; 5 (methoxycarbonylmethyl)-2-(thio)uracil; 5 (methoxycarbonyl-methyl)uracil; 5 (methyl) 2 (thio)uracil; 5 (methyl) 2,4 (dithio)uracil; 5 (methyl) 4 (thio)uracil; 5 (methylaminomethyl)-2 (thio)uracil; 5 (methylaminomethyl)-2,4 (dithio)uracil; 5 (methylaminomethyl)-4 (thio) uracil; 5 (propynyl)uracil; 5 (trifluoromethyl)uracil; 5-(2aminopropyl)uracil; 5-(alkyl)-2-(thio)pseudouracil; 5-(alkyl)-2,4 (dithio)pseudouracil; 5-(alkyl)-4 5-(alkyl)uracil; pseudouracil; 5-(alkyl)pseudouracil; 5-(alkynyl)uracil: 5-(allylamino)uracil: 5-(cyanoalkyl)uracil; 5-(dialkylaminoalkyl)uracil; 5-(dimethylaminoalkyl) uracil; 5-(guanidiniumalkyl)uracil; 5-(halo)uracil; 5-(1,3-diazole-1-alkyl)uracil; 5-(methoxy)uracil; 5-(methoxycarbonylmethyl)-2-(thio)uracil; 5-(methoxycarbonyl-methyl)uracil; 5-(methyl) 2(thio)uracil; 5-(methyl) 2,4 (dithio)uracil; 5-(methyl) 4 (thio)uracil; 5-(methyl)-2-(thio)pseudouracil; 5-(methyl)-2,4 (dithio)pseudouracil; 5-(methyl)-4 (thio)pseudouracil; 5-(methyl)pseudouracil; 5-(methylaminomethyl)-2 (thio)uracil; 5-(methylaminomethyl)-2,4(dithio)uracil; 5-(methylaminomethyl)-4-(thio) uracil; 5-(propynyl)uracil; 5-(trifluoromethyl)uracil; 5-aminoallyl-uridine; 5-bromo-uridine; 5-iodo-uridine; 5-uracil; 6 (azo)uracil; 6-(azo)uracil; 6-aza-uridine; allyamino-uracil; aza uracil; deaza uracil; N3 (methyl)uracil; Pseudo-UTP-1-2-ethanoic acid; Pseudouracil; 4-Thio-pseudo-UTP; 1-carboxymethyl-pseudouridine; 1-methyl-1-deaza-pseudouri-1-propynyl-uridine; 1-taurinomethyl-1-methyldine: uridine; 1-taurinomethyl-4-thio-uridine; 1-taurinomethylpseudouridine; 2-methoxy-4-thio-pseudouridine; 2-thio-1methyl-1-deaza-pseudouridine; 2-thio-1-methyl-

56

pseudouridine: 2-thio-5-aza-uridine; 2-thio-2-thio-dihydrouridine; dihydropseudouridine; 2-thiopseudouridine; 4-methoxy-2-thio-pseudouridine; 4-methoxy-pseudouridine; 4-thio-1-methyl-pseudouridine; 4-thio-pseudouridine; 5-aza-uridine; Dihydropseudouridine; 5 (±)1-(2-Hydroxypropyl)pseudouridine TP; (2R)-1-(2-Hydroxypropyl)pseudouridine TP; (2S)-1-(2-Hydroxypropyl) pseudouridine TP; (E)-5-(2-Bromo-vinyl)ara-uridine TP; (E)-5-(2-Bromo-vinyl)uridine TP; (Z)-5-(2-Bromo-vinyl) ara-uridine TP; (Z)-5-(2-Bromo-vinyl)uridine TP; 1-(2,2,2-Trifluoroethyl)-pseudo-UTP; 1-(2,2,3,3,3-Pentafluoropropyl)pseudouridine TP; 1-(2,2-Diethoxyethyl)pseudouridine TP; 1-(2,4,6-Trimethylbenzyl)pseudouridine TP; 1-(2,4,6-Trimethyl-benzyl)pseudo-UTP; 1-(2,4,6-Trimethyl-phenyl) pseudo-UTP: 1-(2-Amino-2-carboxyethyl)pseudo-UTP; 15 1-(2-Amino-ethyl)pseudo-UTP; 1-(2-Hydroxyethyl) pseudouridine TP; 1-(2-Methoxyethyl)pseudouridine TP; 1-(3,4-Bis-trifluoromethoxybenzyl)pseudouridine TP; 1-(3, 4-Dimethoxybenzyl)pseudouridine TP; 1-(3-Amino-3-carboxypropyl)pseudo-UTP; 1-(3-Amino-propyl)pseudo-UTP; 20 1-(3-Cyclopropyl-prop-2-ynyl)pseudouridine TP; Amino-4-carboxybutyl)pseudo-UTP; 1-(4-Amino-benzyl) pseudo-UTP; 1-(4-Amino-butyl)pseudo-UTP; 1-(4-Aminophenyl)pseudo-UTP; 1-(4-Azidobenzyl)pseudouridine TP; 1-(4-Bromobenzyl)pseudouridine TP; 1-(4-Chlorobenzyl) 25 pseudouridine TP; 1-(4-Fluorobenzyl)pseudouridine TP; 1-(4-Iodobenzyl)pseudouridine TP; 1-(4-Methanesulfonylbenzyl)pseudouridine TP; 1-(4-Methoxybenzyl)pseudouridine TP; 1-(4-Methoxy-benzyl)pseudo-UTP; 1-(4-Methoxyphenyl)pseudo-UTP; 1-(4-Methylbenzyl)pseudouridine TP; 30 1-(4-Nitrobenzyl) 1-(4-Methyl-benzyl)pseudo-UTP; pseudouridine TP; 1-(4-Nitro-benzyl)pseudo-UTP; 1(4-Nitro-phenyl)pseudo-UTP; 1-(4-Thiomethoxybenzyl) pseudouridine 1-(4-Trifluoromethoxybenzyl) pseudouridine TP; 1-(4-Trifluoromethylbenzyl) 35 pseudouridine TP; 1-(5-Amino-pentyl)pseudo-UTP; 1-(6-Amino-hexyl)pseudo-UTP: 1,6-Dimethyl-pseudo-UTP; 1-[3-(2-{2-[2-(2-Aminoethoxy)-ethoxy}-ethoxy}-ethoxy)propionyl]pseudouridine TP; 1-{3-[2-(2-Aminoethoxy)ethoxy]-propionyl}pseudouridine TP; 1-Acetylpseudouri- 40 dine TP; 1-Alkyl-6-(1-propynyl)-pseudo-UTP; 1-Alkyl-6-(2-propynyl)-pseudo-UTP; 1-Alkyl-6-allyl-pseudo-UTP; 1-Alkyl-6-ethynyl-pseudo-UTP; 1-Alkyl-6-homoallylpseudo-UTP; 1-Alkyl-6-vinyl-pseudo-UTP; 1-Allylpseudouridine TP; 1-Aminomethyl-pseudo-UTP; 1-Benzo- 45 ylpseudouridine TP; 1-Benzyloxymethylpseudouridine TP; 1-Benzyl-pseudo-UTP: 1-Biotinyl-PEG2-pseudouridine TP: 1-Biotinylpseudouridine TP; 1-Butyl-pseudo-UTP; 1-Cyanomethylpseudouridine TP; 1-Cyclobutylmethyl-pseudo-UTP; 1-Cyclobutyl-pseudo-UTP; 1-Cycloheptylmethyl- 50 1-Cycloheptyl-pseudo-UTP: pseudo-UTP: 1-Cyclohexylmethyl-pseudo-UTP; 1-Cyclohexyl-pseudo-1-Cyclooctylmethyl-pseudo-UTP; 1-Cyclooctylpseudo-UTP; 1-Cyclopentylmethyl-pseudo-UTP; 1-Cyclo-1-Cyclopropylmethyl-pseudo-UTP; 55 pentyl-pseudo-UTP; 1-Cyclopropyl-pseudo-UTP; 1-Ethyl-pseudo-UTP; 1-Hexyl-pseudo-UTP; 1-Homoallylpseudouridine 1-Hydroxymethylpseudouridine TP; 1-iso-propyl-pseudo-UTP; 1-Me-2-thio-pseudo-UTP; 1-Me-4-thio-pseudo-UTP; 1-Me-alpha-thio-pseudo-UTP; 1-Methanesulfonylmethylp- 60 seudouridine TP; 1-Methoxymethylpseudouridine TP; 1-Methyl-6-(2,2,2-Trifluoroethyl)pseudo-UTP; 1-Methyl-6-(4-morpholino)-pseudo-UTP; 1-Methyl-6-(4-thiomorpholino)-pseudo-UTP; 1-Methyl-6-(substituted phenyl) pseudo-UTP; 1-Methyl-6-amino-pseudo-UTP; 1-Methyl-6- 65 azido-pseudo-UTP; 1-Methyl-6-bromo-pseudo-UTP; 1-Methyl-6-butyl-pseudo-UTP; 1-Methyl-6-chloro-pseudo-

UTP; 1-Methyl-6-cyano-pseudo-UTP; 1-Methyl-6-dimeth-1-Methyl-6-ethoxy-pseudo-UTP; ylamino-pseudo-UTP; 1-Methyl-6-ethylcarboxylate-pseudo-UTP; 1-Methyl-6-1-Methyl-6-fluoro-pseudo-UTP; ethyl-pseudo-UTP; 1-Methyl-6-formyl-pseudo-UTP; 1-Methyl-6-hydroxyamino-pseudo-UTP; 1-Methyl-6-hydroxy-pseudo-UTP; 1-Methyl-6-iodo-pseudo-UTP; 1-Methyl-6-iso-propyl-pseudo-UTP; 1-Methyl-6-methoxy-pseudo-UTP; 1-Methyl-6-methylamino-pseudo-UTP; 1-Methyl-6-phenylpseudo-UTP; 1-Methyl-6-propyl-pseudo-UTP; 1-Methyl-6tert-butyl-pseudo-UTP; 1-Methyl-6-trifluoromethoxypseudo-UTP; 1-Methyl-6-trifluoromethyl-pseudo-UTP; 1-Morpholinomethylpseudouridine TP; 1-Pentyl-pseudo-UTP; 1-Phenyl-pseudo-UTP; 1-Pivaloylpseudouridine TP; 1-Propargylpseudouridine TP; 1-Propyl-pseudo-UTP; 1-propynyl-pseudouridine; 1-p-tolyl-pseudo-UTP; 1-tert-Butyl-pseudo-UTP; 1-Thiomethoxymethylpseudouridine TP; 1-Thiomorpholinomethylpseudouridine TP; 1-Trifluoroacetylpseudouridine TP; 1-Trifluoromethyl-pseudo-UTP; 1-Vinylpseudouridine TP: 2.2'-anhydro-uridine 2'-bromo-deoxyuridine TP; 2'-F-5-Methyl-2'-deoxy-UTP; 2'-OMe-5-Me-UTP; 2'-OMe-pseudo-UTP; 2'-a-Ethynyluridine TP; 2'-a-Trifluoromethyluridine TP; 2'-b-Ethynyluridine TP; 2'-b-Trifluoromethyluridine TP; 2'-Deoxy-2',2'-difluorouridine TP; 2'-Deoxy-2'-a-mercaptouridine TP; 2'-Deoxy-2'-a-thiomethoxyuridine TP; 2'-Deoxy-2'-b-aminouridine TP; 2'-Deoxy-2'-b-azidouridine TP; 2'-Deoxy-2'b-bromouridine TP; 2'-Deoxy-2'-b-chlorouridine TP; 2'-Deoxy-2'-b-fluorouridine TP; 2'-Deoxy-2'-b-iodouridine TP; TP; 2'-Deoxy-2'-b-thio-2'-Deoxy-2'-b-mercaptouridine methoxyuridine TP: 2-methoxy-4-thio-uridine; 2-methoxyuridine; 2'-O-Methyl-5-(1-propynyl)uridine TP; 3-Alkyl-pseudo-UTP; 4'-Azidouridine TP; 4'-Carbocyclic uridine TP; 4'-Ethynyluridine TP; 5-(1-Propynyl)ara-uridine TP; 5-(2-Furanyl)uridine TP; 5-Cyanouridine TP; 5-Dimethylaminouridine TP; 5'-Homo-uridine TP; 5-iodo-2'fluoro-deoxyuridine TP; 5-Phenylethynyluridine TP; 5-Trideuteromethyl-6-deuterouridine TP; 5-Trifluoromethyl-Uridine TP; 5-Vinylarauridine TP; 6-(2,2,2-Trifluoroethyl)-6-(4-Morpholino)-pseudo-UTP; pseudo-UTP; Thiomorpholino)-pseudo-UTP; 6-(Substituted-Phenyl)pseudo-UTP; 6-Amino-pseudo-UTP; 6-Azido-pseudo-UTP; 6-Bromo-pseudo-UTP; 6-Butyl-pseudo-UTP; 6-Chloropseudo-UTP; 6-Cyano-pseudo-UTP; 6-Dimethylaminopseudo-UTP; 6-Ethoxy-pseudo-UTP; 6-Ethylcarboxylatepseudo-UTP; 6-Ethyl-pseudo-UTP; 6-Fluoro-pseudo-UTP; 6-Formyl-pseudo-UTP: 6-Hydroxyamino-pseudo-UTP: 6-Hydroxy-pseudo-UTP; 6-Iodo-pseudo-UTP; 6-iso-Pro-6-Methoxy-pseudo-UTP; pyl-pseudo-UTP; 6-Methylamino-pseudo-UTP; 6-Methyl-pseudo-UTP; 6-Phenvlpseudo-UTP; 6-Phenyl-pseudo-UTP; 6-Propyl-pseudo-6-tert-Butyl-pseudo-UTP; 6-Trifluoromethoxy-UTP; pseudo-UTP; 6-Trifluoromethyl-pseudo-UTP; Alpha-thiopseudo-UTP; Pseudouridine 1-(4-methylbenzenesulfonic acid) TP; Pseudouridine 1-(4-methylbenzoic acid) TP; Pseudouridine TP 1-[3-(2-ethoxy)]propionic acid; Pseudou- $1-[3-{2-(2-(2-ethoxy)-ethoxy)-ethoxy}]$ ethoxy}|propionic acid; Pseudouridine TP 1-[3-{2-(2-[2-{2}]) (2-ethoxy)-ethoxy}-ethoxy}-ethoxy}|propionic acid; Pseudouridine TP 1-[3-{2-(2-[2-ethoxy]-ethoxy)ethoxy}]propionic acid; Pseudouridine TP 1-[3-{2-(2ethoxy)-ethoxy}] propionic acid; Pseudouridine TP 1-methylphosphonic acid; Pseudouridine TP 1-methylphosphonic acid diethyl ester; Pseudo-UTP-N1-3-propionic acid; Pseudo-UTP-N1-4-butanoic acid; Pseudo-UTP-N1-5-pentanoic acid; Pseudo-UTP-N1-6-hexanoic acid; Pseudo-UTP-N1-7-heptanoic acid; Pseudo-UTP-N1-methyl-p-ben-

zoic acid; Pseudo-UTP-N1-p-benzoic acid; Wybutosine; Hydroxywybutosine; Isowyosine; Peroxywybutosine; undermodified hydroxywybutosine; 4-demethylwyosine; 2,6-(diamino)purine; 1-(aza)-2-(thio)-3-(aza)-phenoxazin-1-yl: 1.3-(diaza)-2-(oxo)-phenthiazin-1-yl; 1.3-(diaza)-2-(oxo)-phenoxazin-1-vl; 1.3.5-(triaza)-2.6-(dioxa)-naphthalene: 2 (amino)purine: 2,4,5-(trimethyl)phenyl: 2' methyl. 2'amino, 2'azido, 2'fluro-cytidine; 2' methyl, 2'amino, 2'azido, 2'fluro-adenine; 2'methyl, 2'amino, 2'azido, 2'flurouridine; 2'-amino-2'-deoxyribose; 2-amino-6-Chloro-purine; 2-aza-inosinyl; 2'-azido-2'-deoxyribose; 2'fluoro-2'-deoxyribose; 2'-fluoro-modified bases; 2'-O-methyl-ribose; 2-oxo-7-aminopyridopyrimidin-3-yl; 2-oxo-pyridopyrimidine-3yl; 2-pyridinone; 3 nitropyrrole; 3-(methyl)-7-(propynyl) isocarbostyrilyl; 3-(methyl)isocarbostyrilyl; 4-(fluoro)-6-(methyl)benzimidazole; 4-(methyl)benzimidazole; 4-(methyl)indolyl; 4,6-(dimethyl)indolyl; 5 nitroindole; 5 substituted pyrimidines; 5-(methyl)isocarbostyrilyl; 5-nitroindole; 6-(aza)pyrimidine; 6-(azo)thymine; 6-(methyl)-7- 20 (aza)indolyl; 6-chloro-purine; 6-phenyl-pyrrolo-pyrimidin-2-on-3-yl; 7-(aminoalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)phenthiazin-1-yl; 7-(aminoalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)-phenoxazin-1-yl; 7-(aminoalkylhydroxy)-1,3-(diaza)-2-(oxo)-phenoxazin-1-yl; 7-(aminoalkylhydroxy)-1, 25 3-(diaza)-2-(oxo)-phenthiazin-1-yl; 7-(aminoalkylhydroxy)-1,3-(diaza)-2-(oxo)-phenoxazin-1yl; 7-(aza)indolyl; 7-(guanidiniumalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)-phenoxazinl-yl; 7-(guanidiniumalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)-phenthiazin-1-yl; 7-(guanidiniumalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)-phenoxazin-1-yl; 7-(guanidiniumalkylhydroxy)-1,3-(diaza)-2-

7-(guanidiniumalkylhydroxy)-1-(aza)-2-(thio)-3-(aza)-phenoxazin-1-yl; 7-(guanidiniumalkylhydroxy)-1,3-(diaza)-2-(oxo)-phenoxazin-1-yl; 7-(guanidiniumalkyl-hydroxy)-1,3-(diaza)-2-(oxo)-phenthiazin-1-yl;

7-(guanidiniumalkylhydroxy)-1,3-(diaza)-2-(oxo)-phenoxazin-1-yl; 7-(propynyl)isocarbostyrilyl; 7-(propynyl)isocarbostyrilyl, propynyl-7-(aza)indolyl; 7-deaza-inosinyl; 7-substituted 1-(aza)-2-(thio)-3-(aza)-phenoxazin-1-yl; 7-substituted 1,3-(diaza)-2-(oxo)-phenoxazin-1-yl; 9-(methyl)-imidizopyridinyl; Aminoindolyl; Anthracenyl; 40 bis-ortho-(aminoalkylhydroxy)-6-phenyl-pyrrolo-pyrimidin-2-on-3-yl; bis-ortho-substituted-6-phenyl-pyrrolo-pyrimidin-2-on-3-yl; Difluorotolyl; Hypoxanthine; Imidizopyridinyl; Inosinyl; Isocarbostyrilyl; Isoguanisine; N2-substituted N6-methyl-2-amino-purine; 45 purines; N6-substituted purines; N-alkylated derivative; Napthalenyl: Nitrobenzimidazolyl: Nitroimidazolyl: Nitroindazolyl: Nitropyrazolyl; Nubularine; 06-substituted purines; O-alkylated derivative; ortho-(aminoalkylhydroxy)-6-phenyl-pyrrolo-pyrimidin-2-on-3-yl; ortho-substituted-6-phenyl-pyr- 50 rolo-pyrimidin-2-on-3-yl; Oxoformycin TP: (aminoalkylhydroxy)-6-phenyl-pyrrolo-pyrimidin-2-on-3yl; para-substituted-6-phenyl-pyrrolo-pyrimidin-2-on-3-yl; Pentacenyl; Phenanthracenyl; Phenyl; propynyl-7-(aza)indolyl; Pyrenyl; pyridopyrimidin-3-yl; pyridopyrimidin-3-yl, 55 2-oxo-7-amino-pyridopyrimidin-3-yl; pyrrolo-pyrimidin-2on-3-yl; Pyrrolopyrimidinyl; Pyrrolopyrizinyl; Stilbenzyl; substituted 1,2,4-triazoles; Tetracenyl; Tubercidine; Xanthine; Xanthosine-5'-TP; 2-thio-zebularine; 5-aza-2-thio-zebularine; 7-deaza-2-amino-purine; pyridin-4-one ribo- 60 nucleoside; 2-Amino-riboside-TP; Formycin A Formycin B TP; Pyrrolosine TP; 2'-OH-ara-adenosine TP; 2'-OH-ara-cytidine TP; 2'-OH-ara-uridine TP; 2'-OH-araguanosine TP; 5-(2-carbomethoxyvinyl)uridine TP; and N6-(19-Amino-pentaoxanonadecyl)adenosine TP.

In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) include a com-

58

bination of at least two (e.g., 2, 3, 4 or more) of the aforementioned modified nucleobases.

In some embodiments, modified nucleobases in polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) are selected from the group consisting of pseudouridine ( $\psi$ ), N1-methylpseudouridine ( $m^1\psi$ ), N1-ethylpseudouridine, 2-thiouridine, 4'-thiouridine, 5-methylcytosine, 2-thio-1-methyl-1-deaza-pseudouridine, 2-thio-1methyl-pseudouridine, 2-thio-5-aza-uridine, 2-thiodihydropseudouridine, 2-thio-dihydrouridine, 2-thiopseudouridine, 4-methoxy-2-thio-pseudouridine, 4-methoxy-pseudouridine, 4-thio-1-methyl-pseudouridine, 4-thio-pseudouridine, 5-aza-uridine, dihydropseudouridine, 5-methoxyuridine and 2'-O-methyl uridine. In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) include a combination of at least two (e.g., 2, 3, 4 or more) of the aforementioned modified nucleobases.

In some embodiments, modified nucleobases in polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) are selected from the group consisting of 1-methyl-pseudouridine ( $m^1\psi$ ), 5-methoxy-uridine ( $m^5U$ ), 5-methyl-cytidine ( $m^5C$ ), pseudouridine ( $\psi$ ),  $\alpha$ -thio-guanosine and  $\alpha$ -thio-adenosine. In some embodiments, polynucleotides includes a combination of at least two (e.g., 2, 3, 4 or more) of the aforementioned modified nucleobases.

In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise pseudouridine ( $\psi$ ) and 5-methyl-cytidine (m<sup>5</sup>C). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 1-methylpseudouridine  $(m^1\psi)$ . In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 1-methyl-pseudouridine ( $m^1\psi$ ) and 35 5-methyl-cytidine (m<sup>5</sup>C). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 2-thiouridine (s<sup>2</sup>U). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 2-thiouridine and 5-methyl-cytidine (m<sup>5</sup>C). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise methoxy-uridine (mo<sup>5</sup>U). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 5-methoxy-uridine (mo<sup>5</sup>U) and 5-methyl-cytidine (m<sup>5</sup>C). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 2'-O-methyl uridine. In some embodiments polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise 2'-Omethyl uridine and 5-methyl-cytidine (m<sup>5</sup>C). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise N6-methyl-adenosine (m<sup>6</sup>A). In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) comprise N6-methyl-adenosine (m<sup>6</sup>A) and 5-methyl-cytidine (m<sup>5</sup>C).

In some embodiments, polynucleotides (e.g., RNA polynucleotides, such as mRNA polynucleotides) are uniformly modified (e.g., fully modified, modified throughout the entire sequence) for a particular modification. For example, a polynucleotide can be uniformly modified with 5-methylcytidine (m<sup>5</sup>C), meaning that all cytosine residues in the mRNA sequence are replaced with 5-methyl-cytidine (m<sup>5</sup>C). Similarly, a polynucleotide can be uniformly modified for any type of nucleoside residue present in the sequence by replacement with a modified residue such as those set forth above.

Exemplary nucleobases and nucleosides having a modified cytosine include N4-acetyl-cytidine (ac4C), 5-methyl-cytidine (m5C), 5-halo-cytidine (e.g., 5-iodo-cytidine), 5-hydroxymethyl-cytidine (hm5C), 1-methyl-pseudoisocytidine, 2-thio-cytidine (s2C), and 2-thio-5-methyl-cytidine.

In some embodiments, a modified nucleobase is a modified uridine. Exemplary nucleobases and In some embodiments, a modified nucleobase is a modified cytosine. nucleosides having a modified uridine include 5-cyano uridine, and 4'-thio uridine.

In some embodiments, a modified nucleobase is a modified adenine. Exemplary nucleobases and nucleosides having a modified adenine include 7-deaza-adenine, 1-methyladenosine (m1A), 2-methyladenine (m2A), and N6-methyladenosine (m6A).

In some embodiments, a modified nucleobase is a modified guanine. Exemplary nucleobases and nucleosides having a modified guanine include inosine (I), 1-methyl-inosine (m1I), wyosine (imG), methylwyosine (mimG), 7-deazaguanosine, 7-cyano-7-deaza-guanosine (preQ0), 7-aminomethyl-7-deaza-guanosine (preQ1), 7-methyl-guanosine (m7G), 1-methyl-guanosine (m1G), 8-oxo-guanosine, 7-methyl-8-oxo-guanosine.

The polynucleotides of the present disclosure may be partially or fully modified along the entire length of the 25 molecule. For example, one or more or all or a given type of nucleotide (e.g., purine or pyrimidine, or any one or more or all of A, G, U, C) may be uniformly modified in a polynucleotide of the disclosure, or in a given predetermined sequence region thereof (e.g., in the mRNA including or 30 excluding the polyA tail). In some embodiments, all nucleotides X in a polynucleotide of the present disclosure (or in a given sequence region thereof) are modified nucleotides, wherein X may any one of nucleotides A, G, U, C, or any one of the combinations A+G, A+U, A+C, G+U, G+C, U+C, 35 A+G+U, A+G+C, G+U+C or A+G+C.

The polynucleotide may contain from about 1% to about 100% modified nucleotides (either in relation to overall nucleotide content, or in relation to one or more types of nucleotide, i.e., any one or more of A, G, U or C) or any 40 intervening percentage (e.g., from 1% to 20%, from 1% to 25%, from 1% to 50%, from 1% to 60%, from 1% to 70%, from 1% to 80%, from 1% to 90%, from 1% to 95%, from 10% to 20%, from 10% to 25%, from 10% to 50%, from 10% to 60%, from 10% to 70%, from 10% to 80%, from 45 10% to 90%, from 10% to 95%, from 10% to 100%, from 20% to 25%, from 20% to 50%, from 20% to 60%, from 20% to 70%, from 20% to 80%, from 20% to 90%, from 20% to 95%, from 20% to 100%, from 50% to 60%, from 50% to 70%, from 50% to 80%, from 50% to 90%, from 50 50% to 95%, from 50% to 100%, from 70% to 80%, from 70% to 90%, from 70% to 95%, from 70% to 100%, from 80% to 90%, from 80% to 95%, from 80% to 100%, from 90% to 95%, from 90% to 100%, and from 95% to 100%). Any remaining percentage is accounted for by the presence 55 of unmodified A, G, U, or C.

The polynucleotides may contain at a minimum 1% and at maximum 100% modified nucleotides, or any intervening percentage, such as at least 5% modified nucleotides, at least 10% modified nucleotides, at least 25% modified nucleotides, at least 80% modified nucleotides, or at least 90% modified nucleotides. For example, the polynucleotides may contain a modified pyrimidine such as a modified uracil or cytosine. In some embodiments, at least 5%, at least 10%, at least 25%, at least 50%, at least 25%, at least 50%, at least 25%, at least 25%, at least 65%, at least 90% or 100% of the uracil in the polynucleotide is replaced with a modified uracil (e.g., a

60

5-substituted uracil). The modified uracil can be replaced by a compound having a single unique structure, or can be replaced by a plurality of compounds having different structures (e.g., 2, 3, 4 or more unique structures). n some embodiments, at least 5%, at least 10%, at least 25%, at least 50%, at least 80%, at least 90% or 100% of the cytosine in the polynucleotide is replaced with a modified cytosine (e.g., a 5-substituted cytosine). The modified cytosine can be replaced by a compound having a single unique structure, or can be replaced by a plurality of compounds having different structures (e.g., 2, 3, 4 or more unique structures).

Thus, in some embodiments, the RNA (e.g., mRNA) vaccines comprise a 5'UTR element, an optionally codon optimized open reading frame, and a 3'UTR element, a poly(A) sequence and/or a polyadenylation signal wherein the RNA is not chemically modified.

In some embodiments, the modified nucleobase is a modified uracil. Exemplary nucleobases and nucleosides having a modified uracil include pseudouridine ( $\psi$ ), pyridin-4-one ribonucleoside, 5-aza-uridine, 6-aza-uridine, 2-thio-5-aza-uridine, 2-thio-uridine (s<sup>2</sup>U), 4-thio-uridine (s<sup>4</sup>U), 4-thio-pseudouridine, 2-thio-pseudouridine, 5-hydroxy-uridine (ho<sup>5</sup>U), 5-aminoallyl-uridine, 5-halo-uridine (e.g., 5-iodo-uridineor 5-bromo-uridine), 3-methyl-uridine (m<sup>3</sup>U), 5-methoxy-uridine (mo<sup>5</sup>U), uridine 5-oxyacetic acid (cmo<sup>5</sup>U), uridine 5-oxyacetic acid methyl ester (mcmo<sup>5</sup>U), 5-carboxymethyl-uridine (cm<sup>5</sup>U), 1-carboxymethylpseudouridine, 5-carboxyhydroxymethyl-uridine (chm<sup>5</sup>U), 5-carboxyhydroxymethyl-uridine methyl ester (mchm<sup>5</sup>U), 5-methoxycarbonylmethyl-uridine (mcm<sup>5</sup>U), 5-methoxycarbonylmethyl-2-thio-uridine (mcm<sup>5</sup>s<sup>2</sup>U), 5-aminomethyl-2-thio-uridine (nm<sup>5</sup>s<sup>2</sup>U), 5-methylaminomethyl-uridine (mnm<sup>5</sup>U), 5-methylaminomethyl-2-thio-uridine  $(mnm^5s^2U)$ . 5-methylaminomethyl-2-seleno-uridine (mnm<sup>5</sup>se<sup>2</sup>U), 5-carbamoylmethyl-uridine (ncm<sup>5</sup>U), 5-carboxymethylaminomethyl-uridine (cmnm<sup>5</sup>U), 5-carboxymethylaminomethyl-2-thio-uridine (cmnm<sup>5</sup>s<sup>2</sup>U), 5-propynyluridine, 1-propynyl-pseudouridine, 5-taurinomethyl-uridine (τm<sup>5</sup>U), 1-taurinomethyl-pseudouridine, 5-taurinomethyl-2thio-uridine (tm<sup>5</sup>s<sup>2</sup>U), 1-taurinomethyl-4-thio-pseudouridine, 5-methyl-uridine (m<sup>5</sup>U, i.e., having the nucleobase deoxythymine), 1-methyl-pseudouridine ( $m^1\psi$ ), 5-methyl-2-thio-uridine (m<sup>5</sup>s<sup>2</sup>U), 1-methyl-4-thio-pseudouridine  $(m^1s^4\psi),$ 4-thio-1-methyl-pseudouridine, 3-methylpseudouridine  $(m^3\psi)$ , 2-thio-1-methyl-pseudouridine, 1-methyl-1-deaza-pseudouridine, 2-thio-1-methyl-1-deazapseudouridine, dihydrouridine (D), dihydropseudouridine, 5,6-dihydrouridine, 5-methyldihydrouridine (m<sup>5</sup>D), 2-thiodihydrouridine, 2-thio-dihydropseudouridine, 2-methoxyuridine, 2-methoxy-4-thio-uridine, 4-methoxy-pseudouri-4-methoxy-2-thio-pseudouridine, dine, N1-methyl-3-(3-amino-3-carboxypropyl)uridine (acp<sup>3</sup>U), 1-methyl-3-(3-amino-3-carboxypropyl)pseudouridine (acp<sup>3</sup>  $\psi$ ), 5-(isopentenylaminomethyl)uridine (inm<sup>5</sup>U), 5-(isopentenylaminomethyl)-2-thio-uridine  $(inm^5s^2U)$ , α-thio-uridine, 2'-O-methyl-uridine (Urn), 5,2'-O-dimethyluridine (m<sup>5</sup>Um), 2'-O-methyl-pseudouridine (ψm), 2-thio-2'-O-methyl-uridine (s<sup>2</sup>Um), 5-methoxycarbonylmethyl-2'-O-methyl-uridine (mcm<sup>5</sup>Um), 5-carbamoylmethyl-2'-Omethyl-uridine (ncm5Um), 5-carboxymethylaminomethyl-2'-O-methyl-uridine (cmnm<sup>5</sup>Um), 3,2'-O-dimethyl-uridine (m<sup>3</sup>Um), and 5-(isopentenylaminomethyl)-2'-O-methyl-uridine (inm<sup>5</sup>Um), 1-thio-uridine, deoxythymidine, 2'-F-arauridine. 2'-F-uridine, 2'-0H-ara-uridine, 5-(2-carbomethoxyvinyl) uridine, and 5-[3-(1-E-propenylamino)] uridine.

In some embodiments, the modified nucleobase is a modified cytosine. Exemplary nucleobases and nucleosides having a modified cytosine include 5-aza-cytidine, 6-azacytidine, pseudoisocytidine, 3-methyl-cytidine N4-acetyl-cytidine  $(ac^4C),$ 5-formylcytidine  $(f^5C), 5$ N4-methyl-cytidine (m<sup>4</sup>C), 5-methyl-cytidine  $(m^5C)$ , 5-halo-cytidine (e.g., 5-iodo-cytidine), 5-hydroxymethylcytidine (hm5C), 1-methyl-pseudoisocytidine, pyrrolo-cytidine, pyrrolo-pseudoisocytidine, 2-thio-cytidine (s<sup>2</sup>C), 2-thio-5-methyl-cytidine, 4-thio-pseudoisocytidine, 4-thio-10 1-methyl-pseudoisocytidine, 4-thio-1-methyl-1-deaza-pseudoisocytidine, 1-methyl-1-deaza-pseudoisocytidine, zebularine, 5-aza-zebularine, 5-methyl-zebularine, 5-aza-2-thiozebularine. 2-thio-zebularine. 2-methoxy-cytidine, 2-methoxy-5-methyl-cytidine, 4-methoxy-pseudoisocyti- 15 4-methoxy-1-methyl-pseudoisocytidine, (k<sub>2</sub>C), α-thio-cytidine, 2'-O-methyl-cytidine (Cm), 5,2'-Odimethylcytidine (m<sup>5</sup>Cm), N4-acetyl-2'-O-methyl-cytidine (ac<sup>4</sup>Cm), N4,2'-O-dimethylcytidine (m<sup>4</sup>Cm), 5-formyl-2'-O-methyl-cytidine (f<sup>5</sup>Cm), N4,N4,2'-O-trimethyl-cytidine 20 (m<sup>4</sup>, Cm), 1-thio-cytidine, 2'-F-ara-cytidine, 2'-F-cytidine, and 2'-0H-ara-cytidine.

In some embodiments, the modified nucleobase is a modified adenine. Exemplary nucleobases and nucleosides having a modified adenine include 2-amino-purine, 2,6- 25 diaminopurine, 2-amino-6-halo-purine (e.g., 2-amino-6chloro-purine), 6-halo-purine (e.g., 6-chloro-purine), 2-amino-6-methyl-purine, 8-azido-adenosine, 7-deaza-adenine, 7-deaza-8-aza-adenine, 7-deaza-2-amino-purine, 7-deaza-8-aza-2-amino-purine, 7-deaza-2,6-diaminopurine, 30 7-deaza-8-aza-2,6-diaminopurine, 1-methyl-adenosine (m<sup>1</sup>A), 2-methyl-adenine (m<sup>2</sup>A), N6-methyl-adenosine (m<sup>6</sup>A), 2-methylthio-N6-methyl-adenosine (ms<sup>2</sup> m<sup>6</sup>A), N6-isopentenyl-adenosine (i<sup>6</sup>A), 2-methylthio-N6-isopentenyl-adenosine (ms<sup>2</sup>i<sup>6</sup>A), N6-(cis-hydroxyisopentenyl)ad- 35 enosine (io<sup>6</sup>A), 2-methylthio-N6-(cis-hydroxyisopentenyl) adenosine (ms<sup>2</sup>io<sup>6</sup>A), N6-glycinylcarbamoyl-adenosine (g<sup>6</sup>A), N6-threonylcarbamoyl-adenosine (t<sup>6</sup>A), N6-methyl-N6-threonylcarbamoyl-adenosine (m<sup>6</sup>t<sup>6</sup>A), 2-methylthio-N6-threonylcarbamoyl-adenosine (ms<sup>2</sup>g<sup>6</sup>A), N6,N6-dim- 40 ethyl-adenosine (m<sup>6</sup><sub>2</sub>A), N6-hydroxynorvalylcarbamoyladenosine  $(hn^6A),$ 2-methylthio-N6hydroxynorvalylcarbamoyl-adenosine  $(ms^2hn^6A)$ , N6-acetyl-adenosine (ac<sup>6</sup>A), 7-methyl-adenine, 2-methylthio-adenine, 2-methoxy-adenine, α-thio-adenosine, 2'-O- 45 methyl-adenosine (Am), N6,2'-O-dimethyl-adenosine (m<sup>6</sup>Am), N6,N6,2'-O-trimethyl-adenosine (m<sup>6</sup><sub>2</sub> Am), 1,2'-O-dimethyl-adenosine (m<sup>1</sup>Am), 2'-O-ribosyladenosine (phosphate) (Ar(p)), 2-amino-N6-methyl-purine, 1-thio-adenosine, 8-azido-adenosine, 2'-F-ara-adenosine, 2'-F-ad- 50 enosine, 2'-0H-ara-adenosine, and N6-(19-amino-pentaoxanonadecyl)-adenosine.

In some embodiments, the modified nucleobase is a modified guanine. Exemplary nucleobases and nucleosides having a modified guanine include inosine (I), 1-methyl- 55 inosine (m<sup>1</sup>I), wyosine (imG), methylwyosine (mimG), 4-demethyl-wyosine (imG-14), isowyosine (imG2), wybutosine (yW), peroxywybutosine (o<sub>2</sub>yW), hydroxywybutosine (OhyW), undermodified hydroxywybutosine (OhyW\*), 7-deaza-guanosine, queuosine (Q), epoxyqueuosine (oQ), 60 galactosyl-queuosine (galQ), mannosyl-queuosine (manQ), 7-cyano-7-deaza-guanosine (pre $Q_0$ ), 7-aminomethyl-7deaza-guanosine (preQ<sub>1</sub>), archaeosine (G±), 7-deaza-8-azaguanosine, 6-thio-guanosine, 6-thio-7-deaza-guanosine, 6-thio-7-deaza-8-aza-guanosine, 7-methyl-guanosine 65 6-thio-7-methyl-guanosine, 7-methyl-inosine, 6-methoxy-guanosine, 1-methyl-guanosine  $(m^1G)$ ,

N2-methyl-guanosine (m $^2$ G), N2,N2-dimethyl-guanosine (m $^2$ <sub>2</sub>G), N2,7-dimethyl-guano sine (m $^2$ .7G), N2,N2,7-dimethyl-guanosine (m $^2$ .7G), 8-oxo-guanosine, 7-methyl-8-oxo-guanosine, 1-methyl-6-thio-guanosine, N2-methyl-6-thio-guanosine, N2-methyl-6-thio-guanosine, N2-methyl-guanosine, N2-methyl-guanosine (Gm), N2-methyl-2'-O-methyl-guanosine (m $^2$ Gm), N2,N2-dimethyl-2'-O-methyl-guanosine (m $^2$ Gm), 1-methyl-2'-O-methyl-guanosine (m $^2$ Gm), N2,7-dimethyl-2'-O-methyl-guanosine (m $^2$ I-O-methyl-inosine (Im), 1,2'-O-dimethyl-inosine (m $^1$ Im), 2'-O-ribosylguanosine (phosphate) (Gr(p)), 1-thio-guanosine, 06-methyl-guanosine, 2'-F-ara-guanosine, and 2'-F-guanosine.

62

N-Linked Glycosylation Site Mutants

N-linked glycans of viral proteins play important roles in modulating the immune response. Glycans can be important for maintaining the appropriate antigenic conformations, shielding potential neutralization epitopes, and may alter the proteolytic susceptibility of proteins. Some viruses have putative N-linked glycosylation sites. Deletion or modification of an N-linked glycosylation site may enhance the immune response. Thus, the present disclosure provides, in some embodiments, RNA (e.g., mRNA) vaccines comprising nucleic acids (e.g., mRNA) encoding antigenic polypeptides that comprise a deletion or modification at one or more N-linked glycosylation sites.

In Vitro Transcription of RNA (e.g., mRNA)

Respiratory virus vaccines of the present disclosure comprise at least one RNA polynucleotide, such as a mRNA (e.g., modified mRNA). mRNA, for example, is transcribed in vitro from template DNA, referred to as an "in vitro transcription template." In some embodiments, an in vitro transcription template encodes a 5' untranslated (UTR) region, contains an open reading frame, and encodes a 3' UTR and a polyA tail. The particular nucleic acid sequence composition and length of an in vitro transcription template will depend on the mRNA encoded by the template.

A "5' untranslated region" (5'UTR) refers to a region of an mRNA that is directly upstream (i.e., 5') from the start codon (i.e., the first codon of an mRNA transcript translated by a ribosome) that does not encode a polypeptide.

A "3' untranslated region" (3'UTR) refers to a region of an mRNA that is directly downstream (i.e., 3') from the stop codon (i.e., the codon of an mRNA transcript that signals a termination of translation) that does not encode a polypeptide.

An "open reading frame" is a continuous stretch of DNA beginning with a start codon (e.g., methionine (ATG)), and ending with a stop codon (e.g., TAA, TAG or TGA) and encodes a polypeptide.

A "polyA tail" is a region of mRNA that is downstream, e.g., directly downstream (i.e., 3'), from the 3' UTR that contains multiple, consecutive adenosine monophosphates. A polyA tail may contain 10 to 300 adenosine monophosphates. For example, a polyA tail may contain 10, 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 160, 170, 180, 190, 200, 210, 220, 230, 240, 250, 260, 270, 280, 290 or 300 adenosine monophosphates. In some embodiments, a polyA tail contains 50 to 250 adenosine monophosphates. In a relevant biological setting (e.g., in cells, in vivo) the poly(A) tail functions to protect mRNA from enzymatic degradation, e.g., in the cytoplasm, and aids in transcription termination, export of the mRNA from the nucleus and translation.

In some embodiments, a polynucleotide includes 200 to 3,000 nucleotides. For example, a polynucleotide may include 200 to 500, 200 to 1000, 200 to 1500, 200 to 3000,

500 to 1000, 500 to 1500, 500 to 2000, 500 to 3000, 1000 to 1500, 1000 to 2000, 1000 to 3000, 1500 to 3000, or 2000 to 3000 nucleotides.

Flagellin Adjuvants

Flagellin is an approximately 500 amino acid monomeric 5 protein that polymerizes to form the flagella associated with bacterial motion. Flagellin is expressed by a variety of flagellated bacteria (*Salmonella typhimurium* for example) as well as non-flagellated bacteria (such as *Escherichia coli*). Sensing of flagellin by cells of the innate immune system 10 (dendritic cells, macrophages, etc.) is mediated by the Toll-like receptor 5 (TLR5) as well as by Nod-like receptors (NLRs) Ipaf and Naip5. TLRs and NLRs have been identified as playing a role in the activation of innate immune response and adaptive immune response. As such, flagellin 15 provides an adjuvant effect in a vaccine.

The nucleotide and amino acid sequences encoding known flagellin polypeptides are publicly available in the NCBI GenBank database. The flagellin sequences from S. Typhimurium, H. Pylori, V. Cholera, S. marcesens, S. 20 flexneri, T. Pallidum, L. pneumophila, B. burgdorferei, C. difficile, R. meliloti, A. tumefaciens, R. lupini, B. clarridgeiae, P. Mirabilis, B. subtilus, L. monocytogenes, P. aeruginosa, and E. coli, among others are known.

A flagellin polypeptide, as used herein, refers to a full 25 length flagellin protein, immunogenic fragments thereof, and peptides having at least 50% sequence identify to a flagellin protein or immunogenic fragments thereof. Exemplary flagellin proteins include flagellin from *Salmonella typhi* (UniPro Entry number: Q56086), *Salmonella typhimu-* 30 rium (A0A0C9DG09), *Salmonella enteritidis* (A0A0C9BAB7), and *Salmonella choleraesuis* (Q6V2X8), and SEQ ID NO: 54-56 (Table 17). In some embodiments, the flagellin polypeptide has at least 60%, 70%, 75%, 80%, 90%, 95%, 97%, 98%, or 99% sequence identify to a 35 flagellin protein or immunogenic fragments thereof.

In some embodiments, the flagellin polypeptide is an immunogenic fragment. An immunogenic fragment is a portion of a flagellin protein that provokes an immune response. In some embodiments, the immune response is a 40 TLR5 immune response. An example of an immunogenic fragment is a flagellin protein in which all or a portion of a hinge region has been deleted or replaced with other amino acids. For example, an antigenic polypeptide may be inserted in the hinge region. Hinge regions are the hyper- 45 variable regions of a flagellin. Hinge regions of a flagellin are also referred to as "D3 domain or region, "propeller domain or region," "hypervariable domain or region" and "variable domain or region." "At least a portion of a hinge region," as used herein, refers to any part of the hinge region 50 of the flagellin, or the entirety of the hinge region. In other embodiments an immunogenic fragment of flagellin is a 20, 25, 30, 35, or 40 amino acid C-terminal fragment of flagel-

The flagellin monomer is formed by domains D0 through 55 D3. D0 and D1, which form the stem, are composed of tandem long alpha helices and are highly conserved among different bacteria. The D1 domain includes several stretches of amino acids that are useful for TLR5 activation. The entire D1 domain or one or more of the active regions within 60 the domain are immunogenic fragments of flagellin. Examples of immunogenic regions within the D1 domain include residues 88-114 and residues 411-431 (in *Salmonella typhimurium* FliC flagellin. Within the 13 amino acids in the 88-100 region, at least 6 substitutions are permitted 65 between *Salmonella* flagellin and other flagellins that still preserve TLR5 activation. Thus, immunogenic fragments of

64

flagellin include flagellin like sequences that activate TLR5 and contain a 13 amino acid motif that is 53% or more identical to the *Salmonella* sequence in 88-100 of FliC (LQRVRELAVQSAN; SEQ ID NO: 84).

In some embodiments, the RNA (e.g., mRNA) vaccine includes an RNA that encodes a fusion protein of flagellin and one or more antigenic polypeptides. A "fusion protein" as used herein, refers to a linking of two components of the construct. In some embodiments, a carboxy-terminus of the antigenic polypeptide is fused or linked to an amino terminus of the flagellin polypeptide. In other embodiments, an amino-terminus of the antigenic polypeptide is fused or linked to a carboxy-terminus of the flagellin polypeptide. The fusion protein may include, for example, one, two, three, four, five, six or more flagellin polypeptides linked to one, two, three, four, five, six or more antigenic polypeptides. When two or more flagellin polypeptides and/or two or more antigenic polypeptides are linked such a construct may be referred to as a "multimer."

Each of the components of a fusion protein may be directly linked to one another or they may be connected through a linker. For instance, the linker may be an amino acid linker. The amino acid linker encoded for by the RNA (e.g., mRNA) vaccine to link the components of the fusion protein may include, for instance, at least one member selected from the group consisting of a lysine residue, a glutamic acid residue, a serine residue and an arginine residue. In some embodiments the linker is 1-30, 1-25, 1-25, 5-10, 5, 15, or 5-20 amino acids in length.

In other embodiments the RNA (e.g., mRNA) vaccine includes at least two separate RNA polynucleotides, one encoding one or more antigenic polypeptides and the other encoding the flagellin polypeptide. The at least two RNA polynucleotides may be co-formulated in a carrier such as a lipid nanoparticle.

Broad Spectrum RNA (e.g., mRNA) Vaccines

There may be situations where persons are at risk for infection with more than one strain of hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1). RNA (e.g., mRNA) therapeutic vaccines are particularly amenable to combination vaccination approaches due to a number of factors including, but not limited to, speed of manufacture, ability to rapidly tailor vaccines to accommodate perceived geographical threat, and the like. Moreover, because the vaccines utilize the human body to produce the antigenic protein, the vaccines are amenable to the production of larger, more complex antigenic proteins, allowing for proper folding, surface expression, antigen presentation, etc. in the human subject. To protect against more than one strain of hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1), a combination vaccine can be administered that includes RNA (e.g., mRNA) encoding at least one antigenic polypeptide protein (or antigenic portion thereof) of a first respiratory virus and further includes RNA encoding at least one antigenic polypeptide protein (or antigenic portion thereof) of a second respiratory virus. RNA (e.g., mRNA) can be co-formulated, for example, in a single lipid nanoparticle (LNP) or can be formulated in separate LNPs for co-administration.

Methods of Treatment

Provided herein are compositions (e.g., pharmaceutical compositions), methods, kits and reagents for prevention and/or treatment of respiratory diseases/infections in

humans and other mammals. Respiratory virus RNA (e.g. mRNA) vaccines can be used as therapeutic or prophylactic agents, alone or in combination with other vaccine(s). They may be used in medicine to prevent and/or treat respiratory disease/infection. In exemplary aspects, the RNA (e.g., 5 mRNA) vaccines of the present disclosure are used to provide prophylactic protection from hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1). Prophylactic protection 10 from hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) can be achieved following administration of a RNA (e.g., mRNA) vaccine of the present disclosure. Respiratory virus 15 RNA (e.g., mRNA) vaccines of the present disclosure may be used to treat or prevent viral "co-infections" containing two or more respiratory infections. Vaccines can be administered once, twice, three times, four times or more, but it is likely sufficient to administer the vaccine once (optionally 20 followed by a single booster). It is possible, although less desirable, to administer the vaccine to an infected individual to achieve a therapeutic response. Dosing may need to be adjusted accordingly.

A method of eliciting an immune response in a subject 25 against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) is provided in aspects of the present disclosure. The method involves administering to the subject a respiratory virus 30 RNA (e.g., mRNA) vaccine comprising at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH 35 and/or HCoV-HKU1) antigenic polypeptide thereof, thereby inducing in the subject an immune response specific to hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) 40 antigenic polypeptide or an immunogenic fragment thereof, wherein anti-antigenic polypeptide antibody titer in the subject is increased following vaccination relative to antiantigenic polypeptide antibody titer in a subject vaccinated with a prophylactically effective dose of a traditional vac- 45 cine against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1). An "anti-antigenic polypeptide antibody" is a serum antibody the binds specifically to the antigenic polypeptide. 50

In some embodiments, a RNA (e.g., mRNA) vaccine (e.g., a hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1 RNA vaccine) capable of eliciting an immune response is 55 administered intramuscularly via a composition including a compound according to Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe) (e.g., Compound 3, 18, 20, 25, 26, 29, 30, 60, 108-112, or 122).

A prophylactically effective dose is a therapeutically 60 effective dose that prevents infection with the virus at a clinically acceptable level. In some embodiments the therapeutically effective dose is a dose listed in a package insert for the vaccine. A traditional vaccine, as used herein, refers to a vaccine other than the RNA (e.g., mRNA) vaccines of 65 the present disclosure. For instance, a traditional vaccine includes but is not limited to live/attenuated microorganism

66

vaccines, killed/inactivated microorganism vaccines, subunit vaccines, protein antigen vaccines, DNA vaccines, VLP vaccines, etc. In exemplary embodiments, a traditional vaccine is a vaccine that has achieved regulatory approval and/or is registered by a national drug regulatory body, for example the Food and Drug Administration (FDA) in the United States or the European Medicines Agency (EMA).

In some embodiments the anti-antigenic polypeptide anti-body titer in the subject is increased 1 log to 10 log following vaccination relative to anti-antigenic polypeptide antibody titer in a subject vaccinated with a prophylactically effective dose of a traditional vaccine against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1).

In some embodiments the anti-antigenic polypeptide anti-body titer in the subject is increased 1 log, 2 log, 3 log, 5 log or 10 log following vaccination relative to anti-antigenic polypeptide antibody titer in a subject vaccinated with a prophylactically effective dose of a traditional vaccine against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1).

A method of eliciting an immune response in a subject against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) is provided in other aspects of the disclosure. The method involves administering to the subject a respiratory virus RNA (e.g., mRNA) vaccine comprising at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) antigenic polypeptide or an immunogenic fragment thereof, thereby inducing in the subject an immune response specific to hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) antigenic polypeptide or an immunogenic fragment thereof, wherein the immune response in the subject is equivalent to an immune response in a subject vaccinated with a traditional vaccine against the hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) at 2 times to 100 times the dosage level relative to the RNA (e.g., mRNA) vaccine.

In some embodiments, the immune response in the subject is equivalent to an immune response in a subject vaccinated with a traditional vaccine at 2, 3, 4, 5, 10, 50, 100 times the dosage level relative to the hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) RNA (e.g., mRNA) vaccine.

In some embodiments the immune response in the subject is equivalent to an immune response in a subject vaccinated with a traditional vaccine at 10-100 times, or 100-1000 times, the dosage level relative to the hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) RNA (e.g., mRNA) vaccine.

In some embodiments the immune response is assessed by determining [protein] antibody titer in the subject.

Some aspects of the present disclosure provide a method of eliciting an immune response in a subject against a In some embodiments the immune response in the subject is equivalent to an immune response in a subject vaccinated with a traditional vaccine at 2, 3, 4, 5, 10, 50, 100 times the dosage level relative to the hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) RNA (e.g., mRNA) vaccine by administering to the subject a respiratory virus RNA (e.g., mRNA) vaccine comprising at least one RNA (e.g., mRNA) polynucleotide having an open reading frame encoding at least one hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, 15 HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) antigenic polypeptide, thereby inducing in the subject an immune response specific to the antigenic polypeptide or an immunogenic fragment thereof, wherein the immune response in the subject is induced 2 days to 10 weeks earlier 20 relative to an immune response induced in a subject vaccinated with a prophylactically effective dose of a traditional vaccine against the hMPV, PIV3, RSV, MeV and/or Beta-CoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or 25 HCoV-HKU1). In some embodiments, the immune response in the subject is induced in a subject vaccinated with a prophylactically effective dose of a traditional vaccine at 2 times to 100 times the dosage level relative to the RNA (e.g.,

In some embodiments, the immune response in the subject is induced 2 days earlier, or 3 days earlier, relative to an immune response induced in a subject vaccinated with a prophylactically effective dose of a traditional vaccine.

In some embodiments the immune response in the subject 35 is induced 1 week, 2 weeks, 3 weeks, 5 weeks, or 10 weeks earlier relative to an immune response induced in a subject vaccinated with a prophylactically effective dose of a traditional vaccine.

Also provided herein is a method of eliciting an immune 40 response in a subject against hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1) by administering to the subject a respiratory virus RNA (e.g., mRNA) vaccine having an open 45 reading frame encoding a first antigenic polypeptide, wherein the RNA polynucleotide does not include a stabilization element, and wherein an adjuvant is not co-formulated or co-administered with the vaccine.

Therapeutic and Prophylactic Compositions

Provided herein are compositions (e.g., pharmaceutical compositions), methods, kits and reagents for prevention, treatment or diagnosis of hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH 55 and/or HCoV-HKU1) in humans and other mammals, for example. Respiratory virus RNA (e.g. mRNA) vaccines can be used as therapeutic or prophylactic agents. They may be used in medicine to prevent and/or treat infectious disease. In some embodiments, the respiratory RNA (e.g., mRNA) ovaccines of the present disclosure are used fin the priming of immune effector cells, for example, to activate peripheral blood mononuclear cells (PBMCs) ex vivo, which are then infused (re-infused) into a subject.

In some embodiments, respiratory virus vaccine containing RNA (e.g., mRNA) polynucleotides as described herein can be administered to a subject (e.g., a mammalian subject,

such as a human subject), and the RNA (e.g., mRNA) polynucleotides are translated in vivo to produce an antigenic polypeptide.

The respiratory virus RNA (e.g., mRNA) vaccines may be induced for translation of a polypeptide (e.g., antigen or immunogen) in a cell, tissue or organism. In some embodiments, such translation occurs in vivo, although such translation may occur ex vivo, in culture or in vitro. In some embodiments, the cell, tissue or organism is contacted with an effective amount of a composition containing a respiratory virus RNA (e.g., mRNA) vaccine that contains a polynucleotide that has at least one a translatable region encoding an antigenic polypeptide.

An "effective amount" of an respiratory virus RNA (e.g. mRNA) vaccine is provided based, at least in part, on the target tissue, target cell type, means of administration, physical characteristics of the polynucleotide (e.g., size, and extent of modified nucleosides) and other components of the vaccine, and other determinants. In general, an effective amount of the respiratory virus RNA (e.g., mRNA) vaccine composition provides an induced or boosted immune response as a function of antigen production in the cell, preferably more efficient than a composition containing a corresponding unmodified polynucleotide encoding the same antigen or a peptide antigen. Increased antigen production may be demonstrated by increased cell transfection (the percentage of cells transfected with the RNA, e.g., mRNA, vaccine), increased protein translation from the polynucleotide, decreased nucleic acid degradation (as demonstrated, for example, by increased duration of protein translation from a modified polynucleotide), or altered antigen specific immune response of the host cell.

In some embodiments, RNA (e.g. mRNA) vaccines (including polynucleotides their encoded polypeptides) in accordance with the present disclosure may be used for treatment of hMPV, PIV3, RSV, MeV and/or BetaCoV (including MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH and/or HCoV-HKU1).

Respiratory RNA (e.g. mRNA) vaccines may be administered prophylactically or therapeutically as part of an active immunization scheme to healthy individuals or early in infection during the incubation phase or during active infection after onset of symptoms. In some embodiments, the amount of RNA (e.g., mRNA) vaccine of the present disclosure provided to a cell, a tissue or a subject may be an amount effective for immune prophylaxis.

Respiratory virus RNA (e.g. mRNA) vaccines may be administrated with other prophylactic or therapeutic com-50 pounds. As a non-limiting example, a prophylactic or therapeutic compound may be an adjuvant or a booster. As used herein, when referring to a prophylactic composition, such as a vaccine, the term "booster" refers to an extra administration of the prophylactic (vaccine) composition. A booster (or booster vaccine) may be given after an earlier administration of the prophylactic composition. The time of administration between the initial administration of the prophylactic composition and the booster may be, but is not limited to, 1 minute, 2 minutes, 3 minutes, 4 minutes, 5 minutes, 6 minutes, 7 minutes, 8 minutes, 9 minutes, 10 minutes, 15 minutes, 20 minutes 35 minutes, 40 minutes, 45 minutes, 50 minutes, 55 minutes, 1 hour, 2 hours, 3 hours, 4 hours, 5 hours, 6 hours, 7 hours, 8 hours, 9 hours, 10 hours, 11 hours, 12 hours, 13 hours, 14 hours, 15 hours, 16 hours, 17 hours, 18 hours, 19 hours, 20 hours, 21 hours, 22 hours, 23 hours, 1 day, 36 hours, 2 days, 3 days, 4 days, 5 days, 6 days, 1 week, 10 days, 2 weeks, 3 weeks, 1 month, 2 months, 3

months, 4 months, 5 months, 6 months, 7 months, 8 months, 9 months, 10 months, 11 months, 1 year, 18 months, 2 years, 3 years, 4 years, 5 years, 6 years, 7 years, 8 years, 9 years, 10 years, 11 years, 12 years, 13 years, 14 years, 15 years, 16 years, 17 years, 18 years, 19 years, 20 years, 25 years, 30 years, 35 years, 40 years, 45 years, 50 years, 55 years, 60 years, 65 years, 70 years, 75 years, 80 years, 85 years, 90 years, 95 years or more than 99 years. In some embodiments, the time of administration between the initial administration of the prophylactic composition and the booster may be, but is not limited to, 1 week, 2 weeks, 3 weeks, 1 month, 2 months, 3 months, 6 months or 1 year.

In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines may be administered intramuscularly or intradermally, similarly to the administration of inactivated vaccines known in the art.

Respiratory virus RNA (e.g. mRNA) vaccines may be utilized in various settings depending on the prevalence of the infection or the degree or level of unmet medical need. 20 As a non-limiting example, the RNA (e.g., mRNA) vaccines may be utilized to treat and/or prevent a variety of respiratory infections. RNA (e.g., mRNA) vaccines have superior properties in that they produce much larger antibody titers and produce responses early than commercially available 25 anti-viral agents/compositions.

Provided herein are pharmaceutical compositions including respiratory virus RNA (e.g. mRNA) vaccines and RNA (e.g. mRNA) vaccine compositions and/or complexes optionally in combination with one or more pharmaceuti- 30 cally acceptable excipients.

Respiratory virus RNA (e.g. mRNA) vaccines may be formulated or administered alone or in conjunction with one or more other components. For instance, hMPV/PIV3/RSV RNA (e.g., mRNA) vaccines (vaccine compositions) may 35 comprise other components including, but not limited to, adjuvants.

In some embodiments, respiratory virus (e.g. mRNA) vaccines do not include an adjuvant (they are adjuvant free).

Respiratory virus RNA (e.g. mRNA) vaccines may be 40 formulated or administered in combination with one or more pharmaceutically-acceptable excipients. In some embodiments, vaccine compositions comprise at least one additional active substances, such as, for example, a therapeutically-active substance, a prophylactically-active substance, 45 or a combination of both. Vaccine compositions may be sterile, pyrogen-free or both sterile and pyrogen-free. General considerations in the formulation and/or manufacture of pharmaceutical agents, such as vaccine compositions, may be found, for example, in Remington: The Science and 50 Practice of Pharmacy 21st ed., Lippincott Williams & Wilkins, 2005 (incorporated herein by reference in its entirety).

In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines are administered to humans, human 55 patients or subjects. For the purposes of the present disclosure, the phrase "active ingredient" generally refers to the RNA (e.g., mRNA) vaccines or the polynucleotides contained therein, for example, RNA polynucleotides (e.g., mRNA polynucleotides) encoding antigenic polypeptides. 60

Formulations of the respiratory virus vaccine compositions described herein may be prepared by any method known or hereafter developed in the art of pharmacology. In general, such preparatory methods include the step of bringing the active ingredient (e.g., mRNA polynucleotide) into 65 association with an excipient and/or one or more other accessory ingredients, and then, if necessary and/or desir-

70

able, dividing, shaping and/or packaging the product into a desired single- or multi-dose unit.

Relative amounts of the active ingredient, the pharmaceutically acceptable excipient, and/or any additional ingredients in a pharmaceutical composition in accordance with the disclosure will vary, depending upon the identity, size, and/or condition of the subject treated and further depending upon the route by which the composition is to be administered. By way of example, the composition may comprise between 0.1% and 100%, e.g., between 0.5 and 50%, between 1-30%, between 5-80%, at least 80% (w/w) active ingredient.

Respiratory virus RNA (e.g. mRNA) vaccines can be formulated using one or more excipients to: (1) increase stability; (2) increase cell transfection; (3) permit the sustained or delayed release (e.g., from a depot formulation); (4) alter the biodistribution (e.g., target to specific tissues or cell types); (5) increase the translation of encoded protein in vivo; and/or (6) alter the release profile of encoded protein (antigen) in vivo. In addition to traditional excipients such as any and all solvents, dispersion media, diluents, or other liquid vehicles, dispersion or suspension aids, surface active agents, isotonic agents, thickening or emulsifying agents, preservatives, excipients can include, without limitation, lipidoids, liposomes, lipid nanoparticles, polymers, lipoplexes, core-shell nanoparticles, peptides, proteins, cells transfected with respiratory virus RNA (e.g. mRNA) vaccines (e.g., for transplantation into a subject), hyaluronidase, nanoparticle mimics and combinations thereof.

Stabilizing Elements

Naturally-occurring eukaryotic mRNA molecules have been found to contain stabilizing elements, including, but not limited to untranslated regions (UTR) at their 5'-end (5'UTR) and/or at their 3'-end (3'UTR), in addition to other structural features, such as a 5'-cap structure or a 3'-poly(A) tail. Both the 5'UTR and the 3'UTR are typically transcribed from the genomic DNA and are elements of the premature mRNA. Characteristic structural features of mature mRNA, such as the 5'-cap and the 3'-poly(A) tail are usually added to the transcribed (premature) mRNA during mRNA processing. The 3'-poly(A) tail is typically a stretch of adenine nucleotides added to the 3'-end of the transcribed mRNA. It can comprise up to about 400 adenine nucleotides. In some embodiments the length of the 3'-poly(A) tail may be an essential element with respect to the stability of the individual mRNA.

In some embodiments the RNA (e.g., mRNA) vaccine may include one or more stabilizing elements. Stabilizing elements may include for instance a histone stem-loop. A stem-loop binding protein (SLBP), a 32 kDa protein has been identified. It is associated with the histone stem-loop at the 3'-end of the histone messages in both the nucleus and the cytoplasm. Its expression level is regulated by the cell cycle; it peaks during the S-phase, when histone mRNA levels are also elevated. The protein has been shown to be essential for efficient 3'-end processing of histone premRNA by the U7 snRNP. SLBP continues to be associated with the stem-loop after processing, and then stimulates the translation of mature histone mRNAs into histone proteins in the cytoplasm. The RNA binding domain of SLBP is conserved through metazoa and protozoa; its binding to the histone stem-loop depends on the structure of the loop. The minimum binding site includes at least three nucleotides 5' and two nucleotides 3' relative to the stem-loop.

In some embodiments, the RNA (e.g., mRNA) vaccines include a coding region, at least one histone stem-loop, and optionally, a poly(A) sequence or polyadenylation signal.

The poly(A) sequence or polyadenylation signal generally should enhance the expression level of the encoded protein. The encoded protein, in some embodiments, is not a histone protein, a reporter protein (e.g. Luciferase, GFP, EGFP, β-Galactosidase, EGFP), or a marker or selection protein (e.g. alpha-Globin, Galactokinase and Xanthine:guanine phosphoribosyl transferase (GPT)).

In some embodiments, the combination of a poly(A) sequence or polyadenylation signal and at least one histone stem-loop, even though both represent alternative mechanisms in nature, acts synergistically to increase the protein expression beyond the level observed with either of the individual elements. It has been found that the synergistic effect of the combination of poly(A) and at least one histone stem-loop does not depend on the order of the elements or 15 the length of the poly(A) sequence.

In some embodiments, the RNA (e.g., mRNA) vaccine does not comprise a histone downstream element (HDE). "Histone downstream element" (HDE) includes a purinerich polynucleotide stretch of approximately 15 to 20 20 nucleotides 3' of naturally occurring stem-loops, representing the binding site for the U7 snRNA, which is involved in processing of histone pre-mRNA into mature histone mRNA. Ideally, the inventive nucleic acid does not include an intron

In some embodiments, the RNA (e.g., mRNA) vaccine may or may not contain a enhancer and/or promoter sequence, which may be modified or unmodified or which may be activated or inactivated. In some embodiments, the histone stem-loop is generally derived from histone genes, 30 and includes an intramolecular base pairing of two neighbored partially or entirely reverse complementary sequences separated by a spacer, including (e.g., consisting of) a short sequence, which forms the loop of the structure. The unpaired loop region is typically unable to base pair with 35 either of the stem loop elements. It occurs more often in RNA, as is a key component of many RNA secondary structures, but may be present in single-stranded DNA as well. Stability of the stem-loop structure generally depends on the length, number of mismatches or bulges, and base 40 composition of the paired region. In some embodiments, wobble base pairing (non-Watson-Crick base pairing) may result. In some embodiments, the at least one histone stemloop sequence comprises a length of 15 to 45 nucleotides.

In other embodiments the RNA (e.g., mRNA) vaccine 45 may have one or more AU-rich sequences removed. These sequences, sometimes referred to as AURES are destabilizing sequences found in the 3'UTR. The AURES may be removed from the RNA (e.g., mRNA) vaccines. Alternatively the AURES may remain in the RNA (e.g., mRNA) 50 vaccine.

## Nanoparticle Formulations

In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines are formulated in a nanoparticle. In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines 55 are formulated in a lipid nanoparticle. In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines are formulated in a lipid-polycation complex, referred to as a cationic lipid nanoparticle. As a non-limiting example, the polycation may include a cationic peptide or a polypeptide 60 such as, but not limited to, polylysine, polyornithine and/or polyarginine. In some embodiments, respiratory virus RNA (e.g., mRNA) vaccines are formulated in a lipid nanoparticle that includes a non-cationic lipid such as, but not limited to, cholesterol or dioleoyl phosphatidylethanolamine (DOPE). 65

A lipid nanoparticle formulation may be influenced by, but not limited to, the selection of the cationic lipid com72

ponent, the degree of cationic lipid saturation, the nature of the PEGylation, ratio of all components and biophysical parameters such as size. In one example by Semple et al. (*Nature Biotech.* 2010 28:172-176), the lipid nanoparticle formulation is composed of 57.1% cationic lipid, 7.1% dipalmitoylphosphatidylcholine, 34.3% cholesterol, and 1.4% PEG-c-DMA. As another example, changing the composition of the cationic lipid can more effectively deliver siRNA to various antigen presenting cells (Basha et al. *Mol Ther.* 2011 19:2186-2200).

In some embodiments, lipid nanoparticle formulations may comprise 35 to 45% cationic lipid, 40% to 50% cationic lipid, 50% to 60% cationic lipid and/or 55% to 65% cationic lipid. In some embodiments, the ratio of lipid to RNA (e.g., mRNA) in lipid nanoparticles may be 5:1 to 20:1, 10:1 to 25:1, 15:1 to 30:1 and/or at least 30:1.

In some embodiments, the ratio of PEG in the lipid nanoparticle formulations may be increased or decreased and/or the carbon chain length of the PEG lipid may be modified from C14 to C18 to alter the pharmacokinetics and/or biodistribution of the lipid nanoparticle formulations. As a non-limiting example, lipid nanoparticle formulations may contain 0.5% to 3.0%, 1.0% to 3.5%, 1.5% to 4.0%, 2.0% to 4.5%, 2.5% to 5.0% and/or 3.0% to 6.0% of the lipid molar ratio of PEG-c-DOMG (R-3-[(ω-methoxy-poly(ethyleneglycol)2000)carbamoyl)]-1,2-dimyristyloxypropyl-3amine) (also referred to herein as PEG-DOMG) as compared to the cationic lipid, DSPC and cholesterol. In some embodiments, the PEG-c-DOMG may be replaced with a PEG lipid such as, but not limited to, PEG-DSG (1,2-Distearoyl-snglycerol, methoxypolyethylene glycol), PEG-DMG (1,2-Dimyristoyl-sn-glycerol) and/or PEG-DPG (1,2-Dipalmitoyl-sn-glycerol, methoxypolyethylene glycol). The cationic lipid may be selected from any lipid known in the art such as, but not limited to, DLin-MC3-DMA, DLin-DMA, C12-200 and DLin-KC2-DMA.

In some embodiments, an respiratory virus RNA (e.g. mRNA) vaccine formulation is a nanoparticle that comprises at least one lipid. The lipid may be selected from, but is not limited to, DLin-DMA, DLin-K-DMA, 98N12-5, C12-200, DLin-MC3-DMA, DLin-KC2-DMA, DODMA, PLGA, PEG, PEG-DMG, PEGylated lipids and amino alcohol lipids. In some embodiments, the lipid may be a cationic lipid such as, but not limited to, DLin-DMA, DLin-D-DMA, DLin-MC3-DMA, DLin-KC2-DMA, DODMA and amino alcohol lipids. The amino alcohol cationic lipid may be the lipids described in and/or made by the methods described in U.S. Patent Publication No. US20130150625, herein incorporated by reference in its entirety. As a non-limiting example, the cationic lipid may be 2-amino-3-[(9Z,12Z)octadeca-9,12-dien-1-yloxy]-2-{[(9Z,2Z)-octadeca-9,12dien-1-yloxy]methyl}propan-1-ol (Compound US20130150625); 2-amino-3-[(9Z)-octadec-9-en-1-yloxy]-2-{[(9Z)-octadec-9-en-1-yloxy]methyl}propan-1-ol (Compound 2 in US20130150625); 2-amino-3-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-2-[(octyloxy)methyl]propan-1-ol (Compound 3 in US20130150625); and 2-(dimethylamino)- $3-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-2-\{[(9Z,12Z)-oc-4]((9Z,12Z)-oc-4)\}$ tadeca-9,12-dien-1-yloxy|methyl|propan-1-ol (Compound 4 in US20130150625); or any pharmaceutically acceptable salt or stereoisomer thereof.

Lipid nanoparticle formulations typically comprise a lipid, in particular, an ionizable cationic lipid, for example, 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), or di((Z)-non-2-en-1-yl) 9-((4-

(dimethylamino)butanoyl)oxy)heptadecanedioate (L319),

and further comprise a neutral lipid, a sterol and a molecule capable of reducing particle aggregation, for example a PEG or PEG-modified lipid.

73

In some embodiments, a lipid nanoparticle formulation consists essentially of (i) at least one lipid selected from the 5 group consisting of 2,2-dilinoleyl-4-dimethylaminoethyl-[1, 3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy) heptadecanedioate (L319); (ii) a neutral lipid selected from 10 DSPC, DPPC, POPC, DOPE and SM; (iii) a sterol, e.g., cholesterol; and (iv) a PEG-lipid, e.g., PEG-DMG or PEG-cDMA, in a molar ratio of 20-60% cationic lipid:5-25% neutral lipid:25-55% sterol; 0.5-15% PEG-lipid.

In some embodiments, a lipid nanoparticle formulation 15 includes 25% to 75% on a molar basis of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate 20 (L319), e.g., 35 to 65%, 45 to 65%, 60%, 57.5%, 50% or 40% on a molar basis.

In some embodiments, a lipid nanoparticle formulation includes 0.5% to 15% on a molar basis of the neutral lipid, e.g., 3 to 12%, 5 to 10% or 15%, 10%, or 7.5% on a molar 25 basis. Examples of neutral lipids include, without limitation, DSPC, POPC, DPPC, DOPE and SM. In some embodiments, the formulation includes 5% to 50% on a molar basis of the sterol (e.g., 15 to 45%, 20 to 40%, 40%, 38.5%, 35%, or 31% on a molar basis. A non-limiting example of a sterol 30 is cholesterol. In some embodiments, a lipid nanoparticle formulation includes 0.5% to 20% on a molar basis of the PEG or PEG-modified lipid (e.g., 0.5 to 10%, 0.5 to 5%, 1.5%, 0.5%, 1.5%, 3.5%, or 5% on a molar basis. In some embodiments, a PEG or PEG modified lipid comprises a 35 PEG molecule of an average molecular weight of 2,000 Da. In some embodiments, a PEG or PEG modified lipid comprises a PEG molecule of an average molecular weight of less than 2,000, for example around 1,500 Da, around 1,000 Da, or around 500 Da. Non-limiting examples of PEG- 40 MC3-DMA), modified lipids include PEG-distearoyl glycerol (PEG-DMG) (also referred herein as PEG-C14 or C14-PEG), PEG-cDMA (further discussed in Reyes et al. J. Controlled Release, 107, 276-287 (2005) the contents of which are herein incorporated by reference in their entirety).

In some embodiments, lipid nanoparticle formulations include 25-75% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-50 (dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 0.5-15% of the neutral lipid, 5-50% of the sterol, and 0.5-20% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 35-65% of a cationic lipid selected from 2,2-dili-55 noleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 3-12% of the neutral lipid, 15-45% of the sterol, and 60 0.5-10% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 45-65% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-65 MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319),

74

5-10% of the neutral lipid, 25-40% of the sterol, and 0.5-10% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 60% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 7.5% of the neutral lipid, 31% of the sterol, and 1.5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 50% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 10% of the neutral lipid, 38.5% of the sterol, and 1.5% of the PEG or PEGmodified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 50% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 10% of the neutral lipid, 35% of the sterol, 4.5% or 5% of the PEG or PEG-modified lipid, and 0.5% of the targeting lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 40% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 15% of the neutral lipid, 40% of the sterol, and 5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 57.2% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 7.1% of the neutral lipid, 34.3% of the sterol, and 1.4% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations include 57.5% of a cationic lipid selected from the PEG lipid is PEG-cDMA (PEG-cDMA is further discussed in Reyes et al. (J. Controlled Release, 107, 276-287 (2005), the contents of which are herein incorporated by reference in their entirety), 7.5% of the neutral lipid, 31.5% of the sterol, and 3.5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulations consists essentially of a lipid mixture in molar ratios of 20-70% cationic lipid:5-45% neutral lipid:20-55% cholesterol: 0.5-15% PEG-modified lipid. In some embodiments, lipid nanoparticle formulations consists essentially of a lipid mixture in a molar ratio of 20-60% cationic lipid:5-25% neutral lipid: 25-55% cholesterol: 0.5-15% PEG-modified lipid.

In some embodiments, the molar lipid ratio is 50/10/38.5/1.5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG, PEG-DSG or PEG-DPG), 57.2/7.1134.3/1.4 (mol % cationic lipid/neutral lipid, e.g., DPPC/Chol/PEG-modified lipid, e.g., PEG-cDMA), 40/15/40/5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG), 50/10/35/4.5/0.5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DSG), 50/10/35/5 (cationic

lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG), 40/10/40/10 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA), 35/15/40/10 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA) or 52/13/30/5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA).

Non-limiting examples of lipid nanoparticle compositions and methods of making them are described, for example, in 10 Semple et al. (2010) *Nat. Biotechnol.* 28:172-176; Jayarama et al. (2012), *Angew. Chem. Int. Ed.*, 51: 8529-8533; and Maier et al. (2013) *Molecular Therapy* 21, 1570-1578 (the contents of each of which are incorporated herein by reference in their entirety).

In some embodiments, lipid nanoparticle formulations may comprise a cationic lipid, a PEG lipid and a structural lipid and optionally comprise a non-cationic lipid. As a non-limiting example, a lipid nanoparticle may comprise 40-60% of cationic lipid, 5-15% of a non-cationic lipid, 20 1-2% of a PEG lipid and 30-50% of a structural lipid. As another non-limiting example, the lipid nanoparticle may comprise 50% cationic lipid, 10% non-cationic lipid, 1.5% PEG lipid and 38.5% structural lipid. As yet another non-limiting example, a lipid nanoparticle may comprise 55% cationic lipid, 10% non-cationic lipid, 2.5% PEG lipid and 32.5% structural lipid. In some embodiments, the cationic lipid may be any cationic lipid described herein such as, but not limited to, DLin-KC2-DMA, DLin-MC3-DMA and 1319

In some embodiments, the lipid nanoparticle formulations described herein may be 4 component lipid nanoparticles. The lipid nanoparticle may comprise a cationic lipid, a non-cationic lipid, a PEG lipid and a structural lipid. As a non-limiting example, the lipid nanoparticle may comprise 35 40-60% of cationic lipid, 5-15% of a non-cationic lipid, 1-2% of a PEG lipid and 30-50% of a structural lipid. As another non-limiting example, the lipid nanoparticle may comprise 50% cationic lipid, 10% non-cationic lipid, 1.5% PEG lipid and 38.5% structural lipid. As yet another non- 40 limiting example, the lipid nanoparticle may comprise 55% cationic lipid, 10% non-cationic lipid, 2.5% PEG lipid and 32.5% structural lipid. In some embodiments, the cationic lipid may be any cationic lipid described herein such as, but not limited to, DLin-KC2-DMA, DLin-MC3-DMA and 45 L319.

In some embodiments, the lipid nanoparticle formulations described herein may comprise a cationic lipid, a noncationic lipid, a PEG lipid and a structural lipid. As a non-limiting example, the lipid nanoparticle comprise 50% 50 of the cationic lipid DLin-KC2-DMA, 10% of the noncationic lipid DSPC, 1.5% of the PEG lipid PEG-DOMG and 38.5% of the structural lipid cholesterol. As a nonlimiting example, the lipid nanoparticle comprise 50% of the cationic lipid DLin-MC3-DMA, 10% of the non-cationic 55 lipid DSPC, 1.5% of the PEG lipid PEG-DOMG and 38.5% of the structural lipid cholesterol. As a non-limiting example, the lipid nanoparticle comprise 50% of the cationic lipid DLin-MC3-DMA, 10% of the non-cationic lipid DSPC, 1.5% of the PEG lipid PEG-DMG and 38.5% of the 60 structural lipid cholesterol. As yet another non-limiting example, the lipid nanoparticle comprise 55% of the cationic lipid L319, 10% of the non-cationic lipid DSPC, 2.5% of the PEG lipid PEG-DMG and 32.5% of the structural lipid cholesterol.

Relative amounts of the active ingredient, the pharmaceutically acceptable excipient, and/or any additional ingre76

dients in a vaccine composition may vary, depending upon the identity, size, and/or condition of the subject being treated and further depending upon the route by which the composition is to be administered. For example, the composition may comprise between 0.1% and 99% (w/w) of the active ingredient. By way of example, the composition may comprise between 0.1% and 100%, e.g., between 0.5 and 50%, between 1-30%, between 5-80%, at least 80% (w/w) active ingredient.

In some embodiments, the respiratory virus RNA (e.g. mRNA) vaccine composition may comprise the polynucle-otide described herein, formulated in a lipid nanoparticle comprising MC3, Cholesterol, DSPC and PEG2000-DMG, the buffer trisodium citrate, sucrose and water for injection. As a non-limiting example, the composition comprises: 2.0 mg/mL of drug substance (e.g., polynucleotides encoding H10N8 hMPV), 21.8 mg/mL of MC3, 10.1 mg/mL of cholesterol, 5.4 mg/mL of DSPC, 2.7 mg/mL of PEG2000-DMG, 5.16 mg/mL of trisodium citrate, 71 mg/mL of sucrose and 1.0 mL of water for injection.

In some embodiments, a nanoparticle (e.g., a lipid nanoparticle) has a mean diameter of 10-500 nm, 20-400 nm, 30-300 nm, 40-200 nm. In some embodiments, a nanoparticle (e.g., a lipid nanoparticle) has a mean diameter of 50-150 nm, 50-200 nm, 80-100 nm or 80-200 nm.

Liposomes, Lipoplexes, and Lipid Nanoparticles

The RNA (e.g., mRNA) vaccines of the disclosure can be formulated using one or more liposomes, lipoplexes, or lipid nanoparticles. In some embodiments, pharmaceutical compositions of RNA (e.g., mRNA) vaccines include liposomes. Liposomes are artificially-prepared vesicles which may primarily be composed of a lipid bilayer and may be used as a delivery vehicle for the administration of nutrients and pharmaceutical formulations. Liposomes can be of different sizes such as, but not limited to, a multilamellar vesicle (MLV) which may be hundreds of nanometers in diameter and may contain a series of concentric bilayers separated by narrow aqueous compartments, a small unicellular vesicle (SUV) which may be smaller than 50 nm in diameter, and a large unilamellar vesicle (LUV) which may be between 50 and 500 nm in diameter. Liposome design may include, but is not limited to, opsonins or ligands in order to improve the attachment of liposomes to unhealthy tissue or to activate events such as, but not limited to, endocytosis. Liposomes may contain a low or a high pH in order to improve the delivery of the pharmaceutical formulations.

The formation of liposomes may depend on the physicochemical characteristics such as, but not limited to, the pharmaceutical formulation entrapped and the liposomal ingredients, the nature of the medium in which the lipid vesicles are dispersed, the effective concentration of the entrapped substance and its potential toxicity, any additional processes involved during the application and/or delivery of the vesicles, the optimization size, polydispersity and the shelf-life of the vesicles for the intended application, and the batch-to-batch reproducibility and possibility of large-scale production of safe and efficient liposomal products.

In some embodiments, pharmaceutical compositions described herein may include, without limitation, liposomes such as those formed from 1,2-dioleyloxy-N,N-dimethylaminopropane (DODMA) liposomes, DiLa2 liposomes from Marina Biotech (Bothell, Wash.), 1,2-dilinoleyloxy-3-dimethylaminopropane (DLin-DMA), 2,2-dilinoleyl-4-(2-dimethylaminoethyl)-[1,3]-dioxolane (DLin-KC2-DMA), and MC3 (US20100324120; herein incorporated by reference in its entirety) and liposomes which may deliver small

77 molecule drugs such as, but not limited to, DOXIL® from Janssen Biotech, Inc. (Horsham, Pa.).

In some embodiments, pharmaceutical compositions described herein may include, without limitation, liposomes such as those formed from the synthesis of stabilized plas- 5 mid-lipid particles (SPLP) or stabilized nucleic acid lipid particle (SNALP) that have been previously described and shown to be suitable for oligonucleotide delivery in vitro and in vivo (see Wheeler et al. Gene Therapy. 1999 6:271-281; Zhang et al. Gene Therapy. 1999 6:1438-1447; Jeffs et 10 al. Pharm Res. 2005 22:362-372; Morrissey et al., Nat Biotechnol. 2005 2:1002-1007; Zimmermann et al., Nature. 2006 441:111-114; Heyes et al. J Contr Rel. 2005 107:276-287; Semple et al. Nature Biotech. 2010 28:172-176; Judge et al. J Clin Invest. 2009 119:661-673; deFougerolles Hum 15 Gene Ther. 2008 19:125-132; U.S. Patent Publication No US20130122104; all of which are incorporated herein in their entireties). The original manufacture method by Wheeler et al. was a detergent dialysis method, which was later improved by Jeffs et al. and is referred to as the 20 spontaneous vesicle formation method. The liposome formulations are composed of 3 to 4 lipid components in addition to the polynucleotide. As an example a liposome can contain, but is not limited to, 55% cholesterol, 20% disteroylphosphatidyl choline (DSPC), 10% PEG-S-DSG, 25 1,2-dioleyloxy-N,N-dimethylaminopropane (DODMA), as described by Jeffs et al. As another example, certain liposome formulations may contain, but are not limited to, 48% cholesterol, 20% DSPC, 2% PEG-c-DMA, and 30% cationic lipid, where the cationic lipid can be 30 1,2-distearloxy-N,N-dimethylaminopropane (DSDMA), DODMA, DLin-DMA, or 1,2-dilinolenyloxy-3-dimethylaminopropane (DLenDMA), as described by Heyes et al.

In some embodiments, liposome formulations may comprise from about 25.0% cholesterol to about 40.0% choles- 35 terol, from about 30.0% cholesterol to about 45.0% cholesterol, from about 35.0% cholesterol to about 50.0% cholesterol and/or from about 48.5% cholesterol to about 60% cholesterol. In some embodiments, formulations may comprise a percentage of cholesterol selected from the group 40 consisting of 28.5%, 31.5%, 33.5%, 36.5%, 37.0%, 38.5%, 39.0% and 43.5%. In some embodiments, formulations may comprise from about 5.0% to about 10.0% DSPC and/or from about 7.0% to about 15.0% DSPC.

In some embodiments, the RNA (e.g., mRNA) vaccine 45 pharmaceutical compositions may be formulated in liposomes such as, but not limited to, DiLa2 liposomes (Marina Biotech, Bothell, Wash.), SMARTICLES® (Marina Biotech, Bothell, Wash.), neutral DOPC (1,2-dioleoyl-sn-glycero-3-phosphocholine) based liposomes (e.g., siRNA deliv- 50 ery for ovarian cancer (Landen et al. Cancer Biology & Therapy 2006 5(12)1708-1713); herein incorporated by reference in its entirety) and hyaluronan-coated liposomes (Quiet Therapeutics, Israel).

In some embodiments, the cationic lipid may be a low 55 molecular weight cationic lipid such as those described in U.S. Patent Application No. 20130090372, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in a lipid vesicle, which may have 60 crosslinks between functionalized lipid bilayers.

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in a lipid-polycation complex. The formation of the lipid-polycation complex may be accomplished by methods known in the art and/or as described in 65 U.S. Pub. No. 20120178702, herein incorporated by reference in its entirety. As a non-limiting example, the polyca78

tion may include a cationic peptide or a polypeptide such as, but not limited to, polylysine, polyornithine and/or polyarginine. In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in a lipid-polycation complex, which may further include a non-cationic lipid such as, but not limited to, cholesterol or dioleovl phosphatidylethanolamine (DOPE).

In some embodiments, the ratio of PEG in the lipid nanoparticle (LNP) formulations may be increased or decreased and/or the carbon chain length of the PEG lipid may be modified from C14 to C18 to alter the pharmacokinetics and/or biodistribution of the LNP formulations. As a non-limiting example, LNP formulations may contain from about 0.5% to about 3.0%, from about 1.0% to about 3.5%, from about 1.5% to about 4.0%, from about 2.0% to about 4.5%, from about 2.5% to about 5.0% and/or from about 3.0% to about 6.0% of the lipid molar ratio of PEG-c-DOMG  $(R-3-[(\omega-methoxy-poly(ethyleneglycol)$ 2000)carbamoyl)]-1,2-dimyristyloxypropyl-3-amine) (also referred to herein as PEG-DOMG) as compared to the cationic lipid, DSPC and cholesterol. In some embodiments, the PEG-c-DOMG may be replaced with a PEG lipid such as, but not limited to, PEG-DSG (1,2-Distearoyl-sn-glycerol, methoxypolyethylene glycol), PEG-DMG (1,2-Dimyristoyl-sn-glycerol) and/or PEG-DPG (1,2-Dipalmitoyl-sn-glycerol, methoxypolyethylene glycol). The cationic lipid may be selected from any lipid known in the art such as, but not limited to, DLin-MC3-DMA, DLin-DMA, C12-200 and DLin-KC2-DMA.

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in a lipid nanoparticle.

In some embodiments, the RNA (e.g., mRNA) vaccine formulation comprising the polynucleotide is a nanoparticle which may comprise at least one lipid. The lipid may be selected from, but is not limited to, DLin-DMA, DLin-K-DMA, 98N12-5, C12-200, DLin-MC3-DMA, DLin-KC2-DMA, DODMA, PLGA, PEG, PEG-DMG, PEGylated lipids and amino alcohol lipids. In another aspect, the lipid may be a cationic lipid such as, but not limited to, DLin-DMA, DLin-D-DMA, DLin-MC3-DMA, DLin-KC2-DMA, DODMA and amino alcohol lipids. The amino alcohol cationic lipid may be the lipids described in and/or made by the methods described in U.S. Patent Publication No. US20130150625, herein incorporated by reference in its entirety. As a non-limiting example, the cationic lipid may 2-amino-3-[(9Z,12Z)-octadeca-9,12-dien-1-vloxy]-2-{ [(9Z,2Z)-octadeca-9,12-dien-1-yloxy]methyl}propan-1-ol (Compound 1 in US20130150625); 2-amino-3-[(9Z)-octadec-9-en-1-yloxy]-2- $\{[(9Z)-octadec-9-en-1-yloxy]$ methyl\propan-1-ol (Compound 2 in US20130150625); 2-amino-3-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-2-[(octyloxy)methyl]propan-1-ol (Compound US20130150625); and 2-(dimethylamino)-3-[(9Z,12Z)-oc $tadeca-9,12-dien-1-yloxy]-2-\big\{[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-2-\big\}[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-2-\big][(9Z,12Z)-octadeca-9,12-0-yloxy]-2-\big][(9$ dien-1-yloxy]methyl}propan-1-ol (Compound US20130150625); or any pharmaceutically acceptable salt or stereoisomer thereof.

Lipid nanoparticle formulations typically comprise a lipid, in particular, an ionizable cationic lipid, for example, 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), or di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), and further comprise a neutral lipid, a sterol and a molecule capable of reducing particle aggregation, for example a PEG or PEG-modified lipid.

In some embodiments, the lipid nanoparticle formulation consists essentially of (i) at least one lipid selected from the group consisting of 2,2-dilinoleyl-4-dimethylaminoethyl-[1, 3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-5 en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy) heptadecanedioate (L319); (ii) a neutral lipid selected from DSPC, DPPC, POPC, DOPE and SM; (iii) a sterol, e.g., cholesterol; and (iv) a PEG-lipid, e.g., PEG-DMG or PEG-cDMA, in a molar ratio of about 20-60% cationic lipid:5-10 25% neutral lipid:25-55% sterol; 0.5-15% PEG-lipid.

In some embodiments, the formulation includes from about 25% to about 75% on a molar basis of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), e.g., from about 35 to about 65%, from about 45 to about 65%, about 60%, about 57.5%, about 50% or about 40% on a molar basis.

In some embodiments, the formulation includes from about 0.5% to about 15% on a molar basis of the neutral lipid e.g., from about 3 to about 12%, from about 5 to about 10% or about 15%, about 10%, or about 7.5% on a molar basis. Examples of neutral lipids include, but are not limited to, 25 DSPC, POPC, DPPC, DOPE and SM. In some embodiments, the formulation includes from about 5% to about 50% on a molar basis of the sterol (e.g., about 15 to about 45%, about 20 to about 40%, about 40%, about 38.5%, about 35%, or about 31% on a molar basis. An exemplary sterol is cholesterol. In some embodiments, the formulation includes from about 0.5% to about 20% on a molar basis of the PEG or PEG-modified lipid (e.g., about 0.5 to about 10%, about 0.5 to about 5%, about 1.5%, about 0.5%, about 1.5%, about 3.5%, or about 5% on a molar basis. In some embodiments, 35 the PEG or PEG modified lipid comprises a PEG molecule of an average molecular weight of 2,000 Da. In other embodiments, the PEG or PEG modified lipid comprises a PEG molecule of an average molecular weight of less than 2,000, for example around 1,500 Da, around 1,000 Da, or 40 around 500 Da. Examples of PEG-modified lipids include, but are not limited to, PEG-distearoyl glycerol (PEG-DMG) (also referred herein as PEG-C14 or C14-PEG), PEGcDMA (further discussed in Reyes et al. J. Controlled Release, 107, 276-287 (2005) the contents of which are 45 herein incorporated by reference in their entirety)

In some embodiments, the formulations of the present disclosure include 25-75% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 0.5-15% of the neutral lipid, 5-50% of the sterol, and 0.5-20% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present 55 disclosure include 35-65% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 60 3-12% of the neutral lipid, 15-45% of the sterol, and 0.5-10% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include 45-65% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-

80

(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), 5-10% of the neutral lipid, 25-40% of the sterol, and 0.5-10% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include about 60% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), about 7.5% of the neutral lipid, about 31% of the sterol, and about 1.5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include about 50% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), about 10% of the neutral lipid, about 38.5% of the sterol, and about 1.5% of the PEG or PEG-modified lipid on a molar basis

In some embodiments, the formulations of the present disclosure include about 50% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), about 10% of the neutral lipid, about 35% of the sterol, about 4.5% or about 5% of the PEG or PEG-modified lipid, and about 0.5% of the targeting lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include about 40% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), about 15% of the neutral lipid, about 40% of the sterol, and about 5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include about 57.2% of a cationic lipid selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319), about 7.1% of the neutral lipid, about 34.3% of the sterol, and about 1.4% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, the formulations of the present disclosure include about 57.5% of a cationic lipid selected from the PEG lipid is PEG-cDMA (PEG-cDMA is further discussed in Reyes et al. (J. Controlled Release, 107, 276-287 (2005), the contents of which are herein incorporated by reference in their entirety), about 7.5% of the neutral lipid, about 31.5% of the sterol, and about 3.5% of the PEG or PEG-modified lipid on a molar basis.

In some embodiments, lipid nanoparticle formulation consists essentially of a lipid mixture in molar ratios of about 20-70% cationic lipid:5-45% neutral lipid:20-55% cholesterol: 0.5-15% PEG-modified lipid; more preferably in a molar ratio of about 20-60% cationic lipid:5-25% neutral lipid:25-55% cholesterol: 0.5-15% PEG-modified lipid.

In some embodiments, the molar lipid ratio is approximately 50/10/38.5/1.5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG, PEG-DSG or PEG-DPG), 57.2/7.1134.3/1.4 (mol % cationic lipid/neutral lipid, e.g., DPPC/Chol/PEG-modified lipid,

e.g., PEG-cDMA), 40/15/40/5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG), 50/10/35/4.5/0.5 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DSG), 50/10/35/5 (cationic lipid/neutral lipid, e.g., DSPC/Chol/ 5 PEG-modified lipid, e.g., PEG-DMG), 40/10/40/10 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA), 35/15/40/10 (mol % cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA) or 52/13/30/5 (mol % 10 cationic lipid/neutral lipid, e.g., DSPC/Chol/PEG-modified lipid, e.g., PEG-DMG or PEG-cDMA).

Examples of lipid nanoparticle compositions and methods of making same are described, for example, in Semple et al. (2010) *Nat. Biotechnol.* 28:172-176; Jayarama et al. (2012), 15 *Angew. Chem. Int. Ed.*, 51: 8529-8533; and Maier et al. (2013) *Molecular Therapy* 21, 1570-1578 (the contents of each of which are incorporated herein by reference in their entirety).

In some embodiments, the lipid nanoparticle formulations 20 described herein may comprise a cationic lipid, a PEG lipid and a structural lipid and optionally comprise a non-cationic lipid. As a non-limiting example, the lipid nanoparticle may comprise about 40-60% of cationic lipid, about 5-15% of a non-cationic lipid, about 1-2% of a PEG lipid and about 25 30-50% of a structural lipid. As another non-limiting example, the lipid nanoparticle may comprise about 50% cationic lipid, about 10% non-cationic lipid, about 1.5% PEG lipid and about 38.5% structural lipid. As yet another non-limiting example, the lipid nanoparticle may comprise 30 about 55% cationic lipid, about 10% non-cationic lipid, about 2.5% PEG lipid and about 32.5% structural lipid. In some embodiments, the cationic lipid may be any cationic lipid described herein such as, but not limited to, DLin-KC2-DMA, DLin-MC3-DMA and L319.

In some embodiments, the lipid nanoparticle formulations described herein may be 4 component lipid nanoparticles. The lipid nanoparticle may comprise a cationic lipid, a non-cationic lipid, a PEG lipid and a structural lipid. As a non-limiting example, the lipid nanoparticle may comprise 40 about 40-60% of cationic lipid, about 5-15% of a noncationic lipid, about 1-2% of a PEG lipid and about 30-50% of a structural lipid. As another non-limiting example, the lipid nanoparticle may comprise about 50% cationic lipid, about 10% non-cationic lipid, about 1.5% PEG lipid and 45 about 38.5% structural lipid. As yet another non-limiting example, the lipid nanoparticle may comprise about 55% cationic lipid, about 10% non-cationic lipid, about 2.5% PEG lipid and about 32.5% structural lipid. In some embodiments, the cationic lipid may be any cationic lipid described 50 herein such as, but not limited to, DLin-KC2-DMA, DLin-MC3-DMA and L319.

In some embodiments, the lipid nanoparticle formulations described herein may comprise a cationic lipid, a non-cationic lipid, a PEG lipid and a structural lipid. As a 55 non-limiting example, the lipid nanoparticle comprise about 50% of the cationic lipid DLin-KC2-DMA, about 10% of the non-cationic lipid DSPC, about 1.5% of the PEG lipid PEG-DOMG and about 38.5% of the structural lipid cholesterol. As a non-limiting example, the lipid nanoparticle comprise about 50% of the cationic lipid DLin-MC3-DMA, about 10% of the non-cationic lipid DSPC, about 1.5% of the PEG lipid PEG-DOMG and about 38.5% of the structural lipid cholesterol. As a non-limiting example, the lipid nanoparticle comprise about 50% of the cationic lipid DLin-MC3-DMA, about 10% of the non-cationic lipid DLin-MC3-DMA, about 10% of the non-cationic lipid DSPC, about 1.5% of the PEG lipid PEG-DMG and about 38.5% of

the structural lipid cholesterol. As yet another non-limiting example, the lipid nanoparticle comprise about 55% of the cationic lipid L319, about 10% of the non-cationic lipid DSPC, about 2.5% of the PEG lipid PEG-DMG and about 32.5% of the structural lipid cholesterol.

As a non-limiting example, the cationic lipid may be selected from (20Z,23Z)—N,N-dimethylnonacosa-20,23dien-10-amine, (17Z,20Z)-N,N-dimemylhexacosa-17,20dien-9-amine, (1Z,19Z)—N5N-dimethylpentacosa-16, 19-dien-8-amine, (13Z,16Z)—N,N-dimethyldocosa-13,16-(12Z,15Z)—N,N-dimethylhenicosa-12,15dien-5-amine, dien-4-amine, (14Z,17Z)—N,N-dimethyltricosa-14,17-(15Z,18Z)—N,N-dimethyltetracosa-15,18dien-6-amine, dien-7-amine, (18Z,21Z)—N,N-dimethylheptacosa-18,21dien-10-amine, (15Z,18Z)—N,N-dimethyltetracosa-15,18dien-5-amine, (14Z,17Z)—N,N-dimethyltricosa-14,17dien-4-amine, (19Z,22Z)—N,N-dimeihyloctacosa-19,22dien-9-amine, (18Z,21 Z)-N,N-dimethylheptacosa-18,21dien-8 amine, (17Z,20Z)—N,N-dimethylhexacosa-17,20dien-7-amine, (16Z,19Z)—N.N-dimethylpentacosa-16,19dien-6-amine, (22Z,25Z)—N,N-dimethylhentriaconta-22, 25-dien-10-amine, (21 Z,24Z)—N,N-dimethyltriaconta-21, 24-dien-9-amine, (18Z)—N,N-dimetylheptacos-18-en-10amine, (17Z)—N,N-dimethylhexacos-17-en-9-amine, (19Z, 22Z)—N,N-dimethyloctacosa-19,22-dien-7-amine, dimethylheptacosan-10-amine, (20Z,23Z)—N-ethyl-Nmethylnonacosa-20,23-dien-10-amine, 1-[(11Z,14Z)-1nonylicosa-11,14-dien-1-yl]pyrrolidine, (20Z)-N.Ndimethylheptacos-20-en-10-amine, (15Z)—N,N-dimethyl eptacos-15-en-10-amine, (14Z)—N,N-dimethylnonacos-14en-10-amine, (17Z)—N,N-dimethylnonacos-17-en-10-(24Z)—N,N-dimethyltritriacont-24-en-10-amine, amine, (20Z)—N,N-dimethylnonacos-20-en-10-amine, (22Z)—N, N-dimethylhentriacont-22-en-10-amine, (16Z)—N,N-dim-35 ethylpentacos-16-en-8-amine, (12Z,15Z)—N,N-dimethyl-2-nonylhenicosa-12,15-dien-1-amine, (13Z,16Z)—N,Ndimethyl-3-nonyldocosa-13,16-dien-1 N,Namine. dimethyl-1-[(1S,2R)-2-octylcyclopropyl]eptadecan-8-1-[(1S,2R)-2-hexylcyclopropyl]-N,Namine, dimethylnonadecan-10-amine, N,N-dimethyl-1-[(1S,2R)-2octylcyclopropyl]nonadecan-10-amine, N,N-dimethyl-21-[(1S,2R)-2-octylcyclopropyl]henicosan-10-amine, dimethyl-1-[(1S,2S)-2-{[(1R,2R)-2-pentylcyclopropyl] methyl\cyclopropyl\nonadecan-10-amine,N,N-dimethyl-1-[(1S,2R)-2-octylcyclopropyl]hexadecan-8-amine, N,Ndimethyl-[(1R,2S)-2-undecylcyclopropyl]tetradecan-5-N,N-dimethyl-3-{7-[(1S,2R)-2-octylcyclopropyl] heptyl\dodecan-1-amine, 1-[(1R,2S)-2-heptylcyclopropyl]-N,N-dimethyloctadecan-9-amine, 1-[(1S,2R)-2decylcyclopropyl]-N,N-dimethylpentadecan-6-amine, N,Ndimethyl-1-R1S,2R)-2-octylcyclopropyllpentadecan-8amine, R—N,N-dimethyl-1-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-3-(octyloxy)propan-2-amine, S—N,N-dimethyl-1-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-3-(octyloxy) 1-{2-[(9Z,12Z)-octadeca-9,12-dien-1propan-2-amine, yloxy]-1-[(octyloxy)methyl]ethyl}pyrrolidine, (2S)—N,Ndimethyl-1-[(9Z,12Z)-octadeca-9,12-dien-1-yloxy]-3-[(5Z)-oct-5-en-1-yloxy]propan-2-amine,  $1-\{2-[(9Z,12Z)-(9Z,12Z)$ octadeca-9,12-dien-1-yloxy]-1-[(octyloxy)methyl] ethyl}azetidine, (2S)-1-(hexyloxy)-N,N-dimethyl-3-R9Z, 12Z)-octadeca-9,12-dien-1-yloxylpropan-2-amine, (2S)-1-(heptyloxy)-N,N-dimethyl-3-R9Z,12Z)-octadeca-9,12dien-1-yloxylpropan-2-amine, N,N-dimethyl-1-(nonyloxy)-3-R9Z,12Z)-octadeca-9,12-dien-1-yloxylpropan-2-amine, N,N-dimethyl-1-[(9Z)-octadec-9-en-1-yloxy]-3-(octyloxy) propan-2-amine; (2S)-N,N-dimethyl-1-[(6Z,9Z,12Z)-octa-

deca-6,9,12-trien-1-yloxy]-3-(octyloxy)propan-2-amine,

(2S)-1-[(11Z,14Z)-icosa-11,14-dien-1-yloxy]-N,N-dimethyl-3-(pentyloxy)propan-2-amine, (2S)-1-(hexyloxy)-3-[(11Z,14Z)-icosa-11,14-dien-1-yloxy]-N,N-dimethylpropan-2-amine, 1-[(11Z,14Z)-icosa-11,14-dien-1-yloxy]-N,N-1-[(13Z,16Z)- 5 dimethyl-3-(octyloxy)propan-2-amine, docosa-13,16-dien-1-yloxy]-N,N-dimethyl-3-(octyloxy) propan-2-amine, (2S)-1-[(13Z,16Z)-docosa-13,16-dien-1yloxy]-3-(hexyloxy)-N,N-dimethylpropan-2-amine, (2S)-1-[(13Z)-docos-13-en-1-yloxy]-3-(hexyloxy)-N,Ndimethylpropan-2-amine, 1-[(13Z)-docos-13-en-1-yloxy]- 10 N,N-dimethyl-3-(octyloxy)propan-2-amine, 1-[(9Z)hexadec-9-en-1-yloxy]-N,N-dimethyl-3-(octyloxy)propan-2-amine, (2R)-N,N-dimethyl-H(1-metoyloctyl)oxy]-3-[(9Z, 12Z)-octadeca-9,12-dien-1-yloxy]propan-2-amine, (2R)-1-[(3,7-dimethyloctyl)oxy]-N,N-dimethyl-3-R9Z,12Z)octadeca-9,12-dien-1-yloxylpropan-2-amine, N,Ndimethyl-1-(octyloxy)-3-({8-R1S,25})-2-{[(1R,2R)-2pentylcyclopropyl]methyl}cyclopropyl]octyl}oxy)propan-2-amine, N,N-dimethyl-1-1 [8-(2-oc1ylcyclopropyl)octyl] oxy\-3-(octyloxy)propan-2-amine and (11E,20Z,23Z)—N, 20 N-dimethylnonacosa-11,20,2-trien-10-amine pharmaceutically acceptable salt or stereoisomer thereof.

In some embodiments, the LNP formulations of the RNA (e.g., mRNA) vaccines may contain PEG-c-DOMG at 3% lipid molar ratio. In some embodiments, the LNP formulations of the RNA (e.g., mRNA) vaccines may contain PEG-c-DOMG at 1.5% lipid molar ratio.

In some embodiments, the pharmaceutical compositions of the RNA (e.g., mRNA) vaccines may include at least one of the PEGylated lipids described in International Publication No. WO2012099755, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the LNP formulation may contain PEG-DMG 2000 (1,2-dimyristoyl-sn-glycero-3-phophoethanolamine-N-[methoxy(polyethylene glycol)-2000). In 35 some embodiments, the LNP formulation may contain PEG-DMG 2000, a cationic lipid known in the art and at least one other component. In some embodiments, the LNP formulation may contain PEG-DMG 2000, a cationic lipid known in the art, DSPC and cholesterol. As a non-limiting example, 40 the LNP formulation may contain PEG-DMG 2000, DLin-DMA, DSPC and cholesterol. As another non-limiting example the LNP formulation may contain PEG-DMG 2000, DLin-DMA, DSPC and cholesterol in a molar ratio of 2:40:10:48 (see e.g., Geall et al., Nonviral delivery of 45 self-amplifying RNA (e.g., mRNA) vaccines, PNAS 2012; PMID: 22908294, the contents of each of which are herein incorporated by reference in their entirety).

The lipid nanoparticles described herein may be made in a sterile environment.

In some embodiments, the LNP formulation may be formulated in a nanoparticle such as a nucleic acid-lipid particle. As a non-limiting example, the lipid particle may comprise one or more active agents or therapeutic agents; one or more cationic lipids comprising from about 50 mol % 55 to about 85 mol % of the total lipid present in the particle; one or more non-cationic lipids comprising from about 13 mol % to about 49.5 mol % of the total lipid present in the particle; and one or more conjugated lipids that inhibit aggregation of particles comprising from about 0.5 mol % to 60 about 2 mol % of the total lipid present in the particle.

The nanoparticle formulations may comprise a phosphate conjugate. The phosphate conjugate may increase in vivo circulation times and/or increase the targeted delivery of the nanoparticle. As a non-limiting example, the phosphate 65 conjugates may include a compound of any one of the formulas described in International Application No.

84

WO2013033438, the contents of which are herein incorporated by reference in its entirety.

The nanoparticle formulation may comprise a polymer conjugate. The polymer conjugate may be a water soluble conjugate. The polymer conjugate may have a structure as described in U.S. Patent Application No. 20130059360, the contents of which are herein incorporated by reference in its entirety. In some embodiments, polymer conjugates with the polynucleotides of the present disclosure may be made using the methods and/or segmented polymeric reagents described in U.S. Patent Application No. 20130072709, the contents of which are herein incorporated by reference in its entirety. In some embodiments, the polymer conjugate may have pendant side groups comprising ring moieties such as, but not limited to, the polymer conjugates described in U.S. Patent Publication No. US20130196948, the contents which are herein incorporated by reference in its entirety.

The nanoparticle formulations may comprise a conjugate to enhance the delivery of nanoparticles of the present disclosure in a subject. Further, the conjugate may inhibit phagocytic clearance of the nanoparticles in a subject. In one aspect, the conjugate may be a "self" peptide designed from the human membrane protein CD47 (e.g., the "self" particles described by Rodriguez et al. (Science 2013 339, 971-975), herein incorporated by reference in its entirety). As shown by Rodriguez et al., the self peptides delayed macrophagemediated clearance of nanoparticles which enhanced delivery of the nanoparticles. In another aspect, the conjugate may be the membrane protein CD47 (e.g., see Rodriguez et al. Science 2013 339, 971-975, herein incorporated by reference in its entirety). Rodriguez et al. showed that, similarly to "self" peptides, CD47 can increase the circulating particle ratio in a subject as compared to scrambled peptides and PEG coated nanoparticles.

In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure are formulated in nanoparticles which comprise a conjugate to enhance the delivery of the nanoparticles of the present disclosure in a subject. The conjugate may be the CD47 membrane or the conjugate may be derived from the CD47 membrane protein, such as the "self" peptide described previously. In some embodiments, the nanoparticle may comprise PEG and a conjugate of CD47 or a derivative thereof. In some embodiments, the nanoparticle may comprise both the "self" peptide described above and the membrane protein CD47.

In some embodiments, a "self" peptide and/or CD47 protein may be conjugated to a virus-like particle or pseudovirion, as described herein for delivery of the RNA (e.g., mRNA) vaccines of the present disclosure.

In some embodiments, RNA (e.g., mRNA) vaccine pharmaceutical compositions comprising the polynucleotides of the present disclosure and a conjugate that may have a degradable linkage. Non-limiting examples of conjugates include an aromatic moiety comprising an ionizable hydrogen atom, a spacer moiety, and a water-soluble polymer. As a non-limiting example, pharmaceutical compositions comprising a conjugate with a degradable linkage and methods for delivering such pharmaceutical compositions are described in U.S. Patent Publication No. US20130184443, the contents of which are herein incorporated by reference in their entirety.

The nanoparticle formulations may be a carbohydrate nanoparticle comprising a carbohydrate carrier and a RNA (e.g., mRNA) vaccine. As a non-limiting example, the carbohydrate carrier may include, but is not limited to, an anhydride-modified phytoglycogen or glycogen-type material, phtoglycogen octenyl succinate, phytoglycogen beta-

dextrin, anhydride-modified phytoglycogen beta-dextrin. (See e.g., International Publication No. WO2012109121; the contents of which are herein incorporated by reference in their entirety).

Nanoparticle formulations of the present disclosure may 5 be coated with a surfactant or polymer in order to improve the delivery of the particle. In some embodiments, the nanoparticle may be coated with a hydrophilic coating such as, but not limited to, PEG coatings and/or coatings that have a neutral surface charge. The hydrophilic coatings may help 10 to deliver nanoparticles with larger payloads such as, but not limited to, RNA (e.g., mRNA) vaccines within the central nervous system. As a non-limiting example nanoparticles comprising a hydrophilic coating and methods of making such nanoparticles are described in U.S. Patent Publication 15 No. US20130183244, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the lipid nanoparticles of the present disclosure may be hydrophilic polymer particles. Non-limiting examples of hydrophilic polymer particles and 20 methods of making hydrophilic polymer particles are described in U.S. Patent Publication No. US20130210991, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the lipid nanoparticles of the 25 present disclosure may be hydrophobic polymer particles.

Lipid nanoparticle formulations may be improved by replacing the cationic lipid with a biodegradable cationic lipid which is known as a rapidly eliminated lipid nanoparticle (reLNP). Ionizable cationic lipids, such as, but not 30 limited to, DLinDMA, DLin-KC2-DMA, and DLin-MC3-DMA, have been shown to accumulate in plasma and tissues over time and may be a potential source of toxicity. The rapid metabolism of the rapidly eliminated lipids can improve the tolerability and therapeutic index of the lipid 35 nanoparticles by an order of magnitude from a 1 mg/kg dose to a 10 mg/kg dose in rat. Inclusion of an enzymatically degraded ester linkage can improve the degradation and metabolism profile of the cationic component, while still maintaining the activity of the reLNP formulation. The ester 40 linkage can be internally located within the lipid chain or it may be terminally located at the terminal end of the lipid chain. The internal ester linkage may replace any carbon in the lipid chain.

In some embodiments, the internal ester linkage may be 45 located on either side of the saturated carbon.

In some embodiments, an immune response may be elicited by delivering a lipid nanoparticle which may include a nanospecies, a polymer and an immunogen. (U.S. Publication No. 20120189700 and International Publication No. 50 WO2012099805; each of which is herein incorporated by reference in their entirety). The polymer may encapsulate the nanospecies or partially encapsulate the nanospecies. The immunogen may be a recombinant protein, a modified RNA and/or a polynucleotide described herein. In some 55 embodiments, the lipid nanoparticle may be formulated for use in a vaccine such as, but not limited to, against a pathogen.

Lipid nanoparticles may be engineered to alter the surface properties of particles so the lipid nanoparticles may penetrate the mucosal barrier. Mucus is located on mucosal tissue such as, but not limited to, oral (e.g., the buccal and esophageal membranes and tonsil tissue), ophthalmic, gastrointestinal (e.g., stomach, small intestine, large intestine, colon, rectum), nasal, respiratory (e.g., nasal, pharyngeal, 65 tracheal and bronchial membranes), genital (e.g., vaginal, cervical and urethral membranes). Nanoparticles larger than

86

10-200 nm which are preferred for higher drug encapsulation efficiency and the ability to provide the sustained delivery of a wide array of drugs have been thought to be too large to rapidly diffuse through mucosal barriers. Mucus is continuously secreted, shed, discarded or digested and recycled so most of the trapped particles may be removed from the mucosa tissue within seconds or within a few hours. Large polymeric nanoparticles (200 nm-500 nm in diameter) which have been coated densely with a low molecular weight polyethylene glycol (PEG) diffused through mucus only 4 to 6-fold lower than the same particles diffusing in water (Lai et al. PNAS 2007 104(5):1482-487; Lai et al. Adv Drug Deliv Rev. 2009 61(2): 158-171; each of which is herein incorporated by reference in their entirety). The transport of nanoparticles may be determined using rates of permeation and/or fluorescent microscopy techniques including, but not limited to, fluorescence recovery after photobleaching (FRAP) and high resolution multiple particle tracking (MPT). As a non-limiting example, compositions which can penetrate a mucosal barrier may be made as described in U.S. Pat. No. 8,241,670 or International Patent Publication No. WO2013110028, the contents of each of which are herein incorporated by reference in its entirety.

The lipid nanoparticle engineered to penetrate mucus may comprise a polymeric material (i.e. a polymeric core) and/or a polymer-vitamin conjugate and/or a tri-block co-polymer. The polymeric material may include, but is not limited to, polyamines, polyethers, polyamides, polyesters, polycarbamates, polyureas, polycarbonates, poly(styrenes), polyimides, polysulfones, polyurethanes, polyacetylenes, polyethylenes, polyethyeneimines, polyisocyanates, polyacrylates, polymethacrylates, polyacrylonitriles, and polyarylates. The polymeric material may be biodegradable and/or biocompatible. Non-limiting examples of biocompatible polymers are described in International Patent Publication No. WO2013116804, the contents of which are herein incorporated by reference in their entirety. The polymeric material may additionally be irradiated. As a non-limiting example, the polymeric material may be gamma irradiated (see e.g., International App. No. WO201282165, herein incorporated by reference in its entirety). Non-limiting examples of specific polymers include poly(caprolactone) (PCL), ethylene vinyl acetate polymer (EVA), poly(lactic acid) (PLA), poly(L-lactic acid) (PLLA), poly(glycolic acid) (PGA), poly (lactic acid-co-glycolic acid) (PLGA), poly(L-lactic acidco-glycolic acid) (PLLGA), poly(D,L-lactide) (PDLA), poly (L-lactide) (PLLA), poly(D,L-lactide-co-caprolactone), poly(D,L-lactide-co-caprolactone-co-glycolide), poly(D,Llactide-co-PEO-co-D,L-lactide), poly(D,L-lactide-co-PPOco-D,L-lactide), polyalkyl cyanoacralate, polyurethane, poly-L-lysine (PLL), hydroxypropyl methacrylate (HPMA), polyethyleneglycol, poly-L-glutamic acid, poly(hydroxy acids), polyanhydrides, polyorthoesters, poly(ester amides), polyamides, poly(ester ethers), polycarbonates, polyalkylenes such as polyethylene and polypropylene, polyalkylene glycols such as poly(ethylene glycol) (PEG), polyalkylene oxides (PEO), polyalkylene terephthalates such as poly(ethylene terephthalate), polyvinyl alcohols (PVA), polyvinyl ethers, polyvinyl esters such as poly(vinyl acetate), polyvinyl halides such as poly(vinyl chloride) (PVC), polyvinylpyrrolidone, polysiloxanes, polystyrene (PS), polyurethanes, derivatized celluloses such as alkyl celluloses, hydroxyalkyl celluloses, cellulose ethers, cellulose esters, nitro celluloses, hydroxypropylcellulose, carboxymethylcellulose, polymers of acrylic acids, such as poly(methyl(meth)acrylate) (PMMA), poly(ethyl(meth)

acrylate), poly(butyl(meth)acrylate), poly(isobutyl(meth) acrylate), poly(hexyl(meth)acrylate), poly(isodecyl(meth) acrylate), poly(lauryl(meth)acrylate), poly(phenyl(meth) acrylate), poly(methyl acrylate), poly(isopropyl acrylate), poly(isobutyl acrylate), poly(octadecyl acrylate) and copo- 5 lymers and mixtures thereof, polydioxanone and its copolymers, polyhydroxyalkanoates, polypropylene fumarate, polyoxymethylene, poloxamers, poly(ortho)esters, poly(butyric acid), poly(valeric acid), poly(lactide-co-caprolactone), PEG-PLGA-PEG and trimethylene carbonate, polyvinylpyrrolidone. The lipid nanoparticle may be coated or associated with a co-polymer such as, but not limited to, a block co-polymer (such as a branched polyether-polyamide block copolymer described in International Publication No. WO2013012476, herein incorporated by reference in its 15 entirety), and (poly(ethylene glycol))-(poly(propylene oxide))-(poly(ethylene glycol)) triblock copolymer (see e.g., U.S. Publication 20120121718 and U.S. Publication 20100003337 and U.S. Pat. No. 8,263,665, the contents of each of which is herein incorporated by reference in their 20 entirety). The co-polymer may be a polymer that is generally regarded as safe (GRAS) and the formation of the lipid nanoparticle may be in such a way that no new chemical entities are created. For example, the lipid nanoparticle may comprise poloxamers coating PLGA nanoparticles without 25 forming new chemical entities which are still able to rapidly penetrate human mucus (Yang et al. Angew. Chem. Int. Ed. 2011 50:2597-2600; the contents of which are herein incorporated by reference in their entirety). A non-limiting scalable method to produce nanoparticles which can penetrate 30 human mucus is described by Xu et al. (see, e.g., J Control Release 2013, 170(2):279-86; the contents of which are herein incorporated by reference in their entirety).

The vitamin of the polymer-vitamin conjugate may be vitamin E. The vitamin portion of the conjugate may be 35 substituted with other suitable components such as, but not limited to, vitamin A, vitamin E, other vitamins, cholesterol, a hydrophobic moiety, or a hydrophobic component of other surfactants (e.g., sterol chains, fatty acids, hydrocarbon chains and alkylene oxide chains).

The lipid nanoparticle engineered to penetrate mucus may include surface altering agents such as, but not limited to, polynucleotides, anionic proteins (e.g., bovine serum albumin), surfactants (e.g., cationic surfactants such as for example dimethyldioctadecylammonium bromide), sugars 45 or sugar derivatives (e.g., cyclodextrin), nucleic acids, polymers (e.g., heparin, polyethylene glycol and poloxamer), mucolytic agents (e.g., N-acetylcysteine, mugwort, bromelain, papain, clerodendrum, acetylcysteine, bromhexine, carbocisteine, eprazinone, mesna, ambroxol, sobrerol, domi- 50 odol, letosteine, stepronin, tiopronin, gelsolin, thymosin β4 dornase alfa, neltenexine, erdosteine) and various DNases including rhDNase. The surface altering agent may be embedded or enmeshed in the particle's surface or disposed (e.g., by coating, adsorption, covalent linkage, or other 55 process) on the surface of the lipid nanoparticle. (see e.g., U.S. Publication 20100215580 and U.S. Publication 20080166414 and US20130164343; the contents of each of which are herein incorporated by reference in their entirety).

In some embodiments, the mucus penetrating lipid nanoparticles may comprise at least one polynucleotide described herein. The polynucleotide may be encapsulated in the lipid nanoparticle and/or disposed on the surface of the particle. The polynucleotide may be covalently coupled to the lipid nanoparticle. Formulations of mucus penetrating 65 lipid nanoparticles may comprise a plurality of nanoparticles. Further, the formulations may contain particles which

may interact with the mucus and alter the structural and/or adhesive properties of the surrounding mucus to decrease mucoadhesion, which may increase the delivery of the mucus penetrating lipid nanoparticles to the mucosal tissue.

In some embodiments, the mucus penetrating lipid nanoparticles may be a hypotonic formulation comprising a mucosal penetration enhancing coating. The formulation may be hypotonice for the epithelium to which it is being delivered. Non-limiting examples of hypotonic formulations may be found in International Patent Publication No. WO2013110028, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, in order to enhance the delivery through the mucosal barrier the RNA (e.g., mRNA) vaccine formulation may comprise or be a hypotonic solution. Hypotonic solutions were found to increase the rate at which mucoinert particles such as, but not limited to, mucuspenetrating particles, were able to reach the vaginal epithelial surface (see e.g., Ensign et al. Biomaterials 2013 34(28): 6922-9, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the RNA (e.g., mRNA) vaccine is formulated as a lipoplex, such as, without limitation, the ATUPLEX™ system, the DACC system, the DBTC system and other siRNA-lipoplex technology from Silence Therapeutics (London, United Kingdom), STEMFECT<sup>TM</sup> from STEMGENT® (Cambridge, Mass.), and polyethylenimine (PEI) or protamine-based targeted and non-targeted delivery of nucleic acids acids (Aleku et al. Cancer Res. 2008 68:9788-9798; Strumberg et al. Int J Clin Pharmacol Ther 2012 50:76-78; Santel et al., Gene Ther 2006 13:1222-1234; Santel et al., Gene Ther 2006 13:1360-1370; Gutbier et al., Pulm Pharmacol. Ther. 2010 23:334-344; Kaufmann et al. Microvasc Res 2010 80:286-293 Weide et al. J Immunother. 2009 32:498-507; Weide et al. J Immunother. 2008 31:180-188; Pascolo Expert Opin. Biol. Ther. 4:1285-1294; Fotin-Mleczek et al., 2011 J. Immunother. 34:1-15; Song et al., Nature Biotechnol. 2005, 23:709-717; Peer et al., Proc Natl Acad Sci USA. 2007 6; 104:4095-4100; deFougerolles Hum 40 Gene Ther. 2008 19:125-132, the contents of each of which are incorporated herein by reference in their entirety).

In some embodiments, such formulations may also be constructed or compositions altered such that they passively or actively are directed to different cell types in vivo, including but not limited to hepatocytes, immune cells, tumor cells, endothelial cells, antigen presenting cells, and leukocytes (Akinc et al. Mol Ther. 2010 18:1357-1364; Song et al., Nat Biotechnol. 2005 23:709-717; Judge et al., J Clin Invest. 2009 119:661-673; Kaufmann et al., Microvasc Res 2010 80:286-293; Santel et al., Gene Ther 2006 13:1222-1234; Santel et al., Gene Ther 2006 13:1360-1370; Gutbier et al., Pulm Pharmacol. Ther. 2010 23:334-344; Basha et al., Mol. Ther. 2011 19:2186-2200; Fenske and Cullis, Expert Opin Drug Deliv. 2008 5:25-44; Peer et al., Science. 2008 319:627-630; Peer and Lieberman, Gene Ther. 2011 18:1127-1133, the contents of each of which are incorporated herein by reference in their entirety). One example of passive targeting of formulations to liver cells includes the DLin-DMA, DLin-KC2-DMA and DLin-MC3-DMA-based lipid nanoparticle formulations, which have been shown to bind to apolipoprotein E and promote binding and uptake of these formulations into hepatocytes in vivo (Akinc et al. Mol Ther. 2010 18:1357-1364, the contents of which are incorporated herein by reference in their entirety). Formulations can also be selectively targeted through expression of different ligands on their surface as exemplified by, but not limited by, folate, transferrin, N-acetylga-

lactosamine (GalNAc), and antibody targeted approaches (Kolhatkar et al., Curr Drug Discov Technol. 2011 8:197-206; Musacchio and Torchilin, Front Biosci. 2011 16:1388-1412; Yu et al., Mol Membr Biol. 2010 27:286-298; Patil et al., Crit Rev Ther Drug Carrier Syst. 2008 25:1-61; Benoit 5 et al., Biomacromolecules. 2011 12:2708-2714; Zhao et al., Expert Opin Drug Deliv. 2008 5:309-319; Akinc et al., Mol Ther. 2010 18:1357-1364; Srinivasan et al., Methods Mol Biol. 2012 820:105-116; Ben-Arie et al., Methods Mol Biol. 2012 757:497-507; Peer 2010 J Control Release. 20:63-68; 10 Peer et al., Proc Natl Acad Sci USA. 2007 104:4095-4100; Kim et al., Methods Mol Biol. 2011 721:339-353; Subramanya et al., Mol Ther. 2010 18:2028-2037; Song et al., Nat Biotechnol. 2005 23:709-717; Peer et al., Science. 2008 319:627-630; Peer and Lieberman, Gene Ther. 2011 15 18:1127-1133, the contents of each of which are incorporated herein by reference in their entirety)

In some embodiments, the RNA (e.g., mRNA) vaccine is formulated as a solid lipid nanoparticle. A solid lipid nanoparticle (SLN) may be spherical with an average diameter 20 between 10 to 1000 nm. SLN possess a solid lipid core matrix that can solubilize lipophilic molecules and may be stabilized with surfactants and/or emulsifiers. In some embodiments, the lipid nanoparticle may be a self-assembly lipid-polymer nanoparticle (see Zhang et al., ACS Nano, 25 2008, 2 (8), pp 1696-1702; the contents of which are herein incorporated by reference in their entirety). As a nonlimiting example, the SLN may be the SLN described in International Patent Publication No. WO2013105101, the contents of which are herein incorporated by reference in 30 their entirety. As another non-limiting example, the SLN may be made by the methods or processes described in International Patent Publication No. WO2013105101, the contents of which are herein incorporated by reference in their entirety.

Liposomes, lipoplexes, or lipid nanoparticles may be used to improve the efficacy of polynucleotides directed protein production as these formulations may be able to increase cell transfection by the RNA (e.g., mRNA) vaccine; and/or increase the translation of encoded protein. One such 40 example involves the use of lipid encapsulation to enable the effective systemic delivery of polyplex plasmid DNA (Heyes et al., Mol Ther. 2007 15: 713-720; the contents of which are incorporated herein by reference in their entirety). The liposomes, lipoplexes, or lipid nanoparticles may also 45 be used to increase the stability of the polynucleotide.

In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure can be formulated for controlled release and/or targeted delivery. As used herein, "controlled release" refers to a pharmaceutical composition or com- 50 pound release profile that conforms to a particular pattern of release to effect a therapeutic outcome. In some embodiments, the RNA (e.g., mRNA) vaccines may be encapsulated into a delivery agent described herein and/or known in the art for controlled release and/or targeted delivery. As 55 used herein, the term "encapsulate" means to enclose, surround or encase. As it relates to the formulation of the compounds of the disclosure, encapsulation may be substantial, complete or partial. The term "substantially encapsulated" means that at least greater than 50, 60, 70, 80, 85, 90, 60 95, 96, 97, 98, 99, 99.9, 99.9 or greater than 99.999% of the pharmaceutical composition or compound of the disclosure may be enclosed, surrounded or encased within the delivery agent. "Partially encapsulation" means that less than 10, 10, 20, 30, 40 50 or less of the pharmaceutical composition or 65 compound of the disclosure may be enclosed, surrounded or encased within the delivery agent. Advantageously, encap-

sulation may be determined by measuring the escape or the activity of the pharmaceutical composition or compound of the disclosure using fluorescence and/or electron micrograph. For example, at least 1, 5, 10, 20, 30, 40, 50, 60, 70, 80, 85, 90, 95, 96, 97, 98, 99, 99.9, 99.99 or greater than 99.99% of the pharmaceutical composition or compound of the disclosure are encapsulated in the delivery agent.

In some embodiments, the controlled release formulation may include, but is not limited to, tri-block co-polymers. As a non-limiting example, the formulation may include two different types of tri-block co-polymers (International Pub. No. WO2012131104 and

WO2012131106, the contents of each of which are incorporated herein by reference in their entirety).

In some embodiments, the RNA (e.g., mRNA) vaccines may be encapsulated into a lipid nanoparticle or a rapidly eliminated lipid nanoparticle and the lipid nanoparticles or a rapidly eliminated lipid nanoparticle may then be encapsulated into a polymer, hydrogel and/or surgical sealant described herein and/or known in the art. As a non-limiting example, the polymer, hydrogel or surgical sealant may be PLGA, ethylene vinyl acetate (EVAc), poloxamer, GELSITE® (Nanotherapeutics, Inc. Alachua, Fla.), HYLENEX® (Halozyme Therapeutics, San Diego Calif.), surgical sealants such as fibrinogen polymers (Ethicon Inc. Cornelia, Ga.), TISSELL® (Baxter International, Inc Deerfield, Ill.), PEG-based sealants, and COSEAL® (Baxter International, Inc Deerfield, Ill.).

In some embodiments, the lipid nanoparticle may be encapsulated into any polymer known in the art which may form a gel when injected into a subject. As another non-limiting example, the lipid nanoparticle may be encapsulated into a polymer matrix which may be biodegradable.

In some embodiments, the RNA (e.g., mRNA) vaccine formulation for controlled release and/or targeted delivery may also include at least one controlled release coating. Controlled release coatings include, but are not limited to, OPADRY®, polyvinylpyrrolidone/vinyl acetate copolymer, polyvinylpyrrolidone, hydroxypropyl methylcellulose, hydroxypropyl cellulose, hydroxyethyl cellulose, EUDRAGIT RL®, EUDRAGIT RS® and cellulose derivatives such as ethylcellulose aqueous dispersions (AQUACOAT® and SURELEASE®).

which are incorporated herein by reference in their entirety). The liposomes, lipoplexes, or lipid nanoparticles may also be used to increase the stability of the polynucleotide. In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure can be formulated for controlled release and/or targeted delivery. As used herein, "controlled release and/or targeted delivery. As used herein, "controlled release and/or targeted delivery formulation may comprise at least one degradable polyesters include, but are not limited to, poly(serine ester), poly(L-lactide-co-L-lysine), poly(4-hydroxy-L-proline ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccine controlled release and/or targeted delivery formulation may contain polycationic side chains. Degradable polyesters include, but are not limited to, poly(serine ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccine controlled release and/or targeted delivery formulation may contain polycationic side chains. Degradable polyesters include, but are not limited to, poly(serine ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccine controlled release and/or targeted delivery formulation may contain polycationic side chains. Degradable polyesters include, but are not limited to, poly(serine ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccines of include, but are not limited to, poly(serine ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccines of controlled release and/or targeted delivery formulation may contain polycationic side chains. Degradable polyesters include, but are not limited to, poly(serine ester), and combinations thereof. In some embodiments, the RNA (e.g., mRNA) vaccines controlled release and/or targeted delivery formulation may contain polycationic side chains.

In some embodiments, the RNA (e.g., mRNA) vaccine controlled release and/or targeted delivery formulation comprising at least one polynucleotide may comprise at least one PEG and/or PEG related polymer derivatives as described in U.S. Pat. No. 8,404,222, the contents of which are incorporated herein by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccine controlled release delivery formulation comprising at least one polynucleotide may be the controlled release polymer system described in US20130130348, the contents of which are incorporated herein by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure may be encapsulated in a therapeutic nanoparticle, referred to herein as "therapeutic nanoparticle RNA (e.g., mRNA) vaccines." Therapeutic nanoparticles

may be formulated by methods described herein and known in the art such as, but not limited to, International Pub Nos. WO2010005740, WO2010030763, WO2010005721, WO2010005723, WO2012054923, U.S. Publication Nos. US20110262491. US20100104645, US20100087337, 5 US20100068285, US20110274759, US20100068286, US20120288541, US20130123351 and US20130230567 and U.S. Pat. Nos. 8,206,747, 8,293,276, 8,318,208 and 8,318,211; the contents of each of which are herein incorporated by reference in their entirety. In some embodiments, therapeutic polymer nanoparticles may be identified by the methods described in US Pub No. US20120140790, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the therapeutic nanoparticle RNA 15 (e.g., mRNA) vaccine may be formulated for sustained release. As used herein, "sustained release" refers to a pharmaceutical composition or compound that conforms to a release rate over a specific period of time. The period of time may include, but is not limited to, hours, days, weeks, 20 months and years. As a non-limiting example, the sustained release nanoparticle may comprise a polymer and a therapeutic agent such as, but not limited to, the polynucleotides of the present disclosure (see International Pub No. 2010075072 and US Pub No. US20100216804, 25 US20110217377 and US20120201859, the contents of each of which are incorporated herein by reference in their entirety). In another non-limiting example, the sustained release formulation may comprise agents which permit persistent bioavailability such as, but not limited to, crystals, 30 macromolecular gels and/or particulate suspensions (see U.S. Patent Publication No US20130150295, the contents of each of which are incorporated herein by reference in their

In some embodiments, the therapeutic nanoparticle RNA 35 (e.g., mRNA) vaccines may be formulated to be target specific. As a non-limiting example, the therapeutic nanoparticles may include a corticosteroid (see International Pub. No. WO2011084518, the contents of which are incorporated herein by reference in their entirety). As a non-limiting example, the therapeutic nanoparticles may be formulated in nanoparticles described in International Pub No. WO2008121949, WO2010005726, WO2010005725, WO2011084521 and US Pub No. US20100069426, US20120004293 and US20100104655, the contents of each 45 of which are incorporated herein by reference in their entirety.

In some embodiments, the nanoparticles of the present disclosure may comprise a polymeric matrix. As a nonlimiting example, the nanoparticle may comprise two or 50 more polymers such as, but not limited to, polyethylenes, polycarbonates, polyanhydrides, polyhydroxyacids, polypropylfumerates, polycaprolactones, polyamides, polyacetals, polyethers, polyesters, poly(orthoesters), polycyanopolyurethanes, 55 acrylates, polyvinyl alcohols, polyacrylates, polyphosphazenes, polymethacrylates, polycyanoacrylates, polyureas, polystyrenes, polyamines, polylysine, poly(ethylene imine), poly(serine ester), poly(Llactide-co-L-lysine), poly(4-hydroxy-L-proline ester) or combinations thereof.

In some embodiments, the therapeutic nanoparticle comprises a diblock copolymer. In some embodiments, the diblock copolymer may include PEG in combination with a polymer such as, but not limited to, polyethylenes, polycarbonates, polyanhydrides, polyhydroxyacids, polypropyl-65 fumerates, polycaprolactones, polyamides, polyacetals, polyethers, polyesters, poly(orthoesters), polycyanoacry-

92

lates, polyvinyl alcohols, polyurethanes, polyphosphazenes, polyacrylates, polymethacrylates, polycyanoacrylates, polyureas, polystyrenes, polyamines, polylysine, poly(ethylene imine), poly(serine ester), poly(L-lactide-co-L-lysine), poly (4-hydroxy-L-proline ester) or combinations thereof. In yet another embodiment, the diblock copolymer may be a high-X diblock copolymer such as those described in International Patent Publication No. WO2013120052, the contents of which are incorporated herein by reference in their entirety.

As a non-limiting example the therapeutic nanoparticle comprises a PLGA-PEG block copolymer (see U.S. Publication No. US20120004293 and U.S. Pat. No. 8,236,330, each of which is herein incorporated by reference in their entirety). In another non-limiting example, the therapeutic nanoparticle is a stealth nanoparticle comprising a diblock copolymer of PEG and PLA or PEG and PLGA (see U.S. Pat. No. 8,246,968 and International Publication No. WO2012166923, the contents of each of which are herein incorporated by reference in their entirety). In yet another non-limiting example, the therapeutic nanoparticle is a stealth nanoparticle or a target-specific stealth nanoparticle as described in U.S. Patent Publication No. US20130172406, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the therapeutic nanoparticle may comprise a multiblock copolymer (see e.g., U.S. Pat. Nos. 8,263,665 and 8,287,910 and U.S. Patent Pub. No. US20130195987, the contents of each of which are herein incorporated by reference in their entirety).

In yet another non-limiting example, the lipid nanoparticle comprises the block copolymer PEG-PLGA-PEG (see e.g., the thermosensitive hydrogel (PEG-PLGA-PEG) was used as a TGF-beta1 gene delivery vehicle in Lee et al. Thermosensitive Hydrogel as a Tgf-β1 Gene Delivery Vehicle Enhances Diabetic Wound Healing. Pharmaceutical Research, 2003 20(12): 1995-2000; as a controlled gene delivery system in Li et al. Controlled Gene Delivery System Based on Thermosensitive Biodegradable Hydrogel. Pharmaceutical Research 2003 20(6):884-888; and Chang et al., Non-ionic amphiphilic biodegradable PEG-PLGA-PEG copolymer enhances gene delivery efficiency in rat skeletal muscle. J Controlled Release. 2007 118:245-253, the contents of each of which are herein incorporated by reference in their entirety). The RNA (e.g., mRNA) vaccines of the present disclosure may be formulated in lipid nanoparticles comprising the PEG-PLGA-PEG block copolymer.

In some embodiments, the therapeutic nanoparticle may comprise a multiblock copolymer (see e.g., U.S. Pat. Nos. 8,263,665 and 8,287,910 and U.S. Patent Pub. No. US20130195987, the contents of each of which are herein incorporated by reference in their entirety).

In some embodiments, the block copolymers described herein may be included in a polyion complex comprising a non-polymeric micelle and the block copolymer. (see e.g., U.S. Publication No. 20120076836, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the therapeutic nanoparticle may comprise at least one acrylic polymer. Acrylic polymers include but are not limited to, acrylic acid, methacrylic acid, acrylic acid and methacrylic acid copolymers, methyl methacrylate copolymers, ethoxyethyl methacrylates, cyanoethyl methacrylate, amino alkyl methacrylate copolymer, poly (acrylic acid), poly(methacrylic acid), polycyanoacrylates and combinations thereof.

In some embodiments, the therapeutic nanoparticles may comprise at least one poly(vinyl ester) polymer. The poly

(vinyl ester) polymer may be a copolymer such as a random copolymer. As a non-limiting example, the random copolymer may have a structure such as those described in International Application No. WO2013032829 or U.S. Patent Publication No US20130121954, the contents of each of 5 which are herein incorporated by reference in their entirety. In some embodiments, the poly(vinyl ester) polymers may be conjugated to the polynucleotides described herein.

In some embodiments, the therapeutic nanoparticle may comprise at least one diblock copolymer. The diblock copo- 10 lymer may be, but it not limited to, a poly(lactic) acid-poly (ethylene)glycol copolymer (see, e.g., International Patent Publication No. WO2013044219, the contents of which are herein incorporated by reference in their entirety). As a non-limiting example, the therapeutic nanoparticle may be 15 used to treat cancer (see International publication No. WO2013044219, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the therapeutic nanoparticles may comprise at least one cationic polymer described herein 20 and/or known in the art.

In some embodiments, the therapeutic nanoparticles may comprise at least one amine-containing polymer such as, but not limited to polylysine, polyethylene imine, poly(amidoamine) dendrimers, poly(beta-amino esters) (see, e.g., U.S. 25 Pat. No. 8,287,849, the contents of which are herein incorporated by reference in their entirety) and combinations thereof.

In some embodiments, the nanoparticles described herein may comprise an amine cationic lipid such as those 30 described in International Patent Application No. WO2013059496, the contents of which are herein incorporated by reference in their entirety. In some embodiments, the cationic lipids may have an amino-amine or an aminoamide moiety.

In some embodiments, the therapeutic nanoparticles may comprise at least one degradable polyester which may contain polycationic side chains. Degradeable polyesters include, but are not limited to, poly(serine ester), poly(Llactide-co-L-lysine), poly(4-hydroxy-L-proline ester), and 40 combinations thereof. In some embodiments, the degradable polyesters may include a PEG conjugation to form a PEGylated polymer.

In some embodiments, the synthetic nanocarriers may contain an immunostimulatory agent to enhance the immune 45 response from delivery of the synthetic nanocarrier. As a non-limiting example, the synthetic nanocarrier may comprise a Th1 immunostimulatory agent, which may enhance a Th1-based response of the immune system (see International Pub No. WO2010123569 and U.S. Publication No. 50 US20110223201, the contents of each of which are herein incorporated by reference in their entirety).

In some embodiments, the synthetic nanocarriers may be formulated for targeted release. In some embodiments, the otides at a specified pH and/or after a desired time interval. As a non-limiting example, the synthetic nanoparticle may be formulated to release the RNA (e.g., mRNA) vaccines after 24 hours and/or at a pH of 4.5 (see International Publication Nos. WO2010138193 and WO2010138194 and 60 US Pub Nos. US20110020388 and US20110027217, each of which is herein incorporated by reference in their entireties).

In some embodiments, the synthetic nanocarriers may be formulated for controlled and/or sustained release of the polynucleotides described herein. As a non-limiting 65 example, the synthetic nanocarriers for sustained release may be formulated by methods known in the art, described

herein and/or as described in International Pub No. WO2010138192 and US Pub No. 20100303850, each of which is herein incorporated by reference in their entirety.

94

In some embodiments, the RNA (e.g., mRNA) vaccine may be formulated for controlled and/or sustained release wherein the formulation comprises at least one polymer that is a crystalline side chain (CYSC) polymer. CYSC polymers are described in U.S. Pat. No. 8,399,007, herein incorporated by reference in its entirety.

In some embodiments, the synthetic nanocarrier may be formulated for use as a vaccine. In some embodiments, the synthetic nanocarrier may encapsulate at least one polynucleotide which encode at least one antigen. As a nonlimiting example, the synthetic nanocarrier may include at least one antigen and an excipient for a vaccine dosage form (see International Publication No. WO2011150264 and U.S. Publication No. US20110293723, the contents of each of which are herein incorporated by reference in their entirety). As another non-limiting example, a vaccine dosage form may include at least two synthetic nanocarriers with the same or different antigens and an excipient (see International Publication No. WO2011150249 and U.S. Publication No. US20110293701, the contents of each of which are herein incorporated by reference in their entirety). The vaccine dosage form may be selected by methods described herein, known in the art and/or described in International Publication No. WO2011150258 and U.S. Publication No. US20120027806, the contents of each of which are herein incorporated by reference in their entirety).

In some embodiments, the synthetic nanocarrier may comprise at least one polynucleotide which encodes at least one adjuvant. As non-limiting example, the adjuvant may comprise dimethyldioctadecylammonium-bromide, dimethyldioctadecylammonium-chloride, dimethyldioctadecylam-35 monium-phosphate or dimethyldioctadecylammonium-acetate (DDA) and an apolar fraction or part of said apolar fraction of a total lipid extract of a mycobacterium (see, e.g., U.S. Pat. No. 8,241,610, the content of which is herein incorporated by reference in its entirety). In some embodiments, the synthetic nanocarrier may comprise at least one polynucleotide and an adjuvant. As a non-limiting example, the synthetic nanocarrier comprising and adjuvant may be formulated by the methods described in International Publication No. WO2011150240 and U.S. Publication No. US20110293700, the contents of each of which are herein incorporated by reference in their entirety.

In some embodiments, the synthetic nanocarrier may encapsulate at least one polynucleotide that encodes a peptide, fragment or region from a virus. As a non-limiting example, the synthetic nanocarrier may include, but is not limited to, any of the nanocarriers described in International Publication No. WO2012024621, WO201202629, WO2012024632 and U.S. Publication No. US20120064110, US20120058153 and US20120058154, the contents of each synthetic nanocarrier is formulated to release the polynucle- 55 of which are herein incorporated by reference in their entirety.

> In some embodiments, the synthetic nanocarrier may be coupled to a polynucleotide which may be able to trigger a humoral and/or cytotoxic T lymphocyte (CTL) response (see, e.g., International Publication No. WO2013019669, the contents of which are herein incorporated by reference in their entirety)

> In some embodiments, the RNA (e.g., mRNA) vaccine may be encapsulated in, linked to and/or associated with zwitterionic lipids. Non-limiting examples of zwitterionic lipids and methods of using zwitterionic lipids are described in U.S. Patent Publication No. US20130216607, the con-

tents of which are herein incorporated by reference in their entirety. In some aspects, the zwitterionic lipids may be used in the liposomes and lipid nanoparticles described herein.

In some embodiments, the RNA (e.g., mRNA) vaccine may be formulated in colloid nanocarriers as described in 5 U.S. Patent Publication No. US20130197100, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the nanoparticle may be optimized for oral administration. The nanoparticle may comprise at 10 least one cationic biopolymer such as, but not limited to, chitosan or a derivative thereof. As a non-limiting example, the nanoparticle may be formulated by the methods described in U.S. Publication No. 20120282343, the contents of which are herein incorporated by reference in their 15 entirety.

In some embodiments, LNPs comprise the lipid KL52 (an amino-lipid disclosed in U.S. Application Publication No. 2012/0295832, the contents of which are herein incorporated by reference in their entirety. Activity and/or safety (as 20 measured by examining one or more of ALT/AST, white blood cell count and cytokine induction, for example) of LNP administration may be improved by incorporation of such lipids. LNPs comprising KL52 may be administered intravenously and/or in one or more doses. In some embodiments, administration of LNPs comprising KL52 results in equal or improved mRNA and/or protein expression as compared to LNPs comprising MC3.

In some embodiments, RNA (e.g., mRNA) vaccine may be delivered using smaller LNPs. Such particles may com- 30 prise a diameter from below 0.1 um up to 100 nm such as, but not limited to, less than 0.1 um, less than 1.0 um, less than 5 um, less than 10 um, less than 15 um, less than 20 um, less than 25 um, less than 30 um, less than 35 um, less than 40 um, less than 50 um, less than 55 um, less than 60 um, 35 less than 65 um, less than 70 um, less than 75 um, less than 80 um, less than 85 um, less than 90 um, less than 95 um, less than 100 um, less than 125 um, less than 150 um, less than 175 um, less than 200 um, less than 225 um, less than 250 um, less than 275 um, less than 300 um, less than 325 40 um, less than 350 um, less than 375 um, less than 400 um, less than 425 um, less than 450 um, less than 475 um, less than 500 um, less than 525 um, less than 550 um, less than 575 um, less than 600 um, less than 625 um, less than 650 um, less than 675 um, less than 700 um, less than 725 um, 45 less than 750 um, less than 775 um, less than 800 um, less than 825 um, less than 850 um, less than 875 um, less than 900 um, less than 925 um, less than 950 um, less than 975 um, or less than 1000 um.

In some embodiments, RNA (e.g., mRNA) vaccines may 50 be delivered using smaller LNPs, which may comprise a diameter from about 1 nm to about 100 nm, from about 1 nm to about 10 nm, about 1 nm to about 20 nm, from about 1 nm to about 30 nm, from about 1 nm to about 40 nm, from about 1 nm to about 50 nm, from about 1 nm to about 60 nm, 55 from about 1 nm to about 70 nm, from about 1 nm to about 80 nm, from about 1 nm to about 90 nm, from about 5 nm to about from 100 nm, from about 5 nm to about 10 nm, about 5 nm to about 20 nm, from about 5 nm to about 30 nm, from about 5 nm to about 40 nm, from about 5 nm to about 60 50 nm, from about 5 nm to about 60 nm, from about 5 nm to about 70 nm, from about 5 nm to about 80 nm, from about 5 nm to about 90 nm, about 10 to about 50 nm, from about 20 to about 50 nm, from about 30 to about 50 nm, from about 40 to about 50 nm, from about 20 to about 60 nm, from about 65 30 to about 60 nm, from about 40 to about 60 nm, from about 20 to about 70 nm, from about 30 to about 70 nm, from about

96

40 to about 70 nm, from about 50 to about 70 nm, from about 60 to about 70 nm, from about 20 to about 80 nm, from about 30 to about 80 nm, from about 40 to about 80 nm, from about 50 to about 80 nm, from about 60 to about 80 nm, from about 20 to about 90 nm, from about 30 to about 90 nm, from about 40 to about 90 nm, from about 60 to about 90 nm, from about 50 to about 90 nm, from about 60 to about 90 nm and/or from about 70 to about 90 nm.

In some embodiments, such LNPs are synthesized using methods comprising microfluidic mixers. Examples of microfluidic mixers may include, but are not limited to, a slit interdigital micromixer including, but not limited to those manufactured by Microinnova (Allerheiligen bei Wildon, Austria) and/or a staggered herringbone micromixer (SHM) (Zhigaltsev, I. V. et al., Bottom-up design and synthesis of limit size lipid nanoparticle systems with aqueous and triglyceride cores using millisecond microfluidic mixing have been published (Langmuir. 2012. 28:3633-40; Belliveau, N. M. et al., Microfluidic synthesis of highly potent limit-size lipid nanoparticles for in vivo delivery of siRNA. Molecular Therapy-Nucleic Acids. 2012. 1:e37; Chen. D. et al., Rapid discovery of potent siRNA-containing lipid nanoparticles enabled by controlled microfluidic formulation. J Am Chem Soc. 2012. 134(16):6948-51, the contents of each of which are herein incorporated by reference in their entirety). In some embodiments, methods of LNP generation comprising SHM, further comprise the mixing of at least two input streams wherein mixing occurs by microstructureinduced chaotic advection (MICA). According to this method, fluid streams flow through channels present in a herringbone pattern causing rotational flow and folding the fluids around each other. This method may also comprise a surface for fluid mixing wherein the surface changes orientations during fluid cycling. Methods of generating LNPs using SHM include those disclosed in U.S. Application Publication Nos. 2004/0262223 and 2012/0276209, the contents of each of which are herein incorporated by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccine of the present disclosure may be formulated in lipid nanoparticles created using a micromixer such as, but not limited to, a Slit Interdigital Microstructured Mixer (SIMM-V2) or a Standard Slit Interdigital Micro Mixer (SSIMM) or Caterpillar (CPMM) or Impinging-jet (IJMM) from the Institut für Mikrotechnik Mainz GmbH, Mainz Germany).

In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure may be formulated in lipid nanoparticles created using microfluidic technology (see, e.g., Whitesides, George M. The Origins and the Future of Microfluidics. Nature, 2006 442: 368-373; and Abraham et al. Chaotic Mixer for Microchannels. Science, 2002 295: 647-651; each of which is herein incorporated by reference in its entirety). As a non-limiting example, controlled microfluidic formulation includes a passive method for mixing streams of steady pressure-driven flows in micro channels at a low Reynolds number (see, e.g., Abraham et al. Chaotic Mixer for Microchannels. Science, 2002 295: 647-651, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the RNA (e.g., mRNA) vaccines of the present disclosure may be formulated in lipid nanoparticles created using a micromixer chip such as, but not limited to, those from Harvard Apparatus (Holliston, Mass.) or Dolomite Microfluidics (Royston, UK). A micromixer chip can be used for rapid mixing of two or more fluid streams with a split and recombine mechanism.

In some embodiments, the RNA (e.g., mRNA) vaccines of the disclosure may be formulated for delivery using the drug

encapsulating microspheres described in International Patent Publication No. WO2013063468 or U.S. Pat. No. 8,440, 614, the contents of each of which are herein incorporated by reference in their entirety. The microspheres may comprise a compound of the formula (I), (II), (III), (IV), (V) or 5 (VI) as described in International Patent Publication No. WO2013063468, the contents of which are herein incorporated by reference in their entirety. In some embodiments, the amino acid, peptide, polypeptide, lipids (APPL) are useful in delivering the RNA (e.g., mRNA) vaccines of the 10 disclosure to cells (see International Patent Publication No. WO2013063468, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the RNA (e.g., mRNA) vaccines of the disclosure may be formulated in lipid nanoparticles 15 having a diameter from about 10 to about 100 nm such as, but not limited to, about 10 to about 20 nm, about 10 to about 30 nm, about 10 to about 40 nm, about 10 to about 50 nm, about 10 to about 60 nm, about 10 to about 70 nm, about 10 to about 80 nm, about 10 to about 90 nm, about 20 to about 20 30 nm, about 20 to about 40 nm, about 20 to about 50 nm, about 20 to about 60 nm, about 20 to about 70 nm, about 20 to about 80 nm, about 20 to about 90 nm, about 20 to about 100 nm, about 30 to about 40 nm, about 30 to about 50 nm. about 30 to about 60 nm, about 30 to about 70 nm, about 30 25 to about 80 nm, about 30 to about 90 nm, about 30 to about 100 nm, about 40 to about 50 nm, about 40 to about 60 nm, about 40 to about 70 nm, about 40 to about 80 nm, about 40 to about 90 nm, about 40 to about 100 nm, about 50 to about 60 nm, about 50 to about 70 nm about 50 to about 80 nm, 30 about 50 to about 90 nm, about 50 to about 100 nm, about 60 to about 70 nm, about 60 to about 80 nm, about 60 to about 90 nm, about 60 to about 100 nm, about 70 to about 80 nm, about 70 to about 90 nm, about 70 to about 100 nm, about 80 to about 90 nm, about 80 to about 100 nm and/or 35 about 90 to about 100 nm.

In some embodiments, the lipid nanoparticles may have a diameter from about 10 to 500 nm.

In some embodiments, the lipid nanoparticle may have a diameter greater than 100 nm, greater than 150 nm, greater 40 than 200 nm, greater than 250 nm, greater than 300 nm, greater than 350 nm, greater than 400 nm, greater than 450 nm, greater than 500 nm, greater than 550 nm, greater than 600 nm, greater than 650 nm, greater than 700 nm, greater than 750 nm, greater than 800 nm, greater than 850 nm, 45 greater than 900 nm, greater than 950 nm or greater than 1000 nm.

In some embodiments, the lipid nanoparticle may be a limit size lipid nanoparticle described in International Patent Publication No. WO2013059922, the contents of which are 50 herein incorporated by reference in their entirety. The limit size lipid nanoparticle may comprise a lipid bilayer surrounding an aqueous core or a hydrophobic core; where the lipid bilayer may comprise a phospholipid such as, but not limited to, diacylphosphatidylcholine, a diacylphosphatidylethanolamine, a ceramide, a sphingomyelin, a dihydrosphingomyelin, a cephalin, a cerebroside, a C8-C20 fatty acid diacylphophatidylcholine, and 1-palmitoyl-2-oleoyl phosphatidylcholine (POPC). In some embodiments, the limit size lipid nanoparticle may comprise a polyethylene 60 glycol-lipid such as, but not limited to, DLPE-PEG, DMPE-PEG, DPPC-PEG and DSPE-PEG.

In some embodiments, the RNA (e.g., mRNA) vaccines may be delivered, localized and/or concentrated in a specific location using the delivery methods described in International Patent Publication No. WO2013063530, the contents of which are herein incorporated by reference in their

entirety. As a non-limiting example, a subject may be administered an empty polymeric particle prior to, simultaneously with or after delivering the RNA (e.g., mRNA) vaccines to the subject. The empty polymeric particle undergoes a change in volume once in contact with the subject and becomes lodged, embedded, immobilized or entrapped at a specific location in the subject.

98

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in an active substance release system (see, e.g., U.S. Patent Publication No. US20130102545, the contents of which are herein incorporated by reference in their entirety). The active substance release system may comprise 1) at least one nanoparticle bonded to an oligonucleotide inhibitor strand which is hybridized with a catalytically active nucleic acid and 2) a compound bonded to at least one substrate molecule bonded to a therapeutically active substance (e.g., polynucleotides described herein), where the therapeutically active substance is released by the cleavage of the substrate molecule by the catalytically active nucleic acid.

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in a nanoparticle comprising an inner core comprising a non-cellular material and an outer surface comprising a cellular membrane. The cellular membrane may be derived from a cell or a membrane derived from a virus. As a non-limiting example, the nanoparticle may be made by the methods described in International Patent Publication No. WO2013052167, the contents of which are herein incorporated by reference in their entirety. As another non-limiting example, the nanoparticle described in International Patent Publication No. WO2013052167, the contents of which are herein incorporated by reference in their entirety, may be used to deliver the RNA (e.g., mRNA) vaccines described herein.

In some embodiments, the RNA (e.g., mRNA) vaccines may be formulated in porous nanoparticle-supported lipid bilayers (protocells). Protocells are described in International Patent Publication No. WO2013056132, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccines described herein may be formulated in polymeric nanoparticles as described in or made by the methods described in U.S. Pat. Nos. 8,420,123 and 8,518,963 and European Patent No. EP2073848B1, the contents of each of which are herein incorporated by reference in their entirety. As a non-limiting example, the polymeric nanoparticle may have a high glass transition temperature such as the nanoparticles described in or nanoparticles made by the methods described in U.S. Pat. No. 8,518,963, the contents of which are herein incorporated by reference in their entirety. As another non-limiting example, the polymer nanoparticle for oral and parenteral formulations may be made by the methods described in European Patent No. EP2073848B1, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the RNA (e.g., mRNA) vaccines described herein may be formulated in nanoparticles used in imaging. The nanoparticles may be liposome nanoparticles such as those described in U.S. Patent Publication No US20130129636, herein incorporated by reference in its entirety. As a non-limiting example, the liposome may comprise gadolinium(III)2-{4,7-bis-carboxymethyl-10-[(N, N-distearylamidomethyl-N'-amido-methyl]-1,4,7,10-tetra-azacyclododec-1-yl}-acetic acid and a neutral, fully saturated phospholipid component (see, e.g., U.S. Patent Publication No US20130129636, the contents of which are herein incorporated by reference in their entirety).

In some embodiments, the nanoparticles which may be used in the present disclosure are formed by the methods described in U.S. Patent Application No. US20130130348, the contents of which are herein incorporated by reference in their entirety.

The nanoparticles of the present disclosure may further include nutrients such as, but not limited to, those which deficiencies can lead to health hazards from anemia to neural tube defects (see, e.g., the nanoparticles described in International Patent Publication No WO2013072929, the contents of which are herein incorporated by reference in their entirety). As a non-limiting example, the nutrient may be iron in the form of ferrous, ferric salts or elemental iron, iodine, folic acid, vitamins or micronutrients.

In some embodiments, the RNA (e.g., mRNA) vaccines of 15 the present disclosure may be formulated in a swellable nanoparticle. The swellable nanoparticle may be, but is not limited to, those described in U.S. Pat. No. 8,440,231, the contents of which are herein incorporated by reference in their entirety. As a non-limiting embodiment, the swellable 20 nanoparticle may be used for delivery of the RNA (e.g., mRNA) vaccines of the present disclosure to the pulmonary system (see, e.g., U.S. Pat. No. 8,440,231, the contents of which are herein incorporated by reference in their entirety).

The RNA (e.g., mRNA) vaccines of the present disclosure 25 may be formulated in polyanhydride nanoparticles such as, but not limited to, those described in U.S. Pat. No. 8,449, 916, the contents of which are herein incorporated by reference in their entirety.

The nanoparticles and microparticles of the present dis- 30 closure may be geometrically engineered to modulate macrophage and/or the immune response. In some embodiments, the geometrically engineered particles may have varied shapes, sizes and/or surface charges in order to incorporated the polynucleotides of the present disclosure for targeted 35 delivery such as, but not limited to, pulmonary delivery (see, e.g., International Publication No WO2013082111, the contents of which are herein incorporated by reference in their entirety). Other physical features the geometrically engineering particles may have include, but are not limited to, 40 fenestrations, angled arms, asymmetry and surface roughness, charge which can alter the interactions with cells and tissues. As a non-limiting example, nanoparticles of the present disclosure may be made by the methods described in International Publication No WO2013082111, the contents 45 of which are herein incorporated by reference in their entirety.

In some embodiments, the nanoparticles of the present disclosure may be water soluble nanoparticles such as, but not limited to, those described in International Publication 50 No. WO2013090601, the contents of which are herein incorporated by reference in their entirety. The nanoparticles may be inorganic nanoparticles which have a compact and zwitterionic ligand in order to exhibit good water solubility. The nanoparticles may also have small hydrodynamic diameters (HD), stability with respect to time, pH, and salinity and a low level of non-specific protein binding.

In some embodiments the nanoparticles of the present disclosure may be developed by the methods described in U.S. Patent Publication No. US20130172406, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the nanoparticles of the present disclosure are stealth nanoparticles or target-specific stealth nanoparticles such as, but not limited to, those described in 65 U.S. Patent Publication No. US20130172406, the contents of which are herein incorporated by reference in their

100

entirety. The nanoparticles of the present disclosure may be made by the methods described in U.S. Patent Publication No. US20130172406, the contents of which are herein incorporated by reference in their entirety.

In some embodiments, the stealth or target-specific stealth nanoparticles may comprise a polymeric matrix. The polymeric matrix may comprise two or more polymers such as, but not limited to, polyethylenes, polycarbonates, polyanhydrides, polyhydroxyacids, polypropylfumerates, polycaprolactones, polyamides, polyacetals, polyethers, polyesters, poly(orthoesters), polycyanoacrylates, polyvinyl alcohols, polyurethanes, polyphosphazenes, polyacrylates, polymethacrylates, polycyanoacrylates, polyureas, polystyrenes, polyamines, polyesters, polyamydrides, polyethers, polyurethanes, polymethacrylates, polyacrylates, polycyanoacrylates or combinations thereof.

In some embodiments, the nanoparticle may be a nanoparticle-nucleic acid hybrid structure having a high density nucleic acid layer. As a non-limiting example, the nanoparticle-nucleic acid hybrid structure may made by the methods described in U.S. Patent Publication No. US20130171646, the contents of which are herein incorporated by reference in their entirety. The nanoparticle may comprise a nucleic acid such as, but not limited to, polynucleotides described herein and/or known in the art.

At least one of the nanoparticles of the present disclosure may be embedded in in the core a nanostructure or coated with a low density porous 3-D structure or coating which is capable of carrying or associating with at least one payload within or on the surface of the nanostructure. Non-limiting examples of the nanostructures comprising at least one nanoparticle are described in International Patent Publication No. WO2013123523, the contents of which are herein incorporated by reference in their entirety.

In some embodiments the RNA (e.g., mRNA) vaccine may be associated with a cationic or polycationic compounds, including protamine, nucleoline, spermine or spermidine, or other cationic peptides or proteins, such as poly-L-lysine (PLL), polyarginine, basic polypeptides, cell penetrating peptides (CPPs), including HIV-binding peptides, HIV-1 Tat (HIV), Tat-derived peptides, Penetratin, VP<sup>22</sup> derived or analog peptides, Pestivirus Ems, HSV, VP<sup>22</sup> (Herpes simplex), MAP, KALA or protein transduction domains (PTDs), PpT620, prolin-rich peptides, arginine-rich peptides, lysine-rich peptides, MPG-peptide(s), Pep-1, L-oligomers, Calcitonin peptide(s), Antennapedia-derived peptides (particularly from *Drosophila* antennapedia), pAntp, plsl, FGF, Lactoferrin, Transportan, Buforin-2, Bac715-24, SynB, SynB(1), pVEC, hCT-derived peptides, SAP, histones, cationic polysaccharides, for example chitosan, polybrene, cationic polymers, e.g. polyethyleneimine (PEI), cationic lipids, e.g. DOTMA: [1-(2,3-sioleyloxy) propyl)]-N,N,N-trimethylammonium chloride, DMRIE, di-C14-amidine, DOTIM, SAINT, DC-Chol, BGTC, CTAP, DOPC, DODAP, DOPE: Dioleyl phosphatidylethanolamine, DOSPA, DODAB, DOIC, DMEPC, DOGS: Dioctadecylamidoglicylspermin, DIMRI: Dimyristooxypropyl dimethyl hydroxyethyl ammonium bromide, DOTAP: dioleoyloxy-3-(trimethylammonio)propane, DC-6-14: O,Oditetradecanoyl-N-.alpha.-trimethylammonioacetyl)diethanolamine chloride, CLIP 1: rac-[(2,3-dioctadecyloxypropyl) (2-hydroxyethyl)]-dimethylammonium chloride, CLIP6: rac-[2(2,3-dihexadecyloxypropyloxymethyloxy)ethyl]trimethylammonium, CLIP9: rac-[2(2,3-dihexadecyloxypropyloxysuccinyloxy)ethyl]-trimethylammonium, oligofectamine, or cationic or polycationic polymers, e.g.

modified polyaminoacids, such as beta-aminoacid-polymers

or reversed polyamides, etc., modified polyethylenes, such as PVP (poly(N-ethyl-4-vinylpyridinium bromide)), etc., modified acrylates, such as pDMAEMA (poly(dimethylaminoethyl methylacrylate)), etc., modified amidoamines such as pAMAM (poly(amidoamine)), etc., modified polybetaminoester (PBAE), such as diamine end modified 1.4 butanediol diacrylate-co-5-amino-1-pentanol polymers, etc., dendrimers, such as polypropylamine dendrimers or pAMAM based dendrimers, etc., polyimine(s), such as PEI: poly (ethyleneimine), poly(propyleneimine), etc., polyallylamine, sugar backbone based polymers, such as cyclodextrin based polymers, dextran based polymers, chitosan, etc., silan backbone based polymers, such as PMOXA-PDMS copolymers, etc., blockpolymers consisting of a combination of one or more cationic blocks (e.g. selected from a cationic polymer as mentioned above) and of one or more hydrophilic or hydrophobic blocks (e.g. polyethyleneglycole), etc.

In other embodiments the RNA (e.g., mRNA) vaccine is 20 not associated with a cationic or polycationic compounds.

In some embodiments, a nanoparticle comprises compounds of Formula (I):

$$R_4$$
 $N$ 
 $R_1$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 

or a salt or isomer thereof, wherein:

 $R_1$  is selected from the group consisting of  $C_{\text{5-30}}$  alkyl,  $C_{\text{5-20}}$  alkenyl, —R\*YR", —YR", and —R"M'R';

 $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl,  $C_{2-14}$  alkenyl, -R\*YR", -YR", and -R\*OR", or  $R_2$  and  $R_3$ , together with the atom to which they are attached, form a heterocycle or carbocycle;

 $R_4$  is selected from the group consisting of a  $C_{3-6}$  carbocycle, — $(CH_2)_nQ$ , — $(CH_2)_nCHQR$ , —CHQR, — $CQ(R)_2$ , and unsubstituted  $C_{1-6}$  alkyl, where Q is selected from a carbocycle, heterocycle, -OR,  $-O(CH_2)_nN(R)_2$ , -C(O)OR, -OC(O)R, -CX<sub>3</sub>, -CX<sub>2</sub>H, -CXH<sub>2</sub>, -CN, -N  $(R)_2$ ,  $-C(O)N(R)_2$ , -N(R)C(O)R,  $-N(R)S(O)_2R$ , -N(R) $C(O)N(R)_2$ ,  $--N(R)C(S)N(R)_2$ ,  $--N(R)R_8$ ,  $--O(CH_2)_nOR$ ,  $-N(R)C(=NR_9)N(R)_2$ ,  $-N(R)C(=CHR_9)N(R)_2$ , -OC $(O)N(R)_2$ , -N(R)C(O)OR, -N(OR)C(O)R, -N(OR)S $(\mathrm{O})_2 \ \mathrm{R}, \ --\mathrm{N}(\mathrm{OR})\mathrm{C}(\mathrm{O})\mathrm{OR}, \ --\mathrm{N}(\mathrm{OR})\mathrm{C}(\mathrm{O})\mathrm{N}(\mathrm{R})_2, \ --\mathrm{N}(\mathrm{OR}) \ _{50}$  $C(S)N(R)_2$  $-N(OR)C(=NR_9)N(R)_2$ -N(OR)C $(=CHR_9)N(R)_2$  $-C(=NR_o)N(R)_2$  $-C(=NR_o)R$ , -C(O)N(R)O R, and  $C(R)N(R)_2C(O)OR$ , and each n is independently selected from 1, 2, 3, 4, and 5;

each  $R_5$  is independently selected from the group consisting of  $C_{1\text{--}3}$  alkyl,  $C_{2\text{--}3}$  alkenyl, and H;

each  $R_6$  is independently selected from the group consisting of  $C_{1,3}$  alkyl,  $C_{2,3}$  alkenyl, and H;

M and M' are independently selected from —C(O)O—, —OC(O)—, —C(O)N(R')—,

-N(R')C(O), -C(O), -C(S), -C(S)S, -SC(S), -C(S)S, -SC(S), -C(S)S, -S(O), -S(O), -S(O), an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1\mbox{-}3}$  alkyl,  $C_{2\mbox{-}3}$  alkenyl, and H;

 $R_8$  is selected from the group consisting of  $C_{3-6}$  carbocycle and heterocycle;

 $R_9$  is selected from the group consisting of H, CN,  $NO_2$ ,  $C_{1-6}$  alkyl, -OR,  $-S(O)_2R$ ,  $-S(O)_2N(R)_2$ ,  $C_{2-6}$  alkenyl,  $C_{3-6}$  carbocycle and heterocycle;

each R is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each R' is independently selected from the group consisting of  $C_{1-18}$  alkyl,  $C_{2-18}$  alkenyl, —R\*YR", —YR", and H; each R" is independently selected from the group consisting of  $C_{3-14}$  alkyl and  $C_{3-14}$  alkenyl;

each R\* is independently selected from the group consisting of  $C_{1-12}$  alkyl and  $C_{2-12}$  alkenyl;

each Y is independently a  $C_{3-6}$  carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13.

In some embodiments, a subset of compounds of Formula (I) includes those in which when  $R_4$  is  $-(CH_2)_nQ$ ,  $-(CH_2)_nCHQR$ , -CHQR, or  $-CQ(R)_2$ , then (i) Q is not  $-N(R)_2$  when n is 1, 2, 3, 4 or 5, or (ii) Q is not 5, 6, or 7-membered heterocycloalkyl when n is 1 or 2.

In some embodiments, another subset of compounds of Formula (I) includes those in which

 $R_1$  is selected from the group consisting of  $C_{5-30}$  alkyl,  $C_{5-20}$  alkenyl, —R\*YR", —YR", and —R"M'R';

 $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl,  $C_{2-14}$  alkenyl,  $-R^*YR^*$ ,  $-YR^*$ , and  $-R^*OR^*$ , or  $R_2$  and  $R_3$ , together with the atom to which they are attached, form a heterocycle or carbocycle;

R<sub>4</sub> is selected from the group consisting of a C<sub>3-6</sub> carbo-30 cycle,  $-(CH_2)_nQ$ ,  $-(CH_2)_nCHQR$ , -CHQR,  $-CQ(R)_2$ , and unsubstituted  $C_{1-6}$  alkyl, where Q is selected from a  $C_{3-6}$ carbocycle, a 5- to 14-membered heteroaryl having one or more heteroatoms selected from N, O, and S, -OR,  $-O(CH_2)_nN(R)_2$ , -C(O)OR, -OC(O)R, 35  $-CX_2H$ ,  $-CXH_2$ , -CN,  $-C(O)N(R)_2$ , -N(R)C(O)R,  $-N(R)S(O)_2R$ ,  $-N(R)C(O)N(R)_2$ ,  $-N(R)C(S)N(R)_2$ ,  $-\text{CRN}(R)_2\text{C}(O)\text{OR}, -\text{N}(R)R_8, -\text{O}(CH_2)_n\text{OR}, -\text{N}(R)\text{C}$  $(=NR_9)N(R)_2$ ,  $-N(R)C(=CHR_9)N(R)_2$ ,  $-OC(O)N(R)_2$ , -N(OR)C(O)R,  $-N(OR)S(O)_2R$ , -N(R)C(O)OR-N(OR)C(O)OR,  $-N(OR)C(O)N(R)_2$ , -N(OR)C(S)N $(R)_2$ ,  $-N(OR)C(=NR_9)N(R)_2$ ,  $-N(OR)C(=CHR_9)N$  $(R)_2$ ,  $-C(=NR_9)N(R)_2$ ,  $-C(=NR_9)R$ , -C(O)N(R)OR, and a 5- to 14-membered heterocycloalkyl having one or more heteroatoms selected from N, O, and S which is substituted with one or more substituents selected from oxo =O), OH, amino, mono- or di-alkylamino, and C<sub>1-3</sub> alkyl, and each n is independently selected from 1, 2, 3, 4, and 5; each R<sub>5</sub> is independently selected from the group consist-

ing of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H; each  $R_6$  is independently selected from the group consist-

ing of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H; M and M' are independently selected from —C(O)O—, —OC(O)—, —C(O)N(R')—, —N(R')C(O)—, —C(O)—, —C(O)—, —C(S)—, —SC(S)—, —CH(OH)—, —P(O) (OR')O—, — $S(O)_2$ —, —S—S—, an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

 $\rm R_8$  is selected from the group consisting of  $\rm C_{3\text{-}6}$  carbo- cycle and heterocycle;

 $R_9$  is selected from the group consisting of H, CN,  $NO_2$ ,  $C_{1-6}$  alkyl, -OR,  $-S(O)_2R$ ,  $-S(O)_2N(R)_2$ ,  $C_{2-6}$  alkenyl,  $C_{3-6}$  carbocycle and heterocycle;

each R is independently selected from the group consist-65 ing of C<sub>1-3</sub> alkyl, C<sub>2-3</sub> alkenyl, and H;

each R' is independently selected from the group consisting of  $C_{1-18}$  alkyl,  $C_{2-18}$  alkenyl, —R\*YR", —YR", and H;

each R" is independently selected from the group consisting of  $\rm C_{3-14}$  alkyl and  $\rm C_{3-14}$  alkenyl;

each  $R^*$  is independently selected from the group consisting of  $C_{1-12}$  alkyl and  $C_{2-12}$  alkenyl;

each Y is independently a C<sub>3-6</sub> carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13, or salts or isomers thereof.

In some embodiments, another subset of compounds of Formula (I) includes those in which

 $R_1$  is selected from the group consisting of  $C_{5-30}$  alkyl,  $C_{5-20}$  alkenyl, -R\*YR", -YR", and -R"M'R';

 $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl,  $C_{2-14}$  alkenyl, — $R^*YR^*$ , — $YR^*$ , and — $R^*OR^*$ , or  $R_2$  and  $R_3$ , together with the atom to which they are attached, form a heterocycle or carbocycle;

 $R_4$  is selected from the group consisting of a  $C_{3-6}$  carbocycle,  $-(CH_2)_nQ$ ,  $-(CH_2)_nCHQR$ , -CHQR,  $-CQ(R)_2$ , 20 and unsubstituted  $C_{1-6}$  alkyl, where Q is selected from a  $C_{3-6}$ carbocycle, a 5- to 14-membered heterocycle having one or more heteroatoms selected from N, O, and S, -OR, —C(O)OR,  $-CX_3$ ,  $-O(CH_2)_nN(R)_2$ --OC(O)R,  $-CX_2H$ ,  $-CXH_2$ , -CN,  $-C(O)N(R)_2$ , -N(R)C(O)R, 25  $-N(R)S(O)_2R$ ,  $-N(R)C(O)N(R)_2$  $-N(R)C(S)N(R)_2$  $-CRN(R)_2C(O)OR$ ,  $-N(R)R_8$ ,  $-O(CH_2)_nOR$ , -N(R)C $(=NR_9)N(R)_2$ ,  $-N(R)C(=CHR_9)N(R)_2$ ,  $-OC(O)N(R)_2$ , -N(R)C(O)OR-N(OR)C(O)R $-N(OR)S(O)_2R$ , -N(OR)C(O)OR, -N(OR)C(O)N(R)<sub>2</sub>, -N(OR)C(S)N-N(OR)C(=CHR<sub>o</sub>)N  $-N(OR)C(=NR_9)N(R)_2$  $(R)_2$ , — $C(=NR_9)R$ , —C(O)N(R)OR,

and  $-C(=NR_9)N(R)_2$ , and each n is independently selected from 1, 2, 3, 4, and 5; and when Q is a 5- to 14-membered heterocycle and (i)  $R_4$  is  $-(CH_2)_nQ$  in which n is 1 or 2, or 35 (ii)  $R_4$  is  $-(CH_2)_nCHQR$  in which n is 1, or (iii)  $R_4$  is  $-(CHQR, and -CQ(R)_2, then Q is either a 5- to 14-membered heteroaryl or 8- to 14-membered heterocycloalkyl;$ 

each  $R_5$  is independently selected from the group consisting of  $C_{1\text{--}3}$  alkyl,  $C_{2\text{--}3}$  alkenyl, and H;

each  $R_6$  is independently selected from the group consisting of  $C_{1\text{--}3}$  alkyl,  $C_{2\text{--}3}$  alkenyl, and H;

M and M' are independently selected from —C(O)O—, —OC(O)—, —C(O)N(R')—, —N(R')C(O)—, —C(O)—, —C(O)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

 $R_8$  is selected from the group consisting of  $C_{3-6}$  carbo- 50 cycle and heterocycle;

 $R_9$  is selected from the group consisting of H, CN, NO<sub>2</sub>,  $C_{1-6}$  alkyl, —OR, —S(O)<sub>2</sub>R, —S(O)<sub>2</sub>N(R)<sub>2</sub>,  $C_{2-6}$  alkenyl,  $C_{3-6}$  carbocycle and heterocycle;

each R is independently selected from the group consist-  $^{55}$  Formula (I) includes those in which ing of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;  $R_1$  is selected from the group consist-  $R_1$  is selected from the group consist-  $R_2$  is selected from the group consist-  $R_3$  is selected from  $R_3$  is selected from the  $R_3$  is the first  $R_3$  is the group

each R' is independently selected from the group consisting of  $C_{1-18}$  alkyl,  $C_{2-18}$  alkenyl, —R\*YR", —YR", and H; each R" is independently selected from the group consisting of  $C_{3-14}$  alkyl and  $C_{3-14}$  alkenyl;

each R\* is independently selected from the group consisting of  $C_{1-12}$  alkyl and  $C_{2-12}$  alkenyl;

each Y is independently a C<sub>3-6</sub> carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13, or salts or isomers thereof.

104

In some embodiments, another subset of compounds of Formula (I) includes those in which

 $R_1$  is selected from the group consisting of  $C_{5-30}$  alkyl,  $C_{5-20}$  alkenyl, -R\*YR", -YR", and -R"M'R';

 $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl,  $C_{2-14}$  alkenyl, —R\*YR", —YR", and —R\*OR", or  $R_2$  and  $R_3$ , together with the atom to which they are attached, form a heterocycle or carbocycle;

 $R_4$  is selected from the group consisting of a  $C_{3-6}$ carbocycle,  $-(CH_2)_nQ$ ,  $-(CH_2)_nCHQR$ , -CHQR,  $(R)_2$ , and unsubstituted  $C_{1-6}$  alkyl, where Q is selected from a C<sub>3-6</sub> carbocycle, a 5- to 14-membered heteroaryl having one or more heteroatoms selected from N, O, and S, —OR,  $-\mathrm{O(CH}_2)_n\mathrm{N(R)}_2,$ --OC(O)R, --C(O)OR,  $-CX_2H$ ,  $-CXH_2$ , -CN,  $-C(O)N(R)_2$ , -N(R)C(O)R,  $-N(R)C(O)N(R)_2$ ,  $-N(R)C(S)N(R)_2$ ,  $-N(R)S(O)_{2}R$  $-CRN(R)_2C(O)OR$ ,  $-N(R)R_8$ ,  $-O(CH_2)_nOR$ , -N(R)C $(=NR_9)N(R)_2$ ,  $-N(R)C(=CHR_9)N(R)_2$ ,  $-OC(O)N(R)_2$ ,  $-N(OR)S(O)_2R$ -N(R)C(O)OR-N(OR)C(O)R, -N(OR)C(O)OR, --N(OR)C(O)N(R)<sub>2</sub>, --N(OR)C(S)N $(R)_2$ ,  $-N(OR)C(=NR_9)N(R)_2$ ,  $-N(OR)C(=CHR_o)N$  $(R)_2$ ,  $-C(=NR_9)R$ , -C(O)N(R)OR, and  $-C(=NR_9)N$  $(R)_2$ , and each n is independently selected from 1, 2, 3, 4, and 5:

each  $R_5$  is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each  $R_6$  is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

M and M' are independently selected from —C(O)O—, —OC(O)—, —C(O)N(R')—, —N(R')C(O)—, —C(O)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, —C(S)—, an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1\text{--}3}$  alkyl,  $C_{2\text{--}3}$  alkenyl, and H;

 $R_8$  is selected from the group consisting of  $C_{3-6}$  carbocycle and heterocycle;

 $R_9$  is selected from the group consisting of H, CN, NO $_2,$   $C_{1\mbox{-}6}$  alkyl, —OR, —S(O) $_2$ R, —S(O) $_2$ N(R) $_2$ ,  $C_{2\mbox{-}6}$  alkenyl,  $_4$ O  $C_{3\mbox{-}6}$  carbocycle and heterocycle;

each R is independently selected from the group consisting of C<sub>1-3</sub> alkyl, C<sub>2-3</sub> alkenyl, and H;

each R' is independently selected from the group consisting of  $C_{1\text{-}18}$  alkyl,  $C_{2\text{-}18}$  alkenyl, —R\*YR", —YR", and H; each R" is independently selected from the group consisting of  $C_{3\text{-}14}$  alkyl and  $C_{3\text{-}14}$  alkenyl;

each  $R^*$  is independently selected from the group consisting of  $C_{1-12}$  alkyl and  $C_{2-12}$  alkenyl;

each Y is independently a C<sub>3-6</sub> carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13, or salts or isomers thereof.

In some embodiments, another subset of compounds of Formula (I) includes those in which

 $R_1$  is selected from the group consisting of  $C_{5-30}$  alkyl,  $C_{5-20}$  alkenyl, -R\*YR", -YR", and -R"M'R';

R<sub>2</sub> and R<sub>3</sub> are independently selected from the group consisting of H, C<sub>2-14</sub> alkyl, C<sub>2-14</sub> alkenyl, —R\*YR", 60 —YR", and —R\*OR", or R<sub>2</sub> and R<sub>3</sub>, together with the atom to which they are attached, form a heterocycle or carbocycle; R<sub>4</sub> is —(CH<sub>2</sub>)<sub>n</sub>Q or —(CH<sub>2</sub>)<sub>n</sub>CHQR, where Q is —N (R)<sub>2</sub>, and n is selected from 3, 4, and 5;

each  $R_5$  is independently selected from the group consist-65 ing of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each  $R_6$  is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

M and M' are independently selected from —C(O)O—, —OC(O)—, —C(O)N(R')—, —N(R')C(O)—, —C(O)—, —C(S)—, —C(S)S—, —SC(S)—, —CH(OH)—, —P(O) (OR')O—, —S(O) $_2$ —, —S—S—, an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each R is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each R' is independently selected from the group consisting of  $C_{1-18}$  alkyl,  $C_{2-18}$  alkenyl, —R\*YR", —YR", and H; each R" is independently selected from the group consisting of  $C_{3-14}$  alkyl and  $C_{3-14}$  alkenyl;

each  $R^*$  is independently selected from the group consisting of  $C_{1\text{-}12}$  alkyl and  $C_{1\text{-}12}$  alkenyl;

each Y is independently a C<sub>3-6</sub> carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13, or salts or isomers thereof.

In some embodiments, another subset of compounds of Formula (I) includes those in which

 $R_1$  is selected from the group consisting of  $C_{5-30}$  alkyl,  $C_5$ -20 alkenyl, -R\*YR", -YR", and -R"M'R';

 $R_2$  and  $R_3$  are independently selected from the group consisting of  $C_{1-14}$  alkyl,  $C_{2-14}$  alkenyl, —R\*YR", —YR", and —R\*OR", or  $R_2$  and  $R_3$ , together with the atom to which they are attached, form a heterocycle or carbocycle;

R<sub>4</sub> is selected from the group consisting of —(CH<sub>2</sub>)<sub>n</sub>Q, —(CH<sub>2</sub>)<sub>n</sub>CHQR, —CHQR, and —CQ(R)<sub>2</sub>, where Q is —N(R)<sub>2</sub>, and n is selected from 1, 2, 3, 4, and 5;

each  $R_5$  is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each  $R_6$  is independently selected from the group consisting of  $C_{1\text{--}3}$  alkyl,  $C_{2\text{--}3}$  alkenyl, and H;

M and M' are independently selected from -C(O)O, -OC(O), -C(O)N(R'), -N(R')C(O), -C(O), -C(O), -C(S), -C(S)S, -SC(S), -CH(OH), -P(O) -(OR')O,  $-S(O)_2$ , -S, an aryl group, and a heteroaryl group;

 $R_7$  is selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each R is independently selected from the group consisting of  $C_{1-3}$  alkyl,  $C_{2-3}$  alkenyl, and H;

each R' is independently selected from the group consisting of  $C_{1-18}$  alkyl,  $C_{2-18}$  alkenyl, —R\*YR", —YR", and H; each R" is independently selected from the group consisting of  $C_{3-14}$  alkyl and  $C_{3-14}$  alkenyl;

each  $R^*$  is independently selected from the group consisting of  $C_{1-12}$  alkyl and  $C_{1-12}$  alkenyl;

each Y is independently a C<sub>3-6</sub> carbocycle;

each X is independently selected from the group consisting of F, Cl, Br, and I; and

m is selected from 5, 6, 7, 8, 9, 10, 11, 12, and 13, or salts or isomers thereof.

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (IA):

(IA)

$$R_4$$
 $N$ 
 $M_1$ 
 $R_2$ 
 $R_3$ 

or a salt or isomer thereof, wherein 1 is selected from 1, 2, 3, 4, and 5; m is selected from 5, 6, 7, 8, and 9;  $M_1$  is a bond or M';  $R_4$  is unsubstituted  $C_{1-3}$  alkyl, or  $-(CH_2)_nQ$ , in which Q is OH,  $-NHC(S)N(R)_2$ ,  $-NHC(O)N(R)_2$ , -N(R) C(O)R,  $-N(R)S(O)_2R$ ,  $-N(R)R_8$ ,  $-NHC(=NR_9)N(R)_2$ ,  $-NHC(=CHR_9)N(R)_2$ ,  $-OC(O)N(R)_2$ , -N(R)C(O)OR, heteroaryl or heterocycloalkyl; M and M' are independently selected

from —C(O)O—, —OC(O)—, —C(O)N(R')—, —P(O) (OR')O—, —S—S—, an aryl group, and a heteroaryl group; and  $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl, and  $C_{2-14}$  alkenyl.

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (II):

or a salt or isomer thereof, wherein 1 is selected from 1, 2, 3, 4, and 5;  $M_1$  is a bond or M';  $R_4$  is unsubstituted  $C_{1-3}$  alkyl, or  $-(CH_2)_nQ$ , in which n is 2, 3, or 4, and Q is

OH, —NHC(S)N(R)<sub>2</sub>, —NHC(O)N(R)<sub>2</sub>, —N(R)C(O)R, —N(R)S(O)<sub>2</sub>R, —N(R)R<sub>8</sub>, —NHC(=NR<sub>9</sub>)N(R)<sub>2</sub>, —NHC (=CHR<sub>9</sub>)N(R)<sub>2</sub>, —OC(O)N(R)<sub>2</sub>, —N(R)C(O)OR, heteroaryl or heterocycloalkyl; M and M' are independently selected

from -C(O)O—, -OC(O)—, -C(O)N(R')—, -P(O)(OR')O—, -S—S—, an aryl group, and a heteroaryl group; and  $R_2$  and  $R_3$  are independently selected from the group consisting of H,  $C_{1-14}$  alkyl, and  $C_{2-14}$  alkenyl.

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (IIa), (IIb), (IIc), or (IIe):

$$\mathbb{R}_4$$
  $\mathbb{N}$   $\mathbb{N}$ 

(IIe)

or a salt or isomer thereof, wherein R<sub>4</sub> is as described herein

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (IId):

O 
$$R'$$
HO  $R'$ 
 $R'$ 
 $R_5$ 
 $R_6$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_8$ 
 $R_8$ 
 $R_8$ 
 $R_8$ 
 $R_8$ 

or a salt or isomer thereof, wherein n is 2, 3, or 4; and m, R', R", and  $R_2$  through  $R_6$  are as described herein. For example, each of  $R_2$  and  $R_3$  may be independently selected from the group consisting of  $C_{5-14}$  alkyl and  $C_{5-14}$  alkenyl.

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (IIa), (IIb), (IIc), or (IIe):

$$(IIa)$$
 $R_4$ 
 $N$ 
 $S_0$ 
 $S_0$ 
 $S_0$ 
 $S_0$ 
 $S_0$ 

$$\begin{array}{c} (IIb) \\ \\ R_4 \end{array}$$

(IIc)

$$\mathbb{R}_4$$
  $\mathbb{N}$   $\mathbb{N}$ 

or a salt or isomer thereof, wherein  $R_4$  is as described herein.

In some embodiments, a subset of compounds of Formula (I) includes those of Formula (IId):

O 
$$R'$$

HO  $R'$ 
 $R_5$ 
 $R_6$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 
 $R_7$ 

or a salt or isomer thereof, wherein n is 2, 3, or 4; and m, R', R", and  $R_2$  through  $R_6$  are as described herein. For example, each of  $R_2$  and  $R_3$  may be independently selected from the group consisting of  $C_{5-14}$  alkyl and  $C_{5-14}$  alkenyl.

In some embodiments, the compound of Formula (I) is selected from the group consisting of:

40

$$(Compound 37)$$

30

In further embodiments, the compound of Formula (I) is selected from the group consisting of:

In some embodiments, the compound of Formula (I) is selected from the group consisting of:

$$\begin{array}{c} \text{(Compound 71)} \\ \text{N} \\ \text{O} \\ \text{O}$$

$$\begin{array}{c} \text{HO} \\ \text{N} \\ \\ \text{O} \\ \\ \text{O} \\ \end{array}, \\ \begin{array}{c} \text{(Compound 88)} \\ \\ \text{O} \\ \\ \end{array}$$

$$F = \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{j=1}^{N} \sum_{j=1}^{N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sum_{j=1}^{N}$$

HO 
$$\stackrel{\text{N}}{\longrightarrow}$$
  $\stackrel{\text{O}}{\longrightarrow}$   $\stackrel{\text{O}}{\longrightarrow}$ 

$$\begin{array}{c} O \\ O \\ O \\ N \end{array}$$

$$\begin{array}{c} O_2N \\ N \\ H \end{array}$$

$$(Compound 175)$$

$$\begin{array}{c} O \\ N \\ H \end{array}$$

$$\bigcup_{\mathbf{N}} \bigvee_{\mathbf{N}} \bigvee$$

$$\begin{array}{c} NH \\ H_2N \\ \end{array} \\ N \\ O \\ O \\ \end{array} \\ O \\ O \\ O \\ \end{array}$$

$$(Compound 208)$$

$$N$$

$$N$$

$$N$$

$$O$$

$$O$$

$$O$$

$$O$$

$$\begin{array}{c} O_2N \\ N \\ N \\ H \end{array}$$

$$(Compound\ 210)$$

$$\begin{array}{c} H_2N \\ O = S \\ N \\ H \end{array}$$

$$H_2N$$
 (Compound 221)

 $H_2N$  (Compound 221)

In some embodiments, a nanoparticle comprises the following compound:

or salts and isomers thereof.

In some embodiments, the disclosure features a nanoparticle composition including a lipid component comprising a compound as described herein (e.g., a compound according to Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe)).

In some embodiments, the disclosure features a pharmaceutical composition comprising a nanoparticle composition according to the preceding embodiments and a pharmaceutically acceptable carrier. For example, the pharmaceutically acceptable carrier composition is refrigerated or frozen for storage and/or shipment (e.g., being stored at a temperature of 4° C. or lower, such as a temperature between about -150° C. and about 0° C. or between about -80° C. and about -20° C. (e.g., about -5° C., -10° C., -15° C., -20° C., -25° C., -30° C., -40° C., -50° C., -60° C., -70° C., -80° C., -90° C., 35 -130° C. or -150° C.). For example, the pharmaceutical composition is a solution that is refrigerated for storage and/or shipment at, for example, about -20° C., -30° C., -40° C., -50° C., -60° C., -70° C., or -80° C.

In some embodiments, the disclosure provides a method of delivering a therapeutic and/or prophylactic (e.g., RNA, such as mRNA) to a cell (e.g., a mammalian cell). This method includes the step of administering to a subject (e.g., a mammal, such as a human) a nanoparticle composition including (i) a lipid component including a phospholipid 45 (such as a polyunsaturated lipid), a PEG lipid, a structural lipid, and a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (lie) and (ii) a therapeutic and/or prophylactic, in which administering involves contacting the cell with the nanoparticle composition, whereby the therapeutic and/or 50 prophylactic is delivered to the cell.

In some embodiments, the disclosure provides a method of producing a polypeptide of interest in a cell (e.g., a mammalian cell). The method includes the step of contacting the cell with a nanoparticle composition including (i) a 55 lipid component including a phospholipid (such as a polyunsaturated lipid), a PEG lipid, a structural lipid, and a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe) and (ii) an mRNA encoding the polypeptide of interest, whereby the mRNA is capable of being translated 60 in the cell to produce the polypeptide.

In some embodiments, the disclosure provides a method of treating a disease or disorder in a mammal (e.g., a human) in need thereof. The method includes the step of administering to the mammal a therapeutically effective amount of 65 a nanoparticle composition including (i) a lipid component including a phospholipid (such as a polyunsaturated lipid),

a PEG lipid, a structural lipid, and a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe) and (ii) a therapeutic and/or prophylactic (e.g., an mRNA). In some embodiments, the disease or disorder is characterized by dysfunctional or aberrant protein or polypeptide activity. For example, the disease or disorder is selected from the group consisting of rare diseases, infectious diseases, cancer and proliferative diseases, genetic diseases (e.g., cystic fibrosis), autoimmune diseases, diabetes, neurodegenerative diseases, cardio- and reno-vascular diseases, and metabolic diseases.

In some embodiments, the disclosure provides a method of delivering (e.g., specifically delivering) a therapeutic and/or prophylactic to a mammalian organ (e.g., a liver, spleen, lung, or femur). This method includes the step of administering to a subject (e.g., a mammal) a nanoparticle composition including (i) a lipid component including a phospholipid, a PEG lipid, a structural lipid, and a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe) and (ii) a therapeutic and/or prophylactic (e.g., an mRNA), in which administering involves contacting the cell with the nanoparticle composition, whereby the therapeutic and/or prophylactic is delivered to the target organ (e.g., a liver, spleen, lung, or femur).

In some embodiments, the disclosure features a method for the enhanced delivery of a therapeutic and/or prophylactic (e.g., an mRNA) to a target tissue (e.g., a liver, spleen, lung, or femur). This method includes administering to a subject (e.g., a mammal) a nanoparticle composition, the composition including (i) a lipid component including a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe), a phospholipid, a structural lipid, and a PEG lipid; and (ii) a therapeutic and/or prophylactic, the administering including contacting the target tissue with the nanoparticle composition, whereby the therapeutic and/or prophylactic is delivered to the target tissue.

In some embodiments, the disclosure features a method of lowering immunogenicity comprising introducing the nanoparticle composition of the disclosure into cells, wherein the nanoparticle composition reduces the induction of the cellular immune response of the cells to the nanoparticle composition, as compared to the induction of the cellular immune response in cells induced by a reference composition which comprises a reference lipid instead of a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe). For example, the cellular immune response is an innate immune response, an adaptive immune response, or both.

The disclosure also includes methods of synthesizing a compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe) and methods of making a nanoparticle composition including a lipid component comprising the compound of Formula (I), (IA), (II), (IIa), (IIb), (IIc), (IId) or (IIe). Modes of Vaccine Administration

Respiratory virus RNA (e.g. mRNA) vaccines may be administered by any route which results in a therapeutically effective outcome. These include, but are not limited, to intradermal, intramuscular, and/or subcutaneous administration. The present disclosure provides methods comprising administering RNA (e.g., mRNA) vaccines to a subject in need thereof. The exact amount required will vary from subject to subject, depending on the species, age, and general condition of the subject, the severity of the disease, the particular composition, its mode of administration, its mode of activity, and the like. Respiratory virus RNA (e.g., mRNA) vaccines compositions are typically formulated in dosage unit form for ease of administration and uniformity 20 of dosage. It will be understood, however, that the total daily usage of RNA (e.g., mRNA) vaccine compositions may be decided by the attending physician within the scope of sound medical judgment. The specific therapeutically effective, prophylactically effective, or appropriate imaging dose level 25 for any particular patient will depend upon a variety of factors including the disorder being treated and the severity of the disorder; the activity of the specific compound employed; the specific composition employed; the age, body weight, general health, sex and diet of the patient; the time 30 of administration, route of administration, and rate of excretion of the specific compound employed; the duration of the treatment; drugs used in combination or coincidental with the specific compound employed; and like factors well known in the medical arts.

In some embodiments, respiratory virus RNA (e.g. mRNA) vaccines compositions may be administered at dosage levels sufficient to deliver 0.0001 mg/kg to 100 mg/kg, 0.001 mg/kg to 0.05 mg/kg, 0.005 mg/kg to 0.05 mg/kg, 0.001 mg/kg to 0.005 mg/kg, 0.05 mg/kg to 0.5 40 mg/kg, 0.01 mg/kg to 50 mg/kg, 0.1 mg/kg to 40 mg/kg, 0.5 mg/kg to 30 mg/kg, 0.01 mg/kg to 10 mg/kg, 0.1 mg/kg to 10 mg/kg, or 1 mg/kg to 25 mg/kg, of subject body weight per day, one or more times a day, per week, per month, etc. to obtain the desired therapeutic, diagnostic, prophylactic, or 45 imaging effect (see, e.g., the range of unit doses described in International Publication No WO2013078199, the contents of which are herein incorporated by reference in their entirety). The desired dosage may be delivered three times a day, two times a day, once a day, every other day, every 50 third day, every week, every two weeks, every three weeks, every four weeks, every 2 months, every three months, every 6 months, etc. In some embodiments, the desired dosage may be delivered using multiple administrations (e.g., two, three, four, five, six, seven, eight, nine, ten, eleven, twelve, 55 thirteen, fourteen, or more administrations). When multiple administrations are employed, split dosing regimens such as those described herein may be used. In exemplary embodiments, respiratory virus RNA (e.g., mRNA) vaccines compositions may be administered at dosage levels sufficient to 60 deliver 0.0005 mg/kg to 0.01 mg/kg, e.g., about 0.0005 mg/kg to about 0.0075 mg/kg, e.g., about 0.0005 mg/kg, about 0.001 mg/kg, about 0.002 mg/kg, about 0.003 mg/kg, about 0.004 mg/kg or about 0.005 mg/kg.

In some embodiments, respiratory virus RNA (e.g., 65 mRNA) vaccine compositions may be administered once or twice (or more) at dosage levels sufficient to deliver 0.025

**188** mg/kg to 0.250 mg/kg, 0.025 mg/kg to 0.500 mg/kg, 0.025 mg/kg to 0.750 mg/kg, or 0.025 mg/kg to 1.0 mg/kg.

In some embodiments, respiratory virus RNA (e.g., mRNA) vaccine compositions may be administered twice (e.g., Day 0 and Day 7, Day 0 and Day 14, Day 0 and Day 21, Day 0 and Day 28, Day 0 and Day 60, Day 0 and Day 90, Day 0 and Day 120, Day 0 and Day 150, Day 0 and Day 180, Day 0 and 3 months later, Day 0 and 6 months later, Day 0 and 9 months later, Day 0 and 12 months later, Day 0 and 18 months later, Day 0 and 2 years later, Day 0 and 5 years later, or Day 0 and 10 years later) at a total dose of or at dosage levels sufficient to deliver a total dose of 0.0100 mg, 0.025 mg, 0.050 mg, 0.075 mg, 0.100 mg, 0.125 mg, 0.150 mg, 0.175 mg, 0.200 mg, 0.225 mg, 0.250 mg, 0.275 mg, 0.300 mg, 0.325 mg, 0.350 mg, 0.375 mg, 0.400 mg, 0.425 mg, 0.450 mg, 0.475 mg, 0.500 mg, 0.525 mg, 0.550 mg, 0.575 mg, 0.600 mg, 0.625 mg, 0.650 mg, 0.675 mg, 0.700 mg, 0.725 mg, 0.750 mg, 0.775 mg, 0.800 mg, 0.825 mg, 0.850 mg, 0.875 mg, 0.900 mg, 0.925 mg, 0.950 mg, 0.975 mg, or 1.0 mg. Higher and lower dosages and frequency of administration are encompassed by the present disclosure. For example, a respiratory virus RNA (e.g., mRNA) vaccine composition may be administered three or four times.

In some embodiments, respiratory virus RNA (e.g., mRNA) vaccine compositions may be administered twice (e.g., Day 0 and Day 7, Day 0 and Day 14, Day 0 and Day 21, Day 0 and Day 28, Day 0 and Day 60, Day 0 and Day 90, Day 0 and Day 120, Day 0 and Day 150, Day 0 and Day 180, Day 0 and 3 months later, Day 0 and 6 months later, Day 0 and 9 months later, Day 0 and 12 months later, Day 0 and 18 months later, Day 0 and 2 years later, Day 0 and 5 years later, or Day 0 and 10 years later) at a total dose of or at dosage levels sufficient to deliver a total dose of 0.010 mg, 0.025 mg, 0.100 mg or 0.400 mg.

In some embodiments, the respiratory virus RNA (e.g., mRNA) vaccine for use in a method of vaccinating a subject is administered to the subject as a single dosage of between 10 μg/kg and 400 μg/kg of the nucleic acid vaccine (in an effective amount to vaccinate the subject). In some embodiments the RNA (e.g., mRNA) vaccine for use in a method of vaccinating a subject is administered to the subject as a single dosage of between 10 µg and 400 µg of the nucleic acid vaccine (in an effective amount to vaccinate the subject). In some embodiments, a respiratory virus RNA (e.g., mRNA) vaccine for use in a method of vaccinating a subject is administered to the subject as a single dosage of 25-1000 μg (e.g., a single dosage of mRNA encoding hMPV, PIV3, RSV, MeV and/or BetaCoV antigen). In some embodiments, a respiratory virus RNA (e.g., mRNA) vaccine is administered to the subject as a single dosage of 25, 50, 100, 150, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950 or 1000 µg. For example, a respiratory virus RNA (e.g., mRNA) vaccine may be administered to a subject as a single dose of 25-100, 25-500, 50-100, 50-500, 50-1000, 100-500, 100-1000, 250-500, 250-1000, or 500-1000 μg. In some embodiments, a respiratory virus RNA (e.g., mRNA) vaccine for use in a method of vaccinating a subject is administered to the subject as two dosages, the combination of which equals 25-1000 µg of the respiratory virus RNA (e.g., mRNA) vaccine.

A respiratory virus RNA (e.g. mRNA) vaccine pharmaceutical composition described herein can be formulated into a dosage form described herein, such as an intranasal, intratracheal, or injectable (e.g., intravenous, intraocular, intravitreal, intramuscular, intradermal, intracardiac, intraperitoneal, and subcutaneous).

Respiratory Virus RNA (e.g., mRNA) Vaccine Formulations and Methods of Use

Some aspects of the present disclosure provide formulations of the respiratory virus RNA (e.g., mRNA) vaccine, wherein the RNA (e.g., mRNA) vaccine is formulated in an 5 effective amount to produce an antigen specific immune response in a subject (e.g., production of antibodies specific to an hMPV, PIV3, RSV, MeV and/or BetaCoV antigenic polypeptide). "An effective amount" is a dose of an RNA (e.g., mRNA) vaccine effective to produce an antigenspecific immune response. Also provided herein are methods of inducing an antigen-specific immune response in a subject.

In some embodiments, the antigen-specific immune response is characterized by measuring an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide antibody titer produced in a subject administered a respiratory virus RNA (e.g., mRNA) vaccine as provided herein. An antibody titer is a measurement of the amount of antibodies within a subject, for example, antibodies that are specific to a particular antigen (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) or epitope of an antigen. Antibody titer is typically expressed as the inverse of the greatest dilution that provides a positive result. Enzymelinked immunosorbent assay (ELISA) is a common assay for determining antibody titers, for example.

In some embodiments, an antibody titer is used to assess whether a subject has had an infection or to determine whether immunizations are required. In some embodiments, 30 an antibody titer is used to determine the strength of an autoimmune response, to determine whether a booster immunization is needed, to determine whether a previous vaccine was effective, and to identify any recent or prior infections. In accordance with the present disclosure, an 35 antibody titer may be used to determine the strength of an immune response induced in a subject by the respiratory virus RNA (e.g., mRNA) vaccine.

In some embodiments, an anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-40 BetaCoV antigenic polypeptide) antibody titer produced in a subject is increased by at least 1 log relative to a control. For example, anti-antigenic polypeptide antibody titer produced in a subject may be increased by at least 1.5, at least 2, at least 2.5, or at least 3 log relative to a control. In some 45 embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased by 1, 1.5, 2, 2.5 or 3 log relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased by 1-3 log relative to a control. For example, the 50 anti-antigenic polypeptide antibody titer produced in a subject may be increased by 1-1.5, 1-2, 1-2.5, 1-3, 1.5-2, 1.5-2.5, 1.5-3, 2-2.5, 2-3, or 2.5-3 log relative to a control.

In some embodiments, the anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or 55 anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject is increased at least 2 times relative to a control. For example, the anti-antigenic polypeptide antibody titer produced in a subject may be increased at least 3 times, at least 4 times, at least 5 times, at least 6 times, at least 7 times, at least 8 times, at least 9 times, or at least 10 times relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is increased 2, 3, 4, 5, 6, 7, 8, 9, or 10 times relative to a control. In some embodiments, the anti-antigenic polypeptide antibody titer produced in a subject is increased 2-10 times relative to a control. For example, the anti-antigenic

190

polypeptide antibody titer produced in a subject may be increased 2-10, 2-9, 2-8, 2-7, 2-6, 2-5, 2-4, 2-3, 3-10, 3-9, 3-8, 3-7, 3-6, 3-5, 3-4, 4-10, 4-9, 4-8, 4-7, 4-6, 4-5, 5-10, 5-9, 5-8, 5-7, 5-6, 6-10, 6-9, 6-8, 6-7, 7-10, 7-9, 7-8, 8-10, 8-9, or 9-10 times relative to a control.

A control, in some embodiments, is the anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject who has not been administered a respiratory virus RNA (e.g., mRNA) vaccine of the present disclosure. In some embodiments, a control is an antiantigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject who has been administered a live attenuated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine. An attenuated vaccine is a vaccine produced by reducing the virulence of a viable (live). An attenuated virus is altered in a manner that renders it harmless or less virulent relative to live, unmodified virus. In some embodiments, a control is an anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject administered inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine. In some embodiments, a control is an anti-antigenic polypeptide (e.g., an antihMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject administered a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine. Recombinant protein vaccines typically include protein antigens that either have been produced in a heterologous expression system (e.g., bacteria or yeast) or purified from large amounts of the pathogenic organism. In some embodiments, a control is an anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject who has been administered an hMPV, PIV3, RSV, MeV and/or BetaCoV virus-like particle (VLP) vaccine. For example, an hMPV VLP vaccine used as a control may be a hMPV VLPs. comprising (or consisting of) viral matrix (M) and fusion (F) proteins, generated by expressing viral proteins in suspension-adapted human embryonic kidney epithelial (293-F) cells (see, e.g., Cox R G et al., J Virol. 2014 June; 88(11): 6368-6379, the contents of which are herein incorporated by reference).

In some embodiments, an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose that is reduced compared to the standard of care dose of a recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine. A "standard of care," as provided herein, refers to a medical or psychological treatment guideline and can be general or specific. "Standard of care" specifies appropriate treatment based on scientific evidence and collaboration between medical professionals involved in the treatment of a given condition. It is the diagnostic and treatment process that a physician/clinician should follow for a certain type of patient, illness or clinical circumstance. A "standard of care dose," as provided herein, refers to the dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, or a live attenuated or inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine, that a physician/ clinician or other medical professional would administer to a subject to treat or prevent hMPV, PIV3, RSV, MeV and/or BetaCoV, or a hMPV-, PIV3-, RSV-, MeV- and/or BetaCoVrelated condition, while following the standard of care

guideline for treating or preventing hMPV, PIV3, RSV, MeV and/or BetaCoV, or a hMPV-, PIV3-, RSV-, MeV- and/or BetaCoV-related condition.

In some embodiments, the anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or 5 anti-BetaCoV antigenic polypeptide) antibody titer produced in a subject administered an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is equivalent to an anti-antigenic polypeptide (e.g., an anti-hMPV, anti-PIV3, anti-RSV, anti-MeV and/or anti-BetaCoV antigenic polypeptide) antibody titer produced in a control subject administered a standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine or a live attenuated or inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine.

In some embodiments, an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose equivalent to an at least 2-fold reduction in a standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine. For example, an effective amount 20 of a respiratory virus RNA (e.g., mRNA) vaccine may be a dose equivalent to an at least 3-fold, at least 4-fold, at least 5-fold, at least 6-fold, at least 7-fold, at least 8-fold, at least 9-fold, or at least 10-fold reduction in a standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or 25 BetaCoV protein vaccine. In some embodiments, an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose equivalent to an at least at least 100-fold, at least 500-fold, or at least 1000-fold reduction in a standard of care dose of a recombinant or purified hMPV, PIV3, RSV, 30 MeV and/or BetaCoV protein vaccine. In some embodiments, an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose equivalent to a 2-, 3-, 4-, 5-, 6-, 7-, 8-, 9-, 10-, 20-, 50-, 100-, 250-, 500-, or 1000-fold reduction in a standard of care dose of a recombinant or 35 purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine. In some embodiments, the anti-antigenic polypeptide antibody titer produced in a subject administered an effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is equivalent to an anti-antigenic polypeptide anti- 40 body titer produced in a control subject administered the standard of care dose of a recombinant or protein hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine or a live attenuated or inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine. In some embodiments, an effective 45 amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose equivalent to a 2-fold to 1000-fold (e.g., 2-fold to 100-fold, 10-fold to 1000-fold) reduction in the standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine, wherein the anti- 50 antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine or a live attenuated or 55 inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vac-

In some embodiments, the effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a dose equivalent to a 2 to 1000-, 2 to 900-, 2 to 800-, 2 to 700-, 2 to 600-, 2 60 to 500-, 2 to 400-, 2 to 300-, 2 to 200-, 2 to 100-, 2 to 90-, 2 to 80-, 2 to 70-, 2 to 60-, 2 to 50-, 2 to 40-, 2 to 30-, 2 to 20-, 2 to 10-, 2 to 9-, 2 to 8-, 2 to 7-, 2 to 6-, 2 to 5-, 2 to 4-, 2 to 3-, 3 to 1000-, 3 to 900-, 3 to 800-, 3 to 700-, 3 to 600-, 3 to 500-, 3 to 400-, 3 to 500-, 3 to 500-, 3 to 500-, 3 to 50-, 3 to 50-, 3 to 40-, 3 to 30-, 3 to 20-, 3 to 10-, 3 to 9-, 3 to 8-, 3 to 7-, 3 to 6-, 3 to

192

5-, 3 to 4-, 4 to 1000-, 4 to 900-, 4 to 800-, 4 to 700-, 4 to 600-, 4 to 500-, 4 to 400-, 4 to 4 to 00-, 4 to 200-, 4 to 100-, 4 to 90-, 4 to 80-, 4 to 70-, 4 to 60-, 4 to 50-, 4 to 40-, 4 to 30-, 4 to 20-, 4 to 10-, 4 to 9-, 4 to 8-, 4 to 7-, 4 to 6-, 4 to 5-, 4 to 4-, 5 to 1000-, 5 to 900-, 5 to 800-, 5 to 700-, 5 to 600-, 5 to 500-, 5 to 400-, 5 to 300-, 5 to 200-, 5 to 100-, 5 to 90-, 5 to 80-, 5 to 70-, 5 to 60-, 5 to 50-, 5 to 40-, 5 to 30-, 5 to 20-, 5 to 10-, 5 to 9-, 5 to 8-, 5 to 7-, 5 to 6-, 6 to 1000-, 6 to 900-, 6 to 800-, 6 to 700-, 6 to 600-, 6 to 500-, 6 to 400-, 6 to 300-, 6 to 200-, 6 to 100-, 6 to 90-, 6 to 80-, 6 to 70-, 6 to 60-, 6 to 50-, 6 to 40-, 6 to 30-, 6 to 20-, 6 to 10-, 6 to 9-, 6 to 8-, 6 to 7-, 7 to 1000-, 7 to 900-, 7 to 800-, 7 to 700-, 7 to 600-, 7 to 500-, 7 to 400-, 7 to 300-, 7 to 200-, 7 to 100-, 7 to 90-, 7 to 80-, 7 to 70-, 7 to 60-, 7 to 50-, 7 to 40-, 7 to 30-, 7 to 20-, 7 to 10-, 7 to 9-, 7 to 8-, 8 to 1000-, 8 to 900-, 8 to 800-, 8 to 700-, 8 to 600-, 8 to 500-, 8 to 400-, 8 to 300-, 8 to 200-, 8 to 100-, 8 to 90-, 8 to 80-, 8 to 70-, 8 to 60-, 8 to 50-, 8 to 40-, 8 to 30-, 8 to 20-, 8 to 10-, 8 to 9-, 9 to 1000-, 9 to 900-, 9 to 800-, 9 to 700-, 9 to 600-, 9 to 500-, 9 to 400-, 9 to 300-, 9 to 200-, 9 to 100-, 9 to 90-, 9 to 80-, 9 to 70-, 9 to 60-, 9 to 50-, 9 to 40-, 9 to 30-, 9 to 20-, 9 to 10-, 10 to 1000-, 10 to 900-, 10 to 800-, 10 to 700-, 10 to 600-, 10 to 500-, 10 to 400-, 10 to 300-, 10 to 200-, 10 to 100-, 10 to 90-, 10 to 80-, 10 to 70-, 10 to 60-, 10 to 50-, 10 to 40-, 10 to 30-, 10 to 20-, 20 to 1000-, 20 to 900-, 20 to 800-, 20 to 700-, 20 to 600-, 20 to 500-, 20 to 400-, 20 to 300-, 20 to 200-, 20 to 100-, 20 to 90-, 20 to 80-, 20 to 70-, 20 to 60-, 20 to 50-, 20 to 40-, 20 to 30-, 30 to 1000-, 30 to 900-, 30 to 800-, 30 to 700-, 30 to 600-, 30 to 500-, 30 to 400-, 30 to 300-, 30 to 200-, 30 to 100-, 30 to 90-, 30 to 80-, 30 to 70-, 30 to 60-, 30 to 50-, 30 to 40-, 40 to 1000-, 40 to 900-, 40 to 800-, 40 to 700-, 40 to 600-, 40 to 500-, 40 to 400-, 40 to 300-, 40 to 200-, 40 to 100-, 40 to 90-, 40 to 80-, 40 to 70-, 40 to 60-, 40 to 50-, 50 to 1000-, 50 to 900-, 50 to 800-, 50 to 700-, 50 to 600-, 50 to 500-, 50 to 400-, 50 to 300-, 50 to 200-, 50 to 100-, 50 to 90-, 50 to 80-, 50 to 70-, 50 to 60-, 60 to 1000-, 60 to 900-, 60 to 800-, 60 to 700-, 60 to 600-, 60 to 500-, 60 to 400-, 60 to 300-, 60 to 200-, 60 to 100-, 60 to 90-, 60 to 80-, 60 to 70-, 70 to 1000-, 70 to 900-, 70 to 800-, 70 to 700-, 70 to 600-, 70 to 500-, 70 to 400-, 70 to 300-, 70 to 200-, 70 to 100-, 70 to 90-, 70 to 80-, 80 to 1000-, 80 to 900-, 80 to 800-, 80 to 700-, 80 to 600-, 80 to 500-, 80 to 400-, 80 to 300-, 80 to 200-, 80 to 100-, 80 to 90-, 90 to 1000-, 90 to 900-, 90 to 800-, 90 to 700-, 90 to 600-, 90 to 500-, 90 to 400-, 90 to 300-, 90 to 200-, 90 to 100-, 100 to 1000-, 100 to 900-, 100 to 800-, 100 to 700-, 100 to 600-, 100 to 500-, 100 to 400-, 100 to 300-, 100 to 200-, 200 to 1000-, 200 to 900-, 200 to 800-, 200 to 700-, 200 to 600-, 200 to 500-, 200 to 400-, 200 to 300-, 300 to 1000-, 300 to 900-, 300 to 800-, 300 to 700-, 300 to 600-, 300 to 500-, 300 to 400-, 400 to 1000-, 400 to 900-, 400 to 800-, 400 to 700-, 400 to 600-, 400 to 500-, 500 to 1000-, 500 to 900-, 500 to 800-, 500 to 700-, 500 to 600-, 600 to 1000-, 600 to 900-, 600 to 800-, 600 to 700-, 700 to 1000-, 700 to 900-, 700 to 800-, 800 to 1000-, 800 to 900-, or 900 to 1000-fold reduction in the standard of care dose of a recombinant hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine. In some embodiments, the anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant or purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine or a live attenuated or inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine. In some embodiments, the effective amount is a dose equivalent to (or equivalent to an at least) 2-, 3-, 4-, 5-, 6-, 7-, 8-, 9-, 10-, 20-, 30-, 40-, 50-, 60-, 70-, 80-, 90-, 100-, 110-, 120-, 130-,

140-, 150-, 160-, 170-, 1280-, 190-, 200-, 210-, 220-, 230-, 240-, 250-, 260-, 270-, 280-, 290-, 300-, 310-, 320-, 330-, 340-, 350-, 360-, 370-, 380-, 390-, 400-, 410-, 420-, 430-, 440-, 450-, 4360-, 470-, 480-, 490-, 500-, 510-, 520-, 530-, 540-, 550-, 560-, 5760-, 580-, 590-, 600-, 610-, 620-, 630-, 5 640-, 650-, 660-, 670-, 680-, 690-, 700-, 710-, 720-, 730-, 740-, 750-, 760-, 770-, 780-, 790-, 800-, 810-, 820-, 830-, 840-, 850-, 860-, 870-, 880-, 890-, 900-, 910-, 920-, 930-, 940-, 950-, 960-, 970-, 980-, 990-, or 1000-fold reduction in the standard of care dose of a recombinant hMPV, PIV3, 10 RSV, MeV and/or BetaCoV protein vaccine. In some embodiments, an anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant or 15 purified hMPV, PIV3, RSV, MeV and/or BetaCoV protein vaccine or a live attenuated or inactivated hMPV, PIV3, RSV, MeV and/or BetaCoV vaccine.

In some embodiments, the effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a total dose of 20 50-1000 µg. In some embodiments, the effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a total dose of 50-1000, 50-900, 50-800, 50-700, 50-600, 50-500, 50-400, 50-300, 50-200, 50-100, 50-90, 50-80, 50-70, 50-60, 60-1000, 60-900, 60-800, 60-700, 60-600, 60-500, 25 60-400, 60-300, 60-200, 60-100, 60-90, 60-80, 60-70, 70-1000, 70-900, 70-800, 70-700, 70-600, 70-500, 70-400, 70-300, 70-200, 70-100, 70-90, 70-80, 80-1000, 80-900, 80-800, 80-700, 80-600, 80-500, 80-400, 80-300, 80-200, 80-100, 80-90, 90-1000, 90-900, 90-800, 90-700, 90-600, 30 90-500, 90-400, 90-300, 90-200, 90-100, 100-1000, 100-900, 100-800, 100-700, 100-600, 100-500, 100-400, 100-300, 100-200, 200-1000, 200-900, 200-800, 200-700, 200-600, 200-500, 200-400, 200-300, 300-1000, 300-900, 300-800, 300-700, 300-600, 300-500, 300-400, 400-1000, 400-35 900, 400-800, 400-700, 400-600, 400-500, 500-1000, 500-900, 500-800, 500-700, 500-600, 600-1000, 600-900, 600-900, 600-700, 700-1000, 700-900, 700-800, 800-1000, 800-900, or 900-1000 µg. In some embodiments, the effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is 40 a total dose of 50, 100, 150, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950 or 1000 μg. In some embodiments, the effective amount is a dose of 25-500 μg administered to the subject a total of two times. In some embodiments, the effective amount of a respiratory 45 virus RNA (e.g., mRNA) vaccine is a dose of 25-500, 25-400, 25-300, 25-200, 25-100, 25-50, 50-500, 50-400, 50-300, 50-200, 50-100, 100-500, 100-400, 100-300, 100-200, 150-500, 150-400, 150-300, 150-200, 200-500, 200-400, 200-300, 250-500, 250-400, 250-300, 300-500, 300-50 400, 350-500, 350-400, 400-500 or 450-500 μg administered to the subject a total of two times. In some embodiments, the effective amount of a respiratory virus RNA (e.g., mRNA) vaccine is a total dose of 25, 50, 100, 150, 200, 250, 300, 350, 400, 450, or 500 μg administered to the subject a total 55 of two times.

# Examples of Additional Embodiments of the Disclosure

Additional embodiments of the present disclosure are encompassed by the following numbered paragraphs:

1. A respiratory virus vaccine, comprising: at least one ribonucleic acid (RNA) polynucleotide having an open reading frame encoding at least one, at least two, at least 65 three, at least four or at least five antigenic polypeptides selected from human *Metapneumovirus* (hMPV) antigenic

polypeptides or immunogenic fragments thereof, human parainfluenza virus type 3 (PIV3) antigenic polypeptides or immunogenic fragments thereof, respiratory syncytial virus (RSV) antigenic polypeptides or immunogenic fragments thereof, measles virus (MeV) antigenic polypeptides or immunogenic fragments thereof, and *Betacoronavirus* (BetaCoV) antigenic polypeptides or immunogenic fragments thereof.

2. The respiratory virus vaccine of paragraph 1, comprising: at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and a PIV3 antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof.

3. The respiratory virus vaccine of paragraph 2, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13.

4. The respiratory virus vaccine of paragraph 1, comprising: at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and a RSV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof.

5. The respiratory virus vaccine of paragraph 4, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8.

6. The respiratory virus vaccine of paragraph 1, comprising: at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and MeV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

7. The respiratory virus vaccine of paragraph 6, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50.

8. The respiratory virus vaccine of paragraph 1, comprising: at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immu-

nogenic fragment thereof and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

9. The respiratory virus vaccine of paragraph 8, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

 The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and a RSV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open 25 reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof.

11. The respiratory virus vaccine of paragraph 10, wherein 30 the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13.

12. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and a MeV antigenic polypeptide 40 or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a MeV antigenic polypeptide or an 45 immunogenic fragment thereof.

13. The respiratory virus vaccine of paragraph 12, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to 50 an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence 55 identified by any one of SEQ ID NO: 47-50.

14. The respiratory virus vaccine of paragraph 1, comprising

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and a BetaCoV antigenic

polypeptide or an immunogenic fragment thereof; or at least two RNA polynucleotides, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

196

15. The respiratory virus vaccine of paragraph 14, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

16. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof and a MeV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

17. The respiratory virus vaccine of paragraph 16, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50.

18. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

19. The respiratory virus vaccine of paragraph 18, wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

20. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two RNA polynucleotides, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

21. The respiratory virus vaccine of paragraph 20, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

22. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, and a RSV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof.

23. The respiratory virus vaccine of paragraph 22, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13.

24. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or 30 an immunogenic fragment thereof, and a MeV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open 35 reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

25. The respiratory virus vaccine of paragraph 24, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50. 26. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, and one having an open 65 reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

198

27. The respiratory virus vaccine of paragraph 26, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13 and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34. 28. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a MeV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

29. The respiratory virus vaccine of paragraph 28, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50.

30. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

31. The respiratory virus vaccine of paragraph 30, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34.

32. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or

an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open 5 reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

33. The respiratory virus vaccine of paragraph 32, wherein 10 the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34. 34. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a MeV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open 35 reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

35. The respiratory virus vaccine of paragraph 34, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an 40 amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at 45 least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50.

36. The respiratory virus vaccine of paragraph 1, comprising

at least one RNA polynucleotide having an open reading 50 frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an 55 open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or 60 an immunogenic fragment thereof.

37. The respiratory virus vaccine of paragraph 36, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to 65 an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the BetaCoV antigenic polypep-

tide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34.

200

38. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

39. The respiratory virus vaccine of paragraph 38, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34.

40. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two or three RNA polynucleotides, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

41. The respiratory virus vaccine of paragraph 40, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 23-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 23-34.

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a MeV antigenic polypeptide or an immunogenic fragment thereof;

at least two, three or four RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic

polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof.

43. The respiratory virus vaccine of paragraph 42, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% 15 or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50. 44. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, and a Beta-CoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two, three or four RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

45. The respiratory virus vaccine of paragraph 44, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a Beta-CoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two, three or four RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one hav-

ing an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

47. The respiratory virus vaccine of paragraph 46, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 12-13, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEO ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

48. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two, three or four RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

49. The respiratory virus vaccine of paragraph 48, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

at least one RNA polynucleotide having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two, three or four RNA polynucleotides, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one hav-

ing an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

51. The respiratory virus vaccine of paragraph 50, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to 10 an amino acid sequence identified by any one of SEQ ID NO: 12-13, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by 15 any one of SEQ ID NO: 47-50, and/or wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34. 20 52. The respiratory virus vaccine of paragraph 1, comprising:

at least one RNA polynucleotide having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, a PIV3 antigenic polypeptide or an immunogenic fragment thereof, a RSV antigenic polypeptide or an immunogenic fragment thereof, a MeV antigenic polypeptide or an immunogenic fragment thereof, and a BetaCoV antigenic polypeptide or an immunogenic fragment thereof; or

at least two, three, four or five RNA polynucleotides, one having an open reading frame encoding a hMPV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a PIV3 antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a RSV antigenic polypeptide or an immunogenic fragment thereof, one having an open reading frame encoding a MeV antigenic polypeptide or an immunogenic fragment thereof, and one having an open reading frame encoding a BetaCoV antigenic polypeptide or an immunogenic fragment thereof.

53. The respiratory virus vaccine of paragraph 52, wherein the hMPV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 5-8 or an amino acid sequence having at least 90% or 95% identity to 45 an amino acid sequence identified by any one of SEQ ID NO: 5-8, wherein the PIV3 antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 12-13 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any 50 one of SEQ ID NO: 12-13, wherein the MeV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 47-50 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 47-50, and/or 55 wherein the BetaCoV antigenic polypeptide comprises an amino acid sequence identified by any one of SEQ ID NO: 24-34 or an amino acid sequence having at least 90% or 95% identity to an amino acid sequence identified by any one of SEQ ID NO: 24-34.

54. The vaccine of any one of paragraphs 1-53, wherein at least one RNA polynucleotide has less than 80% identity to wild-type mRNA sequence.

55. The vaccine of any one of paragraphs 1-53, wherein at least one RNA polynucleotide has at least 80% identity to 65 wild-type mRNA sequence, but does not include wild-type mRNA sequence.

204

56. The vaccine of any one of paragraphs 1-55, wherein at least one antigenic polypeptide has membrane fusion activity, attaches to cell receptors, causes fusion of viral and cellular membranes, and/or is responsible for binding of the virus to a cell being infected.

57. The vaccine of any one of paragraphs 1-56, wherein at least one RNA polynucleotide comprises at least one chemical modification.

58. The vaccine of paragraph 57, wherein the chemical modification is selected from pseudouridine, N1-methylpseudouridine, N1-ethylpseudouridine, 2-thiouridine, 4'-thiouridine, 5-methylcytosine, 5-methyluridine, 2-thio-1-methyl-1-deaza-pseudouridine, 2-thio-1-methyl-pseudouridine, 2-thio-5-aza-uridine, 2-thio-dihydropseudouridine, 2-thio-pseudouridine, 4-methoxy-2-thio-pseudouridine, 4-methoxy-pseudouridine, 4-thio-1-methyl-pseudouridine, 4-thio-pseudouridine, 5-aza-uridine, dihydropseudouridine, 5-methoxyuridine and 2'-O-methyl uridine.

59. The vaccine of paragraph 57 or 58, wherein the chemical modification is in the 5-position of the uracil.

60. The vaccine of any one of paragraphs 57-59, wherein the chemical modification is a N1-methylpseudouridine or N1-ethylpseudouridine.

61. The vaccine of any one of paragraphs 57-60, wherein at least 80%, at least 90% or 100% of the uracil in the open reading frame have a chemical modification.

62. The vaccine of any one of paragraphs 1-61, wherein at least one RNA polynucleotide further encodes at least one 5' terminal cap, optionally wherein the 5' terminal cap is 7mG(5')ppp(5')NlmpNp.

63. The vaccine of any one of paragraphs 1-62, wherein at least one antigenic polypeptide or immunogenic fragment thereof is fused to a signal peptide selected from: a HuIgGk signal peptide (METPAQLLFLLLWLPDTTG; SEQ ID NO: 15); IgE heavy chain epsilon-1 signal peptide (MDWTWILFLVAAATRVHS; SEQ ID NO: 16); Japanese encephalitis PRM signal sequence (MLG-SNSGQRVVFTILLLLVAPAYS; SEQ ID NO: 17), VSVg protein signal sequence (MKCLLYLAFLFIGVNCA; SEQ ID NO: 18) and Japanese encephalitis JEV signal sequence (MWLVSLAIVTACAGA; SEQ ID NO:19).

64. The vaccine of paragraph 63, wherein the signal peptide is fused to the N-terminus or the C-terminus of at least one antigenic polypeptide.

65. The vaccine of any one of paragraphs 1-64, wherein the antigenic polypeptide or immunogenic fragment thereof comprises a mutated N-linked glycosylation site.

66. The vaccine of any one of paragraphs 1-65 formulated in a nanoparticle, optionally a lipid nanoparticle.

67. The vaccine of paragraph 66, wherein the lipid nanoparticle comprises a cationic lipid, a PEG-modified lipid, a sterol and a non-cationic lipid; optionally wherein the lipid nanoparticle carrier comprises a molar ratio of about 20-60% cationic lipid, 0.5-15% PEG-modified lipid, 25-55% sterol, and 25% non-cationic lipid; optionally wherein the cationic lipid is an ionizable cationic lipid and the non-cationic lipid is a neutral lipid, and the sterol is a cholesterol; and optionally wherein the cationic lipid is selected from 2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane (DLin-KC2-DMA), dilinoleyl-methyl-4-dimethylaminobutyrate (DLin-MC3-DMA), and di((Z)-non-2-en-1-yl) 9-((4-(dimethylamino)butanoyl)oxy)heptadecanedioate (L319). Formula (II)

68. The vaccine of paragraph 66 or 67, wherein the nanoparticle (e.g., lipid nanoparticle) comprises a compound of

Formula (I) and/or Formula (II), optionally Compound 3, 18, 20, 25, 26, 29, 30, 60, 108-112, or 122.

- 69. The vaccine of any one of paragraphs 1-68 further comprising an adjuvant, optionally a flagellin protein or peptide that optionally comprises an amino acid sequence 5 identified by any one of SEQ ID NO: 54-56.
- 70. The vaccine of any one of paragraphs 1-69, wherein the open reading frame is codon-optimized.
- 71. The vaccine of any one of paragraphs 1-70 formulated in an effective amount to produce an antigen-specific immune 10 response.
- 72. A method of inducing an immune response in a subject, the method comprising administering to the subject the vaccine of any one of paragraphs 1-71 in an amount effective to produce an antigen-specific immune response in the 15 subject.
- 73. The method of paragraph 72, wherein the subject is administered a single dose of the vaccine, or wherein the subject is administered a first dose and then a booster dose of the vaccine.
- 74. The method of paragraph 72 or 73, wherein the vaccine is administered to the subject by intradermal injection or intramuscular injection.
- 75. The method of any one of paragraphs 72-74, wherein an anti-antigenic polypeptide antibody titer produced in the 25 subject is increased by at least 1 log relative to a control, and/or wherein the anti-antigenic polypeptide antibody titer produced in the subject is increased at least 2 times relative to a control.
- 76. The method of any one of paragraphs 72-75, wherein the control is an anti-antigenic polypeptide antibody titer produced in a subject who has not been administered a vaccine against the virus, and/or wherein the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a recombinant protein vaccine or purified protein vaccine against the virus, and/or wherein the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a recombinant protein vaccine or purified protein vaccine against the virus, and/or wherein the control is an anti-antigenic polypeptide antibody titer produced in a subject who has been administered a vaccine about 0.5-2 about 0.5-2 (b) com wherein the control is an anti-antigenic polypeptide at least for at least for open reading the virus.
- 77. The method of any one of paragraphs 72-76, wherein the effective amount is a dose equivalent to an at least 2-fold reduction in the standard of care dose of a recombinant 45 protein vaccine or a purified protein vaccine against the virus, and wherein an anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a recombinant 50 protein vaccine or a purified protein vaccine against the virus, respectively; and/or wherein the effective amount is a dose equivalent to an at least 2-fold reduction in the standard of care dose of a live attenuated vaccine or an inactivated vaccine against the virus, and wherein an anti-antigenic 55 polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject administered the standard of care dose of a live attenuated vaccine or an inactivated vaccine against the virus, respectively; and/or wherein the effective amount 60 is a dose equivalent to an at least 2-fold reduction in the standard of care dose of a VLP vaccine against the virus, and wherein an anti-antigenic polypeptide antibody titer produced in the subject is equivalent to an anti-antigenic polypeptide antibody titer produced in a control subject 65 administered the standard of care dose of a VLP vaccine against the virus.

- 78. The method of any one of paragraphs 72-77, wherein the effective amount is a total dose of 50  $\mu$ g-1000  $\mu$ g, optionally wherein the effective amount is a dose of 25  $\mu$ g, 100  $\mu$ g, 400  $\mu$ g, or 500  $\mu$ g administered to the subject a total of two times. 79. The method of any one of paragraphs 72-78, wherein the efficacy of the vaccine against the virus is greater than 65%; and/or wherein the vaccine immunizes the subject against the virus for up to 2 years or wherein the vaccine immunizes the subject against the virus for more than 2 years.
- 80. The method of any one of paragraphs 72-79, wherein the subject has an age of about 5 years old or younger or wherein the subject has an age of about 60 years old or older; and/or wherein the subject has a chronic pulmonary disease; and/or the subject has been exposed to the virus, wherein the subject is infected with the virus, or wherein the subject is at risk of infection by the virus; and/or wherein the subject is immunocompromised.
- 81. The respiratory virus vaccine of any one of paragraphs 1-71, comprising at least one (e.g., at least two, at least three, at least four, or at least five) RNA polynucleotide having an open reading frame encoding at least one (e.g., at least two, at least three, at least four, or at least five) antigenic polypeptide selected from hMPV antigenic polypeptides (SEQ ID NO: 5-8), PIV3 antigenic polypeptides (SEQ ID NO: 12-13), RSV antigenic polypeptides, MeV antigenic polypeptides (SEQ ID NO: 47-50) and BetaCoV antigenic polypeptides (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1; (SEQ ID NO: 24-34)), formulated in a cat-30 ionic lipid nanoparticle
  - (a) having a molar ratio of about 20-60% cationic lipid, about 5-25% non-cationic lipid, about 25-55% sterol, and about 0.5-15% PEG-modified lipid, and/or
  - (b) comprising a compound of Formula (I) and/or Formula (II).
  - wherein the at least one (e.g., at least two, at least three, at least four, or at least five) RNA polynucleotide comprises at least one chemical modification.
  - 82. The respiratory virus vaccine of any one of paragraphs 1-71, comprising at least one (e.g., at least two, at least three, at least four, or at least five) RNA polynucleotide having an open reading frame encoding at least one (e.g., at least two, at least three, at least four, or at least five) antigenic polypeptide selected from hMPV antigenic polypeptides (SEQ ID NO: 5-8), PIV3 antigenic polypeptides (SEQ ID NO: 12-13), RSV antigenic polypeptides, MeV antigenic polypeptides (SEQ ID NO: 47-50) and BetaCoV antigenic polypeptides (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1; (SEQ ID NO: 24-34)), formulated in a cationic lipid nanoparticle
  - (a) having a molar ratio of about 20-60% cationic lipid, about 5-25% non-cationic lipid, about 25-55% sterol, and about 0.5-15% PEG-modified lipid, and/or
  - (b) comprising at least one (e.g., at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, or 14) Compound selected from Compounds 3, 18, 20, 25, 26, 29, 30, 60, 108-112 and 122. 83. The respiratory virus vaccine of paragraphs 81 or 82, wherein the at least one antigenic polypeptide is selected from hMPV antigentic polypeptides (e.g., SEQ ID NO: 5-8). 84. The respiratory virus vaccine of any one of paragraphs 81-83, wherein the at least one antigenic polypeptide is selected from PIV3 antigentic polypeptides (e.g., SEQ ID NO: 12-13).
  - 85. The respiratory virus vaccine of any one of paragraphs 81-84, wherein the at least one antigenic polypeptide is selected from RSV antigentic polypeptides.

86. The respiratory virus vaccine of any one of paragraphs 81-85, wherein the at least one antigenic polypeptide is selected from MeV antigentic polypeptides (e.g., SEQ ID NO: 47-50).

87. The respiratory virus vaccine of any one of paragraphs 5 81-86, wherein the at least one antigenic polypeptide is selected from BetaCoV antigentic polypeptides (e.g., SEQ ID NO: 24-34).

88. The respiratory virus vaccine of paragraph 87, wherein the BetaCoV antigentic polypeptides are MERS antigentic 10 polypeptides.

89. The respiratory virus vaccine of paragraph 87, wherein the BetaCoV antigentic polypeptides are SARS antigentic polypeptides.

90. The respiratory virus vaccine of any one of paragraphs 15 81-89, wherein the at least one (e.g., at least two, at least three, at least four, or at least five) RNA polynucleotide comprises at least one chemical modification (e.g., selected from pseudouridine, N1-methylpseudouridine, N1-ethylpseudouridine, 2-thiouridine, 4'-thiouridine, 5-methylcyto- 20 sine, 5-methyluridine, 2-thio-1-methyl-1-deaza-pseudouridine, 2-thio-1-methyl-pseudouridine, 2-thio-5-aza-uridine, 2-thio-dihydropseudouridine, 2-thio-dihydrouridine, 2-thiopseudouridine, 4-methoxy-2-thio-pseudouridine, 4-thio-pseudouridine, 5-aza-uridine, dihydropseudouridine, 5-methoxyuridine and 2'-O-methyl uridine).

91. A respiratory virus vaccine, comprising:

at least one messenger ribonucleic acid (mRNA) polynucleotide having a 5' terminal cap, an open reading frame 30 encoding at least one respiratory virus antigenic polypeptide, and a 3' polyA tail.

92. The vaccine of paragraph 91, wherein the at least one mRNA polynucleotide comprises a sequence identified by any one of SEQ ID NO: 57-80.

93. The vaccine of paragraph 91 or 92, wherein the 5' terminal cap is or comprises 7mG(5')ppp(5')NlmpNp.

94. The vaccine of any one of paragraphs 91-93, wherein 100% of the uracil in the open reading frame is modified to include N1-methyl pseudouridine at the 5-position of the 40 of a chimeric polynucleotide may be joined or ligated using uracil.

95. The vaccine of any one of paragraphs 91-94, wherein the vaccine is formulated in a lipid nanoparticle comprising: DLin-MC3-DMA; cholesterol; 1,2-Distearoyl-sn-glycero-3phosphocholine (DSPC); and polyethylene glycol (PEG) 45 2000-DMG.

96. The vaccine of paragraph 95, wherein the lipid nanoparticle further comprises trisodium citrate buffer, sucrose and

97. A respiratory syncytial virus (RSV) vaccine, comprising: 50 at least one messenger ribonucleic acid (mRNA) polynucleotide having a 5' terminal cap 7mG(5')ppp(5')NlmpNp, a sequence identified by any one of SEQ ID NO: 57-80 and a 3' polyA tail, formulated in a lipid nanoparticle comprising DLin-MC3-DMA, cholesterol, 1,2-Distearoyl-sn-glycero-3-55 phosphocholine (DSPC), and polyethylene glycol (PEG) 2000-DMG, wherein the uracil nucleotides of the sequence identified by any one of SEQ ID NO: 57-80 are modified to include N1-methyl pseudouridine at the 5-position of the uracil nucleotide.

This disclosure is not limited in its application to the details of construction and the arrangement of components set forth in the following description or illustrated in the drawings. The disclosure is capable of other embodiments and of being practiced or of being carried out in various 65 ways. Also, the phraseology and terminology used herein is for the purpose of description and should not be regarded as

208

limiting. The use of "including," "comprising," or "having," "containing," "involving," and variations thereof herein, is meant to encompass the items listed thereafter and equivalents thereof as well as additional items.

#### **EXAMPLES**

#### Example 1: Manufacture of Polynucleotides

According to the present disclosure, the manufacture of polynucleotides and/or parts or regions thereof may be accomplished utilizing the methods taught in International Publication WO2014/152027, entitled "Manufacturing Methods for Production of RNA Transcripts," the contents of which is incorporated herein by reference in its entirety.

Purification methods may include those taught in International Publication WO2014/152030 and International Publication WO2014/152031, each of which is incorporated herein by reference in its entirety.

Detection and characterization methods of the polynucleotides may be performed as taught in International Publication WO2014/144039, which is incorporated herein by ref-

Characterization of the polynucleotides of the disclosure 4-methoxy-pseudouridine, 4-thio-1-methyl-pseudouridine, 25 may be accomplished using polynucleotide mapping, reverse transcriptase sequencing, charge distribution analysis, detection of RNA impurities, or any combination of two or more of the foregoing. "Characterizing" comprises determining the RNA transcript sequence, determining the purity of the RNA transcript, or determining the charge heterogeneity of the RNA transcript, for example. Such methods are taught in, for example, International Publication WO2014/ 144711 and International Publication WO2014/144767, the content of each of which is incorporated herein by reference 35 in its entirety.

#### Example 2: Chimeric Polynucleotide Synthesis

According to the present disclosure, two regions or parts triphosphate chemistry. A first region or part of 100 nucleotides or less is chemically synthesized with a 5' monophosphate and terminal 3'desOH or blocked OH, for example. If the region is longer than 80 nucleotides, it may be synthesized as two strands for ligation.

If the first region or part is synthesized as a non-positionally modified region or part using in vitro transcription (IVT), conversion the 5'monophosphate with subsequent capping of the 3' terminus may follow.

Monophosphate protecting groups may be selected from any of those known in the art.

The second region or part of the chimeric polynucleotide may be synthesized using either chemical synthesis or IVT methods. IVT methods may include an RNA polymerase that can utilize a primer with a modified cap. Alternatively, a cap of up to 130 nucleotides may be chemically synthesized and coupled to the IVT region or part.

For ligation methods, ligation with DNA T4 ligase, followed by treatment with DNase should readily avoid con-60 catenation.

The entire chimeric polynucleotide need not be manufactured with a phosphate-sugar backbone. If one of the regions or parts encodes a polypeptide, then such region or part may comprise a phosphate-sugar backbone.

Ligation is then performed using any known click chemistry, orthoclick chemistry, solulink, or other bioconjugate chemistries known to those in the art.

Synthetic Route

The chimeric polynucleotide may be made using a series of starting segments. Such segments include:

- (a) a capped and protected 5' segment comprising a normal 3'OH (SEG. 1)
- (b) a 5' triphosphate segment, which may include the coding region of a polypeptide and a normal 3'OH (SEG. 2)  $\,$
- (c) a 5' monophosphate segment for the 3' end of the chimeric polynucleotide (e.g., the tail) comprising cordycepin or no 3'OH (SEG. 3)

After synthesis (chemical or IVT), segment 3 (SEG. 3) may be treated with cordycepin and then with pyrophosphatase to create the 5' monophosphate.

Segment 2 (SEG. 2) may then be ligated to SEG. 3 using RNA ligase. The ligated polynucleotide is then purified and 15 treated with pyrophosphatase to cleave the diphosphate. The treated SEG. 2-SEG. 3 construct may then be purified and SEG. 1 is ligated to the 5' terminus. A further purification step of the chimeric polynucleotide may be performed.

Where the chimeric polynucleotide encodes a polypeptide, the ligated or joined segments may be represented as: 5'UTR (SEG. 1), open reading frame or ORF (SEG. 2) and 3'UTR+PolyA (SEG. 3).

The yields of each step may be as much as 90-95%.

#### Example 3: PCR for cDNA Production

PCR procedures for the preparation of cDNA may be performed using  $2\times KAPA$  HIFITM HotStart ReadyMix by Kapa Biosystems (Woburn, Mass.). This system includes  $_{30}$  2×KAPA ReadyMix 12.5  $\mu l;$  Forward Primer (10  $\mu M)$  0.75  $\mu l;$  Reverse Primer (10  $\mu M)$  0.75  $\mu l;$  Template cDNA 100 ng; and dH $_2O$  diluted to 25.0  $\mu l.$  The reaction conditions may be at 95° C. for 5 min. The reaction may be performed for 25 cycles of 98° C. for 20 sec, then 58° C. for 15 sec, then 72°  $_{35}$  C. for 45 sec, then 72° C. for 5 min, then 4° C. to termination.

The reaction may be cleaned up using Invitrogen's PURELINK<sup>TM</sup> PCR Micro Kit (Carlsbad, Calif.) per manufacturer's instructions (up to 5  $\mu g$ ). Larger reactions may  $_{40}$  require a cleanup using a product with a larger capacity. Following the cleanup, the cDNA may be quantified using the NANODROPTM and analyzed by agarose gel electrophoresis to confirm that the cDNA is the expected size. The cDNA may then be submitted for sequencing analysis before  $_{45}$  proceeding to the in vitro transcription reaction.

### Example 4: In Vitro Transcription (IVT)

The in vitro transcription reaction generates RNA polynucleotides. Such polynucleotides may comprise a region or part of the polynucleotides of the disclosure, including chemically modified RNA (e.g., mRNA) polynucleotides. The chemically modified RNA polynucleotides can be uniformly modified polynucleotides. The in vitro transcription reaction utilizes a custom mix of nucleotide triphosphates (NTPs). The NTPs may comprise chemically modified NTPs, or a mix of natural and chemically modified NTPs, or natural NTPs.

A typical in vitro transcription reaction includes the  $_{60}$  following:

1.0 μg

# 210 -continued

| (i | Custom NTPs (25 mM each) | 0.2           | μl    |
|----|--------------------------|---------------|-------|
| ŀ) | RNase Inhibitor          | 20            | U     |
| i) | T7 RNA polymerase        | 3000          | U     |
| (( | $dH_20$                  | up to 20.0 ul | . and |

Incubation at 37° C. for 3 hr-5 hrs.

The crude IVT mix may be stored at 4° C. overnight for cleanup the next day. 1 U of RNase-free DNase may then be used to digest the original template. After 15 minutes of incubation at 37° C., the mRNA may be purified using Ambion's MEGACLEAR™ Kit (Austin, Tex.) following the manufacturer's instructions. This kit can purify up to 500 μg of RNA. Following the cleanup, the RNA polynucleotide may be quantified using the NanoDrop and analyzed by agarose gel electrophoresis to confirm the RNA polynucleotide is the proper size and that no degradation of the RNA has occurred.

#### Example 5: Enzymatic Capping

Capping of a RNA polynucleotide is performed as follows where the mixture includes: IVT RNA 60  $\mu$ g-180  $\mu$ g and <sup>25</sup> dH<sub>2</sub>O up to 72  $\mu$ l. The mixture is incubated at 65° C. for 5 minutes to denature RNA, and then is transferred immediately to ice.

The protocol then involves the mixing of 10x Capping Buffer (0.5 M Tris-HCl (pH 8.0), 60 mM KCl, 12.5 mM MgCl<sub>2</sub>) (10.0  $\mu$ l); 20 mM GTP (5.0  $\mu$ l); 20 mM S-Adenosyl Methionine (2.5  $\mu$ l); RNase Inhibitor (100 U); 2'-O-Methyltransferase (400U); Vaccinia capping enzyme (Guanylyl transferase) (40 U); dH<sub>2</sub>O (Up to 28  $\mu$ l); and incubation at 37° C. for 30 minutes for 60  $\mu$ g RNA or up to 2 hours for 180  $\mu$ g of RNA.

The RNA polynucleotide may then be purified using Ambion's MEGACLEAR<sup>TM</sup> Kit (Austin, Tex.) following the manufacturer's instructions. Following the cleanup, the RNA may be quantified using the NANODROP<sup>TM</sup> (ThermoFisher, Waltham, Mass.) and analyzed by agarose gel electrophoresis to confirm the RNA polynucleotide is the proper size and that no degradation of the RNA has occurred. The RNA polynucleotide product may also be sequenced by running a reverse-transcription-PCR to generate the cDNA for sequencing.

### Example 6: PolyA Tailing Reaction

Without a poly-T in the cDNA, a poly-A tailing reaction must be performed before cleaning the final product. This is done by mixing capped IVT RNA (100  $\mu$ l); RNase Inhibitor (20 U);  $10\times$  Tailing Buffer (0.5 M Tris-HCl (pH 8.0), 2.5 M NaCl, 100 mM MgCl $_2$ ) (12.0  $\mu$ l); 20 mM ATP (6.0  $\mu$ l); Poly-A Polymerase (20 U); dH $_2$ O up to 123.5  $\mu$ l and incubation at 37° C. for 30 min. If the poly-A tail is already in the transcript, then the tailing reaction may be skipped and proceed directly to cleanup with Ambion's MEGA-CLEARTM kit (Austin, Tex.) (up to 500  $\mu$ g). Poly-A Polymerase may be a recombinant enzyme expressed in yeast.

It should be understood that the processivity or integrity of the polyA tailing reaction may not always result in an exact size polyA tail. Hence, polyA tails of approximately between 40-200 nucleotides, e.g., about 40, 50, 60, 70, 80, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 150-165, 155, 156, 157,

Template cDNA

<sup>2) 10</sup>x transcription buffer (400 mM Tris-HCl pH 8.0, 190 mM MgCl<sub>2</sub>, 50 mM DTT, 10 mM Spermidine)

158, 159, 160, 161, 162, 163, 164 or 165 are within the scope of the present disclosure.

#### Example 7. Natural 5' Caps and 5' Cap Analogues

5'-capping of polynucleotides may be completed concomitantly during the in vitro-transcription reaction using the following chemical RNA cap analogs to generate the 5'-guanosine cap structure according to manufacturer protocols: 3'-O-Me-m7G(5')ppp(5') G [the ARCA cap]; G(5') 10 ppp(5')A; G(5')ppp(5')G; m7G(5')ppp(5')A; m7G(5')ppp (5')G (New England BioLabs, Ipswich, Mass.). 5'-capping of modified RNA may be completed post-transcriptionally using a Vaccinia Virus Capping Enzyme to generate the "Cap 0" structure: m7G(5')ppp(5')G (New England Bio-Labs, Ipswich, Mass.). Cap 1 structure may be generated using both Vaccinia Virus Capping Enzyme and a 2'-O methyl-transferase to generate: m7G(5')ppp(5')G-2'-Omethyl. Cap 2 structure may be generated from the Cap 1 structure followed by the 2'-O-methylation of the 5'-ante-20 penultimate nucleotide using a 2'-O methyl-transferase. Cap 3 structure may be generated from the Cap 2 structure followed by the 2'-O-methylation of the 5'-preantepenultimate nucleotide using a 2'-O methyl-transferase. Enzymes are preferably derived from a recombinant source.

When transfected into mammalian cells, the modified mRNAs have a stability of between 12-18 hours or more than 18 hours, e.g., 24, 36, 48, 60, 72 or greater than 72 hours.

# Example 8: Capping Assays

Protein Expression Assay

Polynucleotides (e.g., mRNA) encoding a polypeptide, containing any of the caps taught herein, can be transfected 35 into cells at equal concentrations. The amount of protein secreted into the culture medium can be assayed by ELISA at 6, 12, 24 and/or 36 hours post-transfection. Synthetic polynucleotides that secrete higher levels of protein into the medium correspond to a synthetic polynucleotide with a 40 higher translationally-competent cap structure.

Purity Analysis Synthesis

RNA (e.g., mRNA) polynucleotides encoding a polypeptide, containing any of the caps taught herein can be compared for purity using denaturing Agarose-Urea gel electrophoresis or HPLC analysis. RNA polynucleotides with a single, consolidated band by electrophoresis correspond to the higher purity product compared to polynucleotides with multiple bands or streaking bands. Chemically modified RNA polynucleotides with a single HPLC peak also correspond to a higher purity product. The capping reaction with a higher efficiency provides a more pure polynucleotide population.

Cytokine Analysis

RNA (e.g., mRNA) polynucleotides encoding a polypeptide, containing any of the caps taught herein can be transfected into cells at multiple concentrations. The amount of pro-inflammatory cytokines, such as TNF-alpha and IFN-beta, secreted into the culture medium can be assayed by ELISA at 6, 12, 24 and/or 36 hours post-transfection. RNA 60 polynucleotides resulting in the secretion of higher levels of pro-inflammatory cytokines into the medium correspond to a polynucleotides containing an immune-activating cap structure.

Capping Reaction Efficiency

RNA (e.g., mRNA) polynucleotides encoding a polypeptide, containing any of the caps taught herein can be ana-

212

lyzed for capping reaction efficiency by LC-MS after nuclease treatment. Nuclease treatment of capped polynucleotides yield a mixture of free nucleotides and the capped 5'-5-triphosphate cap structure detectable by LC-MS. The amount of capped product on the LC-MS spectra can be expressed as a percent of total polynucleotide from the reaction and correspond to capping reaction efficiency. The cap structure with a higher capping reaction efficiency has a higher amount of capped product by LC-MS.

# Example 9: Agarose Gel Electrophoresis of Modified RNA or RT PCR Products

Individual RNA polynucleotides (200-400 ng in a 20 µl volume) or reverse transcribed PCR products (200-400 ng) may be loaded into a well on a non-denaturing 1.2% Agarose E-Gel (Invitrogen, Carlsbad, Calif.) and run for 12-15 minutes, according to the manufacturer protocol.

# Example 10: Nanodrop Modified RNA Quantification and UV Spectral Data

Chemically modified RNA polynucleotides in TE buffer  $(1 \mu l)$  are used for Nanodrop UV absorbance readings to quantitate the yield of each polynucleotide from an chemical synthesis or in vitro transcription reaction.

# Example 11: Formulation of Modified mRNA Using Lipidoids

RNA (e.g., mRNA) polynucleotides may be formulated for in vitro experiments by mixing the polynucleotides with the lipidoid at a set ratio prior to addition to cells. In vivo formulation may require the addition of extra ingredients to facilitate circulation throughout the body. To test the ability of these lipidoids to form particles suitable for in vivo work, a standard formulation process used for siRNA-lipidoid formulations may be used as a starting point. After formation of the particle, polynucleotide is added and allowed to integrate with the complex. The encapsulation efficiency is determined using a standard dye exclusion assays.

#### Example 12: Immunogenicity Study

The instant study is designed to test the immunogenicity in mice of candidate hMPV vaccines comprising a mRNA polynucleotide encoding Fusion (F) glycoprotein, major surface glycoprotein G, or a combination thereof, obtained from hMPV.

Mice are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) with candidate vaccines. Candidate vaccines are chemically modified or unmodified. A total of four immunizations are given at 3-week intervals (i.e., at weeks 0, 3, 6, and 9), and sera are collected after each immunization until weeks 33-51. Serum antibody titers against Fusion (F) glycoprotein or major surface glycoprotein (G) protein are determined by ELISA. Sera collected from each mouse during weeks 10-16 are pooled, and total IgG purified. Purified antibodies are used for immunoelectron microscopy, antibody-affinity testing, and in vitro protection assays.

#### Example 13: hMPV Rodent Challenge

The instant study is designed to test the efficacy in cotton rats of candidate hMPV vaccines against a lethal challenge using an hMPV vaccine comprising mRNA encoding Fusion

(F) glycoprotein, major surface glycoprotein G, or a combination of both antigens obtained from hMPV. Cotton rats are challenged with a lethal dose of the hMPV.

Animals are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) at week 0 and week 3 with 5 candidate hMPV vaccines with and without adjuvant. Candidate vaccines are chemically modified or unmodified. The animals are then challenged with a lethal dose of hMPV on week 7 via IV, IM or ID. Endpoint is day 13 post infection, death or euthanasia. Animals displaying severe illness as 10 determined by >30% weight loss, extreme lethargy or paralysis are euthanized. Body temperature and weight are assessed and recorded daily.

In experiments where a lipid nanoparticle (LNP) formulation is used, the formulation may include a cationic lipid, non-cationic lipid, PEG lipid and structural lipid in the ratios 50:10:1.5:38.5. The cationic lipid is DLin-KC2-DMA (50 mol %) or DLin-MC3-DMA (50 mol %), the non-cationic lipid is DSPC (10 mol %), the PEG lipid is PEG-DOMG (1.5 mol %) and the structural lipid is cholesterol (38.5 mol %), <sup>20</sup> for example.

#### Example 14: Immunogenicity of hMPV mRNA Vaccine in BALB/c Mice

The instant study was designed to test the immunogenicity in BALB/c mice of hMPV vaccines comprising an mRNA polynucleotide encoding the hMPV Fusion (F) glycoprotein. The mRNA polynucleotide encodes the fulllength fusion protein and comprises the wild-type nucleotide 30 sequence obtained from the hMPV A2a strain. Mice were divided into 3 groups (n=8 for each group) and immunized intramuscularly (IM) with PBS, a 10 µg dose of mRNA vaccines encoding hMPV fusion protein, or a 2 µg dose of mRNA vaccines encoding hMPV fusion protein. A total of 35 two immunizations were given at 3-week intervals (i.e., at weeks 0, and 3 weeks), and sera were collected after each immunization according to the schedule described in Table 1. Serum antibody titers against hMPV fusion glycoprotein were determined by ELISA and antibodies were detected in 40 the sera collected on day 14 onward. Both vaccine doses tested induced comparable levels of immune response in mice (FIGS. 2A-2C).

Additionally, mice sera were used for IgG isotyping (FIGS. 3A-3C). Both hMPV fusion protein-specific IgG1 45 and IgG2a were detected in mice sera. hMPV fusion protein mRNA vaccine also induced Th1 and Th2 cytokine responses, with a Th1 bias.

Sera from mice immunized with either  $10\,\mu g$  or  $2\,\mu g$  doses of the hMPV fusion protein mRNA vaccine contain neutralizing antibodies. The ability of these antibodies to neutralize hMPV B2 strain was also tested. The antibody-containing sera successfully neutralized the hMPV B2 virus (FIG. 4).

# Example 15: T-Cell Stimulation

The instant study was designed to test T-cell stimulation in the splenocytes of mice immunized with mRNA vaccines encoding hMPV fusion protein, as described herein. Immunization of BALB/c mice was performed as described in 60 Example 14. The splenocytes for each group were pooled and split into two parts. One part of splenocytes from each group of mice was stimulated with hMPV-free media, Concanavalin A or a hMPV fusion protein peptide pool comprising 15-mers (15 amino acids long); while the other part of splenocytes from each group of mice was stimulated with hMPV-free media, Concanavalin A or inactivated hMPV

214

virus. Secreted mouse cytokines were measured using the Meso Scale Discovery (MSD) assay.

Cytokines specific to Th1 or Th2 responses were measured. For Th1 response, IFN- $\gamma$ , IL2 and IL12 were detected from splenocytes stimulated with the hMPV fusion protein peptide pool at a level comparable to that of Concanavalin A (FIGS. 5A-5C). For a Th2 response, the hMPV fusion protein peptide pool induced the secretion of detectable IL10, TNF- $\alpha$ , IL4 and IL, but not IL5, while Concanavalin A stimulated the secretion of all the above-mentioned Th2 cytokines (FIGS. 6A-6E) at a much higher level.

In contrast, inactivated hMPV virus only induced the secretion of IL2 in the Th1 response comparable to that of Concanavalin A (FIGS. 7A-7C). For the Th2 response, the inactivated hMPV virus induced the secretion of detectable IL10, TNF-α, IL4 and IL6, but not IL5, while Concanavalin A stimulated the secretion of all the above-mentioned Th2 cytokines (FIGS. 8A-8E) at a much higher level.

### Example 16: hMPV Rodent Challenge in Cotton Rats Immunized with mRNA Vaccine Encoding hMPV Fusion Protein

The instant study was designed to test the efficacy in cotton rats of hMPV vaccines against a lethal challenge. mRNA vaccines encoding hMPV fusion protein were used. The mRNA polynucleotide encodes a full-length fusion protein and comprises the wild-type nucleotide sequence obtained from the hMPV A2a strain.

Cotton rats were immunized intramuscularly (IM) at week 0 and week 3 with the mRNA vaccines encoding hMPV fusion protein with either 2  $\mu g$  or 10  $\mu g$  doses for each immunization. The animals were then challenged with a lethal dose of hMPV in week 7 post initial immunization via IV, IM or ID. The endpoint was day 13 post infection, death or euthanasia. Viral titers in the noses and lungs of the cotton rats were measured. The results (FIGS. 9A and 9B) show that a 10  $\mu g$  dose of mRNA vaccine protected the cotton mice 100% in the lung and drastically reduced the viral titer in the nose after challenge (~2 log reduction). Moreover, a 2  $\mu g$  dose of mRNA vaccine showed a 1 log reduction in lung viral titer in the cotton mice challenged.

Further, the histopathology of the lungs of the cotton mice immunized and challenged showed no pathology associated with vaccine-enhanced disease (FIG. 10).

# Example 17. Immunogenicity Study

The instant study is designed to test the immunogenicity in mice of candidate PIV3 vaccines comprising a mRNA polynucleotide encoding hemagglutinin-neuraminidase or fusion protein (F or F0) obtained from PIV3.

Mice are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) with candidate vaccines. Candidate vaccines are chemically modified or unmodified. A total of four immunizations are given at 3-week intervals (i.e., at weeks 0, 3, 6, and 9), and sera are collected after each immunization until weeks 33-51. Serum antibody titers against hemagglutinin-neuraminidase or fusion protein (F or F0) are determined by ELISA. Sera collected from each mouse during weeks 10-16 are, optionally, pooled, and total IgGs are purified. Purified antibodies are used for immunoelectron microscopy, antibody-affinity testing, and in vitro protection assays.

#### Example 18: PIV3 Rodent Challenge

The instant study is designed to test the efficacy in cotton rats of candidate PIV3 vaccines against a lethal challenge

using a PIV3 vaccine comprising mRNA encoding hemagglutinin-neuraminidase or fusion protein (F or F0) obtained from PIV3. Cotton rats are challenged with a lethal dose of the PIV3.

Animals are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) at week 0 and week 3 with candidate PIV3 vaccines with and without adjuvant. Candidate vaccines are chemically modified or unmodified. The animals are then challenged with a lethal dose of PIV3 on week 7 via IV, IM or ID. Endpoint is day 13 post infection, death or euthanasia. Animals displaying severe illness as determined by >30% weight loss, extreme lethargy or paralysis are euthanized. Body temperature and weight are assessed and recorded daily.

In experiments where a lipid nanoparticle (LNP) formulation is used, the formulation may include a cationic lipid, non-cationic lipid, PEG lipid and structural lipid in the ratios 50:10:1.5:38.5. The cationic lipid is DLin-KC2-DMA (50 mol %) or DLin-MC3-DMA (50 mol %), the non-cationic 20 lipid is DSPC (10 mol %), the PEG lipid is PEG-DOMG (1.5 mol %) and the structural lipid is cholesterol (38.5 mol %), for example.

#### Example 19: hMPV/PIV Cotton Rat Challenge

The instant study was designed to test the efficacy in cotton rats of candidate hMPV mRNA vaccines, PIV3 mRNA vaccines, or hMPV/PIV combination mRNA vaccines against a lethal challenge using PIV3 strain or hMPV/ 30 A2 strain. The study design is shown in Table 9.

Cotton rats of 10-12 weeks old were divided into 12 groups (n=5), and each group was vaccinated with mRNA vaccines indicated in Table 9. The PIV3 vaccine comprises mRNA encoding hemagglutinin-neuraminidase or fusion protein (F or F0) obtained from PIV3. The hMPV mRNA vaccine encodes the full-length hMPV fusion protein. The hMPV/PIV combination mRNA vaccine is a mixture of the PIV3 vaccine and hMPV vaccine at a 1:1 ratio.

Cotton rats were immunized intramuscularly (IM) at week 0 and week 3 with candidate vaccines with the doses indicated in Table 9. Cotton rats immunized with hMPV mRNA vaccines or hMPV/PIV combination mRNA vaccines were challenged with a lethal dose of hMPV/A2 strain 45 on week 7 via IM. Cotton rats immunized with PIV mRNA vaccines or hMPV/PIV combination mRNA vaccines were challenged with a lethal dose of PIV3 strain on week 7 via IM

The endpoint was day 13 post infection, death or euthanasia. Animals displaying severe illness as determined by >30% weight loss, extreme lethargy or paralysis were euthanized. Body temperature and weight were assessed and recorded daily.

Lung and nose hMPV/A2 (FIG. 12) or PIV3 (FIG. 13) 55 viral titers were assessed. Lung histopathology of the immunized and challenged cotton rat immunized and challenged were assessed to determine pathology associated with vaccine enhance disease. Neutralization antibody titers in the serum of immunized cotton rats on day 0 and 42 post 60 immunization were assessed (FIG. 11).

hMPV/A2 (FIG. 14) or PIV3 (FIG. 15) neutralizing antibody titers in the serum samples of the immunized cotton rat 42 days post immunization were measured. All mRNA vaccines tested induced strong neutralizing antibodies cotton rats. Lung histopathology of the immunized cotton rats were also evaluated (FIG. 16). Low occurrence of

216

alevolitis and interstitial pneumonia was observed, indicating no antibody-dependent enhancement (ADE) of hMPV or PIV associated diseases.

# Example 20: Betacoronavirus Immunogenicity Study

The instant study is designed to test the immunogenicity in rabbits of candidate *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1 or a combination thereof) vaccines comprising a mRNA polynucleotide encoding the spike (S) protein, the S1 subunit (S1) of the spike protein, or the S2 subunit (S2) of the spike protein obtained from a *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1).

Rabbits are vaccinated on week 0 and 3 via intravenous (IV), intramuscular (IM), or intradermal (ID) routes. One group remains unvaccinated and one is administered inactivated *Betacoronavirus*. Serum is collected from each rabbit on weeks 1, 3 (pre-dose) and 5. Individual bleeds are tested for anti-S, anti-S1 or anti-S2 activity via a virus neutralization assay from all three time points, and pooled samples from week 5 only are tested by Western blot using inactivated *Betacoronavirus* (e.g., inactivated MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1).

In experiments where a lipid nanoparticle (LNP) formulation is used, the formulation may include a cationic lipid, non-cationic lipid, PEG lipid and structural lipid in the ratios 50:10:1.5:38.5. The cationic lipid is DLin-KC2-DMA (50 mol %) or DLin-MC3-DMA (50 mol %), the non-cationic lipid is DSPC (10 mol %), the PEG lipid is PEG-DOMG (1.5 mol %) and the structural lipid is cholesterol (38.5 mol %), for example.

#### Example 21: Betacoronavirus Challenge

The instant study is designed to test the efficacy in rabbits of candidate *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-HKU1 or a combination thereof) vaccines against a lethal challenge using a *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-HKU1 or a combination thereof) vaccine comprising mRNA encoding the spike (S) protein, the S1 subunit (S1) of the spike protein, or the S2 subunit (S2) of the spike protein obtained from *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-1KU1). Rabbits are challenged with a lethal dose (10×LD90; ~100 plaque-forming units; PFU) of *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL or HCoV-HKU1).

The animals used are 6-8 week old female rabbits in groups of 10. Rabbits are vaccinated on weeks 0 and 3 via an IM, ID or IV route of administration. Candidate vaccines are chemically modified or unmodified. Rabbit serum is tested for microneutralization (see Example 14). Rabbits are then challenged with ~1 LD90 of *Betacoronavirus* (e.g., MERS-CoV, SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1) on week 7 via an IN, IM, ID or IV route of administration. Endpoint is day 13 post infection, death or euthanasia. Animals displaying severe illness as determined by >30%

217

weight loss, extreme lethargy or paralysis are euthanized. Body temperature and weight are assessed and recorded daily.

#### Example 22: Microneutralization Assay

Nine serial 2-fold dilutions (1:50-1:12,800) of rabbit serum are made in 50 µl virus growth medium (VGM) with trypsin in 96 well microtiter plates. Fifty microliters of virus containing ~50 pfu of Betacoronavirus (e.g., MERS-CoV, 10 SARS-CoV, HCoV-OC43, HCoV-229E, HCoV-NL63, HCoV-NL, HCoV-NH or HCoV-HKU1) is added to the serum dilutions and allowed to incubate for 60 minutes at room temperature (RT). Positive control wells of virus without sera and negative control wells without virus or sera are included in triplicate on each plate. While the serumvirus mixtures incubate, a single cell suspension of Madin-Darby Canine-Kidney cells are prepared by trypsinizing (Gibco 0.5% bovine pancrease trypsin in EDTA) a confluent monolayer and suspended cells are transferred to a 50 ml 20 centrifuge tube, topped with sterile PBS and gently mixed. The cells are then pelleted at 200 g for 5 minutes, supernatant aspirated and cells resuspended in PBS. This procedure is repeated once and the cells are resuspended at a concentration of 3×10<sup>5</sup>/ml in VGM with porcine trypsin. Then, 100 <sup>25</sup> µl of cells are added to the serum-virus mixtures and the plates incubated at 35° C. in CO<sub>2</sub> for 5 days. The plates are fixed with 80% acetone in phosphate buffered saline (PBS) for 15 minutes at RT, air dried and then blocked for 30 minutes containing PBS with 0.5% gelatin and 2% FCS. An antibody to the S proteins, 51 protein or S2 protein is diluted in PBS with 0.5% gelatin/2% FCS/0.5% Tween 20 and incubated at RT for 2 hours. Wells are washed and horseradish peroxidase-conjugated goat anti-mouse IgG added, followed by another 2 hour incubation. After washing, 0-phenylenediamine dihydrochloride is added and the neutralization titer is defined as the titer of serum that reduced color development by 50% compared to the positive control wells.

# Example 23: MERS CoV Vaccine Immunogenicity Study in Mice

The instant study was designed to test the immunogenicity in mice of candidate MERS-CoV vaccines comprising a 45 mRNA polynucleotide encoding the full-length Spike (S) protein, or the S2 subunit (S2) of the Spike protein obtained from MERS-CoV.

Mice were vaccinated with a 10  $\mu g$  dose of MERS-CoV mRNA vaccine encoding either the full-length MERS-CoV 50 Spike (S) protein, or the S2 subunit (S2) of the Spike protein on days 0 and 21. Sera were collected from each mice on days 0, 21, 42, and 56. Individual bleeds were tested for anti-S, anti-S2 activity via a virus neutralization assay from all four time points.

As shown in FIG. 17, the MERS-CoV vaccine encoding the full-length S protein induced strong immune response after the boost dose on day 21. Further, full-length S protein vaccine generated much higher neutralizing antibody titers as compared to S2 alone (FIG. 18).

#### Example 24: MERS CoV Vaccine Immunogenicity Study in New Zealand White Rabbits

The instant study was designed to test the immunogenicity of candidate MERS-CoV mRNA vaccines encoding the full-length Spike (S) protein. The New Zealand white rabbits 218

used in this study weighed about 4-5 kg. The rabbits were divided into three groups (Group 1a, Group 1b, and Group 2, n=8). Rabbits in Group 1a were immunized intramuscularly (IM) with one 20 µg dose of the MERS-CoV mRNA vaccine encoding the full-length Spike protein on day 0. Rabbits in Group 1b were immunized intramuscularly (IM) with one 20 µg dose of the MERS-CoV mRNA vaccine encoding the full-length Spike protein on day 0, and again on day 21 (booster dose). Group 2 received placebo (PBS). The immunized rabbits were then challenged and samples were collected 4 days after challenge. The viral loads in the lungs, bronchoalveolar lavage (Bal), nose, and throat of the rabbits were determined, e.g., via quantitative PCR. Replicating virus in the lung tissues of the rabbits were also detected. Lung histopathology were evaluated and the neutralizing antibody titers in serum samples of the rabbits were determined.

Two 20 µg doses of MERS-CoV mRNA vaccine resulted in a 3 log reduction of viral load in the nose and led to complete protection in the throat of the New Zealand white rabbits (FIG. 19A). Two 20 µg doses of MERS-CoV mRNA vaccine also resulted in a 4 log reduction of viral load in the BAL of the New Zealand white rabbits (FIG. 19B). One 20 µg dose of MERS-CoV mRNA vaccine resulted in a 2 log reduction of viral load, while two 20 µg doses of MERS-CoV mRNA vaccine resulted in an over 4 log reduction of viral load in the lungs of the New Zealand white rabbits (FIG. 19C).

Quantitative PCR results show that two 20 µg doses of MERS-CoV mRNA vaccine reduced over 99% (2 log) of viruses in the lungs of New Zealand white rabbits (FIG. 20A). No replicating virus were detected in the lungs (FIG. 20B).

Further, as shown in FIG. **21**, two **20**  $\mu$ g doses of MERS-CoV mRNA vaccine induced significant amount of neutralizing antibodies against MERS-CoV (EC<sub>50</sub> between 500-1000). The MERS-CoV mRNA vaccine induced antibody titer is 3-5 fold better than any other vaccines tested in the same model.

### Example 25: Immunogenicity Study

The instant study is designed to test the immunogenicity in mice of candidate MeV vaccines comprising a mRNA polynucleotide encoding MeV hemagglutinin (HA) protein, MeV Fusion (F) protein or a combination of both.

Mice are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) with candidate vaccines. Up to three immunizations are given at 3-week intervals (i.e., at weeks 0, 3, 6, and 9), and sera are collected after each immunization until weeks 33-51. Serum antibody titers against MeV HA protein or MeV F protein are determined by ELISA.

# Example 26: MeV Rodent Challenge

The instant study is designed to test the efficacy in transgenic mice of candidate MeV vaccines against a lethal challenge using a MeV vaccine comprising mRNA encoding MeV HA protein or MeV F protein. The transgenic mice express human receptor CD46 or signaling lymphocyte activation molecule (SLAM) (also referred to as CD150). Humans are the only natural host for MeV infection, thus transgenic lines are required for this study. CD46 is a complement regulatory protein that protects host tissue from complement deposition by binding to complement components C3b and C4b. Its expression on murine fibroblast and

lymphoid cell lines renders these otherwise refractory cells permissive for MeV infection, and the expression of CD46 on primate cells parallels the clinical tropism of MeV infection in humans and nonhuman primates (Rall G F et al. *PNAS USA* 1997; 94(9):4659-63). SLAM is a type 1 membrane glycoprotein belonging to the immunoglobulin superfamily. It is expressed on the surface of activated lymphocytes, macrophages, and dendritic cells and is thought to play an important role in lymphocyte signaling. SLAM is a receptor for both wild-type and vaccine MeV strains (Sellin C I et al. *J Virol.* 2006; 80(13):6420-29).

CD46 or SLAM/CD150 transgenic mice are challenged with a lethal dose of the MeV. Animals are immunized intravenously (IV), intramuscularly (IM), or intradermally (ID) at week 0 and week 3 with candidate MeV vaccines

with and without adjuvant. The animals are then challenged with a lethal dose of MeV on week 7 via IV, IM or ID. Endpoint is day 13 post infection, death or euthanasia. Animals displaying severe illness as determined by >30% weight loss, extreme lethargy or paralysis are euthanized. Body temperature and weight are assessed and recorded daily.

220

In experiments where a lipid nanoparticle (LNP) formulation is used, the formulation may include a cationic lipid, non-cationic lipid, PEG lipid and structural lipid in the ratios 50:10:1.5:38.5. The cationic lipid is DLin-KC2-DMA (50 mol %), the non-cationic lipid is DSPC (10 mol %), the PEG lipid is PEG-DOMG (1.5 mol %) and the structural lipid is cholesterol (38.5 mol %), for example.

TABLE 1

|                                 | hMPV Immunogenicity studies bleeding schedule                  |  |               |       |        |        |                  |        |        |                                     |
|---------------------------------|--|--|---------------|-------|--------|--------|------------------|--------|--------|-------------------------------------|
|                                 | Animal<br>groups   |  |               | Day   |        |        |                  |        |        |                                     |
|                                 | (n = 8)  | vaccine                                      | -2            | 0     | 7      | 14     | 21               | 28     | 35     | 56                                  |
| Placebo  10 µg  Dose 2 µg  Dose | Group 1<br>(n = 8)<br>Group 2<br>(n = 8)<br>Group 3<br>(n = 8) | PBS<br>(IM)<br>10 µg<br>(IM)<br>2 µg<br>(IM) | Pre-<br>Bleed | Prime | Bleeds | Bleeds | Bleeds/<br>Boost | Bleeds | Bleeds | Harvest Spleens/<br>Terminal Bleeds |

Total n = 24

Each of the sequences described herein encompasses a chemically modified sequence or an unmodified sequence which includes no nucleotide modifications.

TABLE 2

|             |          | SEQ |
|-------------|----------|-----|
|             |          | ID  |
| Description | Sequence | NO: |

hMPV Nucleic Acid Sequences

gi|122891979|gb| EF051124.1| Human metapneumo virus isolate TN/92-4 fusion protein gene, complete genome ATGAGCTGGAAGGTGGTGATTATCTTCAGCCTGCTGATTA CACCTCAACACGGCCTGAAGGAGGAGCTACCTGGAAGAGA GCTGCTCCACCATCACCGAGGGCTACCTGAGCGTGCTGC GGACCGGCTGGTACACCAACGTGTTCACCCTGGAGGTGG GCGACGTGGAGAACCTGACCTGCAGCGACGGCCCTAGCC TGATCAAGACCGAGCTGGACCTGACCAAGAGCGCTCTGA GAGAGCTGAAGACCGTGTCCGCCGACCAGCTGGCCAGAG AGGAACAGATCGAGAACCCTCGGCAGAGCAGATTCGTGC  $\tt TGGGCGCCATCGCTCTGGGAGTCGCCGCTGCCGCTGCAG$ TGACAGCTGGAGTGGCCATTGCTAAGACCATCAGACTGG  ${\tt AAAGCGAGGTGACAGCCATCAACAATGCCCTGAAGAAG}$ ACCAACGAGGCCGTGAGCACCCTGGGCAATGGAGTGAGA  $\tt GTGCTGGCCACAGCCGTGCGGGAGCTGAAGGACTTCGTG$ AGCAAGAACCTGACCAGAGCCATCAACAAGAACAAGTG CGACATCGATGACCTGAAGATGGCCGTGAGCTTCTCCCA  $\tt GTTCAACAGACGGTTCCTGAACGTGGTGAGACAGTTCTC$ CGACAACGCTGGAATCACACCTGCCATTAGCCTGGACCT GATGACCGACCCGAGCTGGCTAGAGCCGTGCCCAACAT GCCCACCAGCGCTGGCCAGATCAAGCTGATGCTGGAGAA  ${\tt CAGAGCCATGGTGCGGAGAAAGGGCTTCGGCATCCTGAT}$ TGGGGTGTATGGAAGCTCCGTGATCTACATGGTGCAGCT GCCCATCTTCGGCGTGATCGACACCCCTGCTGGATCGTG AAGGCCGCTCCTAGCTGCTCCGAGAAGAAAGGAAACTAT GCCTGTCTGCTGAGAGAGGACCAGGGCTGGTACTGCCAG AACGCCGGAAGCACAGTGTACTATCCCAACGAGAAGGAC  ${\tt TGCGAGACCAGAGGCGACCACGTGTTCTGCGACACCGCT}$ GCCGGAATCAACGTGGCCGAGCAGGAGCAAGGAGTGCAA CATCAACATCAGCACAACCAACTACCCCTGCAAGGTGAG CACCGGACGCCACCCCATCAGCATGGTGGCTCTGAGCCC TCTGGGCGCTCTGGTGGCCTGCTATAAGGGCGTGTCCTGT AGCATCGGCAGCAATCGGGTGGGCATCATCAAGCAGCTG

TABLE 2 -continued SEO ID NO: Sequence AACAAGGGATGCTCCTACATCACCAACCAGGACGCCGAC

ACCGTGACCATCGACAACACCGTGTACCAGCTGAGCAAG GTGGAGGCGAGCACCTGATCAAGGGCAGACCCGT GAGCTCCAGCTTCGACCCCATCAAGTTCCCTGAGGACCA GTTCAACGTGGCCCTGGACCAGGTGTTTGAGAACATCGA GAACAGCCAGGCCCTGGTGGACCAGAGCAACAGAATCCT GTCCAGCGCTGAGAAGGGCAACACCGGCTTCATCATTGT GATCATTCTGATCGCCGTGCTGGGCAGCTCCATGATCCTG GTGAGCATCTTCATCATTATCAAGAAGACCAAGAAACCC ACCGGAGCCCCTCCTGAGCTGAGCGGCGTGACCAACAAT GGCTTCATTCCCCACAACTGA

qb AY525843.1 : 3065-4684 Human metapneumo virus isolate NL/1/99, complete genome

Description

ATGTCTTGGAAAGTGATGATCATCATTTCGTTACTCATAA CACCCCAGCACGGGCTAAAGGAGAGTTATTTGGAAGAAT CATGTAGTACTATAACTGAGGGATACCTCAGTGTTTTAAG AACAGGCTGGTACACTAATGTCTTCACATTAGAAGTTGGT GATGTTGAAAATCTTACATGTACTGATGGACCTAGCTTAA TCAAAACAGAACTTGATCTAACAAAAAGTGCTTTAAGGG AACTCAAAACAGTCTCTGCTGATCAGTTGGCGAGAGAGG AGCAAATTGAAAATCCCAGACAATCAAGATTTGTCTTAG GTGCGATAGCTCTCGGAGTTGCTACAGCAGCAGCAGTCA CAGCAGGCATTGCAATAGCCAAAACCATAAGGCTTGAGA GTGAGGTGAATGCAATTAAAGGTGCTCTCAAACAAACTA ATGAAGCAGTATCCACATTAGGGAATGGTGTGCGGGTCC TAGCCACTGCAGTGAGAGAGCTAAAAGAATTTGTGAGCA AAAACCTGACTAGTGCAATCAACAGGAACAAATGTGACA TTGCTGATCTGAAGATGGCTGTCAGCTTCAGTCAATTCAA  ${\tt CAGAAGATTTCTAAATGTTGTGCGGCAGTTTTCAGACAAT}$  $\tt GCAGGGATAACACCAGCAATATCATTGGACCTGATGACT$ GATGCTGAGTTGGCCAGAGCTGTATCATACATGCCAACA TCTGCAGGGCAGATAAAACTGATGTTGGAGAACCGCGCA  $\tt ATGGTAAGGAGAAAAGGATTTGGAATCCTGATAGGGGTC$ TACGGAAGCTCTGTGATTTACATGGTTCAATTGCCGATCT  ${\tt TTGGTGTCATAGATACACCTTGTTGGATCATCAAGGCAGC}$  ${\tt TCCCTCTTGCTCAGAAAAAAACGGGAATTATGCTTGCCTC}$ CTAAGAGAGGATCAAGGGTGGTATTGTAAAAATGCAGGA TCTACTGTTTACTACCCAAATGAAAAAGACTGCGAAACA AGAGGTGATCATGTTTTTTGTGACACAGCAGCAGGGATC AATGTTGCTGAGCAATCAAGAGAATGCAACATCAACATA TCTACTACCAACTACCCATGCAAAGTCAGCACAGGAAGA CACCCTATAAGCATGGTTGCACTATCACCTCTCGGTGCTT TGGTGGCTTGCTATAAAGGGGTAAGCTGCTCGATTGGCA GCAATTGGGT

TGGAATCATCAAACAATTACCCAAAGGCTGCTCATACAT AACCAACCAGGATGCAGACACTGTAACAATTGACAATAC  $\tt CGTGTATCAACTAAGCAAAGTTGAAGGTGAACAGCATGT$ AATAAAAGGGAGACCAGTTTCAAGCAGTTTTGATCCAAT CAAGTTTCCTGAGGATCAGTTCAATGTTGCGCTTGATCAA GTCTTCGAAAGCATTGAGAACAGTCAGGCACTAGTGGAC CAGTCAAACAAATTCTAAACAGTGCAGAAAAAGGAAA CACTGGTTTCATTATCGTAGTAATTTTTGGTTGCTGTTCTTG GTCTAACCATGATTTCAGTGAGCATCATCATCATAATCAA GAAAACAAGGAAGCCCACAGGAGCACCTCCAGAGCTGA ATGGTGTCACCAACGGCGGTTTCATACCACATAGTTA

qb|KJ627414.1|: 3015-4634 Human metapneumo virus strain hMPV/Homo sapiens/PER/ CFI0497/2010/B complete genome

ATGTCTTGGAAAGTGATGATTATCATTTCGTTACTCATAA CACCTCAGCATGGACTAAAAGAAAGTTATTTAGAAGAAT CATGTAGTACTATAACTGAAGGATATCTCAGTGTTTTTAAG AACAGGTTGGTACACCAATGTCTTTACATTAGAAGTTGGT GATGTTGAAAATCTTACATGTACTGATGGACCTAGCTTAA TCAAAACAGAACTTGACCTAACCAAAAGTGCTTTAAGAG AACTCAAAACAGTTTCTGCTGATCAGTTAGCGAGAGAAG AACAAATTGAAAATCCCAGACAATCAAGGTTTGTCCTAG GTGCAATAGCTCTTGGAGTTGCCACAGCAGCAGCAGTCA CAGCAGGCATTGCAATAGCCAAAACTATAAGGCTTGAGA GTGAAGTGAATGCAATCAAAGGTGCTCTCAAAACAACCA ATGAGGCAGTATCAACACTAGGAAATGGAGTGCGGGTCC TAGCCACTGCAGTAAGAGAGCTGAAAGAATTTGTGAGCA AAAACCTGACTAGTGCGATCAACAAGAACAAGTGTGACA TTGCTGATTTGAAGATGGCTGTCAGCTTCAGTCAGTTCAA CAGAAGATTCCTAAATGTTGTGCGGCAGTTTTCAGACAAT GCAGGGATAACACCAGCAATATCATTGGACCTGATGAAT GATGCTGAGCTGGCCAGAGCTGTATCATACATGCCAACA  ${\tt TCTGCAGGACAGATAAAACTAATGTTAGAGAACCGTGCA}$ ATGGTGAGGAGAAAAGGATTTGGAATCTTGATAGGGGTC TACGGAAGCTCTGTGATTTACATGGTCCAGCTGCCGATCT

TABLE 2 -continued

|             |          | SEQ |
|-------------|----------|-----|
|             |          | ID  |
| Description | Sequence | NO: |

TTGGTGTCATAAATACACCTTGTTGGATAATCAAGGCAGC TCCCTCTTGTTCAGAAAAAGATGGAAATTATGCTTGCCTC CTAAGAGAGGATCAAGGGTGGTATTGTAAAAATGCAGGA  ${\tt TCCACTGTTTACTACCCAAATGAAAAAGACTGCGAAACA}$ AGAGGTGATCATGTTTTTTGTGACACAGCAGCAGGGATC AATGTTGCTGAGCAATCAAGAGAATGCAACATCAACATA TCTACCACCAACTACCCATGCAAAGTCAGCACAGGAAGA  ${\tt CACCCTATCAGCATGGTTGCACTATCACCTCTCGGTGCTT}$ TGGTAGCTTGCTACAAAGGGGTTAGCTGCTCGACTGGCA GTAATCAGGTTGGAATAATCAAACAACTACCTAAAGGCT GCTCATACATAACTAACCAGGACGCAGACACTGTAACAA TTGACAACACTGTGTATCAACTAAGCAAAGTTGAGGGTG AACAGCATGTAATAAAAGGGAGACCAGTTTCAAGCAGTT TTGATCCAATCAGGTTTCCTGAGGATCAGTTCAATGTTGC GCTTGATCAAGTCTTTGAAAGCATTGAAAACAGTCAAGC ACTAGTGGACCAGTCAAACAAAATTCTGAACAGTGCAGA AAAAGGAAACACTGGT

TTCATTATTGTAATAATTTTGATTGCTGTTTCTTGGGTTAAC CATGATTTCAGTGAGCATCATCATCATAATCAAAAAAAC AAGGAAGCCCACAGGGGCACCTCCGGAGCTGAATGGTGT TACCAACGGCGGTTTCATACCGCATAGTTAG

gb|KJ723483.1|: 5586-7310 Human respiratory syncytial virus strain RSV A/Homo sapiens/USA/84I-215A-01/1984, complete genome ATGGAGTTGCCAATCCTCAAAACAAATGCAATTACCACA ATCCTTGCTGCAGTCACACTCTGTTTCGCTTCCAGTCAAA TTAGCAAAGGCTATCTTAGTGCTCTAAGAACTGGTTGGTA TACTAGTGTTATAACTATAGAATTAAGTAATATCAAGGA AAATAAGTGTAATGGAACAGATGCTAAGGTAAAATTGAT AAAACAAGAATTAGATAAATATAAAAATGCTGTAACAGA ATTGCAGTTGCTCATGCAAAGCACACCAGCAGCCAACAA TCGAGCCAGAAGAGAACTACCAAGGTTTATGAATTATAC ACTCAATAATACCAAAAATACCAATGTAACATTAAGCAA GAAAAGGAAAAGAAGATTTCTTGGCTTTTTGTTAGGTGTT GGATCTGCAATCGCCAGTGGCATTGCTGTATCTAAGGTCC  $\tt TGCACCTAGAAGGGGAAGTGAACAAAATCAAAAGTGCTC$ TACTATCCACAAACAAGGCTGTAGTCAGCTTATCAAATG GAGTTAGTGTCTTAACCAGCAAAGTGTTAGACCTCAAAA ACTATATAGATAAACAGTTGTTACCTATTGTGAACAAGC AAAGCTGCAGCATATCAAACATTGAAACTGTGATAGAGT TCCAACAAAAGAACAACAGACTACTAGAGATTACCAGGG AATTTAGTGTTAATGCAGGTGTAACTACACCTGTAAGCAC TTATATGTTAACTAATAGTGAATTATTATCATTAATCAAT GATATGCCTATAACAAATGATCAGAAAAAGTTAATGTCC AACAATGTTCAAATAGTTAGACAGCAAAGTTACTCTATC ATGTCCATAATAAAGGAGGAAGTCTTAGCATATGTAGTA CAATTACCACTATATGGTGTAATAGATACACCCTGTTGGA AACTGCACACATCCCCTCTATGTACAACCAACACAAAGG AAGGGTCCAACATCTGCTTAACAAGAACCGACAGAGGAT GGTATTGTGACAATGCAGGATCAGTATCTTTCTTCCCACA AGCTGAAACATGTAAAGTTCAATCGAATCGGGTATTTTGT GACACAATGAACAGTTAACATTACCAAGTGAAGTAAAT CTCTGCAACATTGACATATTCAACCCCAAATATGATTGCA AAATTATGACTTCAAAAACAGATGTAAGCAGCTCCGTTA TCACATCTCTAGGAGCCATTGTGTCATGCTATGGCAAAAC TAAATGTACAGCATCCAATAAAAATCGTGGGATCATAAA GACATTTTCTAACGGGTGTGATTATGTATCAAATAAGGG GGTGGATACTGTGTCTGTAGGTAATACATTATATTATGTA AATAAGCAAGAAGGCAAAAGTCTCTATGTAAAAGGTGAA CCAATAATAAATTTCTATGACCCATTAGTGTTCCCCTCTG ATGAATTTGATGCATCAATATCTCAAGTCAATGAGAAGA TTAACCAGAGCCTAGCATTTATTCGTAAATCCGATGAATT ATTACATAATGTAAATGCTGGTAAATCCACCACAAATAT CATGATAACTACTATAATTATAGTGATTATAGTAATATTG TTATCATTAATTGCAGTTGGACTGCTCCTATACTGCAAGG CCAGAAGCACACCAGTCACACTAAGTAAGGATCAACTGA

#### hMPV mRNA Sequences

GTGGTATAAATAATATTGCATTTAGTAACTGA

gi|122891979|gb| EF051124.11 Human metapneumo virus isolate TN/92-4 fusion protein gene, complete genome AUGAGCUGGAAGGUGGUGAUUAUCUUCAGCCUGCUGAU
UACACCUCAACACGGCCUGAAGAGAGAGCUACCUGGAAG
AGAGCUGCUCCACCAUCACCGAGGGCUACCUGAGCGUG
CUGCGGACCGGCUGGUACACCAACGUGUUCACCCUGGA
GGUGGGCGACGUGGAGACCUGACCGACGACGCC
CUAGCCUGAUCAAGACCGAGCUGGACCUGACCAGAGG
GCUCUGAGAGAGCUGAACGACCGACCAGCU

57

TABLE 2 -continued

 $\begin{array}{ccc} & & & & & \\ & & & & \\ & & & & \\ \text{Description} & & \text{Sequence} & & \\ \text{NO:} & & & \\ \end{array}$ 

GGCCAGAGAGCAGAUCGAGAACCCUCGGCAGAGCA GAUUCGUGCUGGGCGCCAUCGCUCUGGGAGUCGCCGCU GCCGCUGCAGUGACAGCUGGAGUGGCCAUUGCUAAGAC CAUCAGACUGGAAAGCGAGGUGACAGCCAUCAACAAUG CCCUGAAGAAGACCAACGAGGCCGUGAGCACCCUGGGC AAUGGAGUGAGAGUGCUGGCCACAGCCGUGCGGGAGCU GAAGGACUUCGUGAGCAAGAACCUGACCAGAGCCAUCA ACAAGAACAAGUGCGACAUCGAUGACCUGAAGAUGGCC GUGAGCUUCUCCCAGUUCAACAGACGGUUCCUGAACGU GGUGAGACAGUUCUCCGACAACGCUGGAAUCACACCUG CCAUUAGCCUGGACCUGAUGACCGACGCCGAGCUGGCU AGAGCCGUGCCCAACAUGCCCACCAGCGCUGGCCAGAU CAAGCUGAUGCUGGAGAACAGAGCCAUGGUGCGGAGAA AGGGCUUCGGCAUCCUGAUUGGGGUGUAUGGAAGCUCC GUGAUCUACAUGGUGCAGCUGCCCAUCUUCGGCGUGAU CGACACCCUGCUGGAUCGUGAAGGCCGCUCCUAGCU GCUCCGAGAAGAAAGGAAACUAUGCCUGUCUGCUGAGA GAGGACCAGGGCUGGUACUGCCAGAACGCCGGAAGCAC AGUGUACUAUCCCAACGAGAAGGACUGCGAGACCAGAG GCGACCACGUGUUCUGCGACACCGCUGCCGGAAUCAAC GUGGCCGAGCAGAGCAAGGAGUGCAACAUCAACAUCAG CACAACCAACUACCCCUGCAAGGUGAGCACCGGACGGC ACCCCAUCAGCAUGGUGGCUCUGAGCCCUCUGGGCGCU CUGGUGGCCUGCUAUAAGGGCGUGUCCUGUAGCAUCGG CAGCAAUCGGGUGGGCAUCAUCAAGCAGCUGAACAAGG GAUGCUCCUACAUCACCAACCAGGACGCCGACACCGUG ACCAUCGACAACACCGUGUACCAGCUGAGCAAGGUGGA GGGCGAGCACGUGAUCAAGGGCAGACCCGUGAGCU CCAGCUUCGACCCCAUCAAGUUCCCUGAGGACCAGUUC AACGUGGCCCUGGACCAGGUGUUUGAGAACAUCGAGAA CAGCCAGGCCCUGGUGGACCAGAGCAACAGAAUCCUGU CCAGCGCUGAGAAGGGCAACACCGGCUUCAUCAUUGUG AUCAUUCUGAUCGCCGUGCUGGGCAGCUCCAUGAUCCU GGUGAGCAUCUUCAUCAUUAUCAAGAAGACCAAGAAAC  $\tt CCACCGGAGCCCCUCCUGAGCUGAGCGGCGUGACCAAC$ AAUGGCUUCAUUCCCCACAACUGA

gb|AY525843.1|: 3065-4684 Human metapneumo virus isolate NL/1/99, complete genome AUGUCUUGGAAAGUGAUGAUCAUCAUUUCGUUACUCAU AACACCCCAGCACGGGCUAAAGGAGAGUUAUUUGGAAG AAUCAUGUAGUACUAUAACUGAGGGAUACCUCAGUGUU UUAAGAACAGGCUGGUACACUAAUGUCUUCACAUUAGA AGUUGGUGAUGUUGAAAAUCUUACAUGUACUGAUGGA CCUAGCUUAAUCAAAACAGAACUUGAUCUAACAAAAAG UGCUUUAAGGGAACUCAAAACAGUCUCUGCUGAUCAGU UGGCGAGAGAGGAGCAAAUUGAAAAUCCCAGACAAUCA AGAUUUGUCUUAGGUGCGAUAGCUCUCGGAGUUGCUAC AGCAGCAGUCACAGCAGGCAUUGCAAUAGCCAAAA CCAUAAGGCUUGAGAGUGAGGUGAAUGCAAUUAAAGG UGCUCUCAAACAAACUAAUGAAGCAGUAUCCACAUUAG GGAAUGGUGUGCGGGUCCUAGCCACUGCAGUGAGAGAG CUAAAAGAAUUUGUGAGCAAAAACCUGACUAGUGCAAU CAACAGGAACAAAUGUGACAUUGCUGAUCUGAAGAUGG CUGUCAGCUUCAGUCAAUUCAACAGAAGAUUUCUAAAU GUUGUGCGGCAGUUUUCAGACAAUGCAGGGAUAACACC AGCAAUAUCAUUGGACCUGAUGACUGAUGCUGAGUUGG CCAGAGCUGUAUCAUACAUGCCAACAUCUGCAGGGCAG AUAAAACUGAUGUUGGAGAACCGCGCAAUGGUAAGGAG AAAAGGAUUUGGAAUCCUGAUAGGGGUCUACGGAAGCU CUGUGAUUUACAUGGUUCAAUUGCCGAUCUUUGGUGUC AUAGAUACACCUUGUUGGAUCAUCAAGGCAGCUCCCUC UUGCUCAGAAAAAAACGGGAAUUAUGCUUGCCUCCUAA GAGAGGAUCAAGGGUGGUAUUGUAAAAAUGCAGGAUC UACUGUUUACUACCCAAAUGAAAAAGACUGCGAAACAA GAGGUGAUCAUGUUUUUUGUGACACAGCAGCAGGGAUC AAUGUUGCUGAGCAAUCAAGAGAAUGCAACAUCAACAU AUCUACUACCAACUACCCAUGCAAAGUCAGCACAGGAA GACACCCUAUAAGCAUGGUUGCACUAUCACCUCUCGGU GCUUUGGUGGCUUGCUAUAAAGGGGUAAGCUGCUCGAU UGGCAGCAAUUGGGU

UGGAAUCAUCAAACAAUUACCCAAAGGCUGCUCAUACA
UAACCAACCAGGAUGCAGACACUGUAACAAUUGACAAU
ACCGUGUAUCAACUAAGCAAGUUGAAGGAGCA
UGUAAUAAAAGGGAGACCAGUUUCAAGCAGUUUUGAUC
CAAUCAAGUUUCCUGAGGAUCAGUUCAAUGUUGGCCUU
GAUCAAGUCUUCGAAACAUGAGAACAGUCAGGCACU
AGUGGACCAGUCAAACAAAAUUCUAAACAGUGCAGAAA

|   | TABLE 2 -continued   |                  |
|---|--|------------------|
| Description   | Sequence   | SEÇ<br>ID<br>NO: |
|   | AAGGAAACACUGGUUUCAUUAUCGUAGUAAUUUUGGU UGCUGUUCUUGGUCUAACCAUGAUUUCAGUGAGCAUCA UCAUCAUAAUCAAGAAAACCAAGGAAGCCCACAGGAGCA CCUCCAGAGCUGAAUGGUGUCACCAACGGCGGUUUCAU ACCACAUAGUUAG  |                  |
| gb KJ627414.1 : 3015-4634 Human metapneumo virus strain hMPV/Homo sapiens/PER/ CFI0497/2010/B, complete genome              | AUGUCUUGGAAAGUGAUUAUCAUUUCGUUACUCAU AACACCUCAGCAUGGACUAAAAGAAAGUUAUUUAGAAG AAUCAUGUAGUACUAUAACUGAAGGAUAUCUCAGUGUU UUAAGAACAGGUUGGUUGAACCCAAUGUCUUUACAUUAGA AGUUGGUGAUGUUGAAAACCGAAUGUCUUUACAUUAGA CCUAGCUUAAUCAAAACAGAACUUGACCUAACCAAAAG UGCUUUAAGAGAACUCAAAACAGAACUUGACCUAACCAAAAG UGCUUUAAGAGAACUCAAAACAGUUUCUGCUGAUCAGU UAGCGAGAGAGAGACUCAAAACAGUUUCUGCUGAUCAGU UAGCGAGAGAGACACAAAUUGAAAAUCCCAGACAAUCA AGGUUUGUCCUAGGUGCAAUAGCUCUUGGAGGUUGCCAC AGCAGCAGCAGCACACACUCAAAACAGUUUCUAGACUAUACA AGGUUUGUCCAAGAGAGAGAGACUUGCAAUAGCCAAAA CUAUAAAGCCUUGAGAGGAGUAGCAAUACAAAC UGCUCUCAAAACAACAAUGAGGCAGUACCAACACUAG GAAAUGGAGUGCGGGUCCUAGCCACUGCAGUAAGAGA CCUGAAAACAACAACAAUGAGGCAGUACCAACACACACAC | 59               |
| gb KJ723483.1 : 5586-7310 Human respiratory syncytial virus strain RSVA/Homo sapiens/USA/84I- 215A-01/1984, complete genome | AUGGAGUUGCCAAUCCUCAAAACAAAUGCAAUUACCAC AAUCCUUGCUGCAGUCACACUCUGUUUCGCUUCCAGUC AAAACACCACUGAAGAAUUUUAUCAAUCAACAUGCAGU GCAGUUAGCAAGAUUUUAUCAAUCAACAUGCAGU GCAGUUAGCAAAGAUUUAAUGAAUUAAAACAAGG GUUGGUAGCAAGAGUUAACAAGAAGUAAAAAAAAAA  | 60               |

UACCACUAUAUGGUGUAAUAGAUACACCCUGUUGGAAA CUGCACACAUCCCCUCUAUGUACAACCAACACAAAGGA

TABLE 2 -continued

| Description | Sequence                               | SEÇ<br>ID<br>NO: |
|-------------|--|------------------|
|             | AGGGUCCAACAUCUGCUUAACAAGAACCGACAGAGGAU |                  |
|             | GGUAUUGUGACAAUGCAGGAUCAGUAUCUUUCUUCCCA |                  |
|             | CAAGCUGAAACAUGUAAAGUUCAAUCGAAUCGGGUAUU |                  |
|             | UUGUGACACAAUGAACAGUUUAACAUUACCAAGUGAAG |                  |
|             | UAAAUCUCUGCAACAUUGACAUAUUCAACCCCAAAUAU |                  |
|             | GAUUGCAAAAUUAUGACUUCAAAAACAGAUGUAAGCAG |                  |
|             | CUCCGUUAUCACAUCUCUAGGAGCCAUUGUGUCAUGCU |                  |
|             | AUGGCAAAACUAAAUGUACAGCAUCCAAUAAAAAUCGU |                  |
|             | GGGAUCAUAAAGACAUUUUCUAACGGGUGUGAUUAUG  |                  |
|             | UAUCAAAUAAGGGGGUGGAUACUGUGUCUGUAGGUAA  |                  |
|             | UACAUUAUAUUAUGUAAAUAAGCAAGAAGGCAAAAGU  |                  |
|             | CUCUAUGUAAAAGGUGAACCAAUAAUAAAUUUCUAUGA |                  |
|             | CCCAUUAGUGUUCCCCUCUGAUGAAUUUGAUGCAUCAA |                  |
|             | UAUCUCAAGUCAAUGAGAAGAUUAACCAGAGCCUAGCA |                  |
|             | UUUAUUCGUAAAUCCGAUGAAUUAUUACAUAAUGUAA  |                  |
|             | AUGCUGGUAAAUCCACCACAAAUAUCAUGAUAACUACU |                  |
|             | AUAAUUAUAGUGAUUAUAGUAAUAUUGUUAUCAUUAA  |                  |
|             | UUGCAGUUGGACUGCUCCUAUACUGCAAGGCCAGAAGC |                  |
|             | ACACCAGUCACACUAAGUAAGGAUCAACUGAGUGGUAU |                  |
|             | AAAUAAUAUUGCAUUUAGUAACUGA              |                  |

## TABLE 3

| TABLE 3   |   |                  |  |
|---|---|------------------|--|
| hMPV Amino Acid Sequences   |   |                  |  |
| Description   | Sequence  | SEQ<br>ID<br>NO: |  |
| gi 122891979 gb <br>EF051124.1  Human<br>metapneumo virus<br>isolate TN/92-4<br>fusion protein gene,<br>complete cds          | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGW YTNVFTLEVGDVENLTCSDGPSLIKTELDLTKSALRELKTVS ADQLAREEQIENPRQSRFVLGAIALGVAAAAAVTAGVAIAK TIRLESEVTAINNALKKTNEAVSTLGNGVRVLATAVRELKD FVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVVRQFS DNAGITPAISLDLMTDAELARAVPNMFTSAGQIKLMLENRA MVRRKGFGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPS CSEKKGNYACLLREDQGWYCQNAGSTVYYPNEKDCETRG DHVFCDTAAGINVAEQSKECNINISTTNYPCKVSTGRHPISM VALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCSYITNQD ADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQF NVALDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAV LGSSMILVSIFIIIKKTKKPTGAPPELSGVTNNGFIPHN | 5                |  |
| gb AY525843.1 :<br>3065-4684 Human<br>metapneumo virus<br>isolate NL/1/99,<br>complete cds                                    | MSWKVMIIISLLITPQHGLKESYLEESCSTITEGYLSVLRTGW YTNVFTLEVGDVENLTCTDGPSLIKTELDLTKSALRELKTVS ADQLAREEQIENPRQSRFVLGAIALGVATAAAVTAGIAIAKT IRLESEVNAIKGALKQTNEAVSTLGNGVRVLATAVRELKEF VSKNLTSAINRNKCDIADLKMAVSFSQFNRRFLNVVRQFSD NAGITPAISLDLMTDAELARAVSYMPTSAGQIKLMLENRAM VRRKGFGILIGVYGSSVIYMVQLPIFGVIDTPCWIIKAAPSCS EKNGNYACLLREDQGWYCKNAGSTVYYPNEKDCETRGDH VFCDTAAGINVAEQSRECNINISTTNYPCKVSTGRHPISMVA LSPLGALVACYKGVSCSIGSNWVGIIKQLPKGCSYITNQDAD TVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFNV ALDQVFESIENSQALVDQSNKILNSAEKGNTGFIIVVILVAVL GLTMISVSIIIIIKKTRKPTGAPPELNGVTNGGFIPHS | 6                |  |
| gb KJ627414.1 :<br>3015-4634 Human<br>metapneumo virus<br>strain hMPV/Homo<br>sapiens/PER/CFI04<br>97/2010/B,<br>complete cds | MSWKVMIIISLLITPQHGLKESYLEESCSTITEGYLSVLRTGW YTNVFTLEVGDVENLTCTDGFSLIKTELDLTKSALRELKTVS ADQLAREEQIENPRQSRFVLGAIALGVATAAAVTAGIAIAKT IRLESEVNAIKGALKTTNEAVSTLGNGVRVLATAVRELKEF VSKNLTSAINKNKCDIADLKMAVSFSQFNRRFLNVVRQFSD NAGITPAISLDLMNDAELARAVSYMPTSAGQIKLMLENRAM VRRKGFGILIGVYGSSVIYMVQLPIFGVINTPCWIIKAAPSCS EKDGNYACLLREDQGMYCKNAGSTVYYPNEKDCETRGDH VFCDTAAGINVAEQSRECNINISTTNYPCKVSTGRHPISMVA LSPLGALVACYKGVSCSTGSNQVGIIKQLPKGCSYITNQDAD TVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIRFPEDQFNV ALDQVFESIENSQALVDQSNKILNSAEKGNTGFIIVIILIAVLG                                       | 7                |  |
| gb KJ723483.1 :<br>5586-7310 Human  | MELPILKTNAITTILAAVTLCFASSQNITEEFYQSTCSAVSKG<br>YLSALRTGWYTSVITIELSNIKENKCNGTDAKVKLIKQELDK   | 8                |  |

TABLE 3 -continued

| hMPV Amino Acid Sequences   |   |                  |
|---|---|------------------|
| Description   | Sequence  | SEQ<br>ID<br>NO: |
| respiratory<br>syncytial virus<br>strain RSVA/Homo<br>sapiens/USA/84I-<br>215A-01/1984,<br>complete cds | YKNAVTELQLLMQSTPAANNRARRELPRFMNYTLNNTKNT  NVTLSKKRKRRPLGFLLGVGSATASGIAVSKVLHLEGEVNKI  KSALLSTNKAVVSLSNGVSVLTSKVLDLKNYIDKQLLPIVN  KQSCSISNIETVIEFQQKNNRLLEITREFSVNAGVTTPVSTYM  LTNSELLSLINDMPITNDQKKLMSNNVQIVRQQSYSIMSIIKE  EVLAYVVQLPLYGVIDTPCWKLHTSPLCTTNTKEGSNICLTR  TDRGWYCDNAGSVSFFPQAETCKVQSNRVFCDTMNSLTLP  SEVNLCNIDIFNPKYDCKIMTSKTDVSSSVITSLGAIVSCYGK  TKCTASNKNRGIIKTFSNGCDYVSNKGVDTVSVGNTLYYVN  KQEGKSLYVKGEPIINFYDPLVFPSDEFDASISQVNEKINQSL  AFIRKSDELLHNVNAGKSTTNIMITTIIIVIIVILLSLIAVGLLL  YCKARSTPVTLSKDQLSGINNIAFSN |                  |

TABLE 4

| hMPV NCBI Accession Numbers (Amino Acid  | l Sequences)             |
|--|--------------------------|
| Virus  | GenBank Accession        |
| F [Human metapneumovirus] [Human metapneumovirus]                                  | AEK26895.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53565.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53566.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53569.1               |
| fusion protein [Human metapneumovirus]   | AEZ52347.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53574.1               |
| fusion glycoprotein [Human metapneumovirus]  | AHV79473.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53570.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53567.1               |
| fusion protein [Human metapneumovirus]   | AAS22125.1               |
| fusion glycoprotein [Human metapneumovirus]  | AHV79795.1               |
| fusion glycoprotein [Human metapneumovirus]  | AHV79455.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53568.1               |
| fusion protein [Human metapneumovirus]   | AAS22109.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGU68417.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGJ74228.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53575.1               |
| fusion protein [Human metapneumovirus]   | AAU25820.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGU68377.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGU68371.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGJ74087.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53560.1               |
| fusion glycoprotein [Human metapneumovirus]  | AHV79858.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53577.1               |
| fusion protein [Human metapneumovirus]   | AAS22085.1               |
| fusion protein [Human metapneumovirus]   | AEZ52348.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGJ74044.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53563.1               |
| fusion glycoprotein precursor [Human metapneumovirus]                              | YP_012608.1              |
| fusion glycoprotein [Human metapneumovirus]  | AGJ74053.1               |
| fusion protein [Human metapneumovirus]   | BAM37562.1               |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53561.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGU68387.1               |
| fusion [Human metapneumovirus]   | AGL74060.1               |
| fusion glycoprotein precursor [Human metapneumovirus]                              | AAV88364.1               |
| fusion protein [Human metapneumovirus]   | AAN52910.1               |
| fusion protein [Human metapneumovirus]   | AAN52915.1               |
| fusion protein [Human metapneumovirus]   | BAM37564.1               |
| fusion glycoprotein precursor [Human metapneumovirus]                              | BAH59618.1               |
| fusion protein [Human metapneumovirus]   | AAQ90144.1               |
| fusion glycoprotein [Human metapneumovirus]  | AHV79446.1               |
| fusion protein [Human metapneumovirus]   | AEL87260.1               |
| fusion glycoprotein [Human metapneumovirus] fusion protein [Human metapneumovirus] | AHV79867.1<br>ABQ66027.2 |
|  | •                        |
| fusion glycoprotein [Human metapneumovirus] fusion protein [Human metapneumovirus] | ACJ53621.1<br>AAN52911.1 |
| fusion glycoprotein [Human metapneumovirus]  | AHV79536.1               |
| fusion glycoprotein [Human metapneumovirus]  | AGU68411.1               |
| fusion protein [Human metapneumovirus]   | AEZ52346.1               |
| fusion protein [Human metapneumovirus]   | AAN52913.1               |
| fusion protein [Human metapneumovirus]   | AAN52913.1<br>AAN52908.1 |
| fusion glycoprotein [Human metapneumovirus]  | ACJ53553.1               |
| ration 5.300protein [trainan metaphetimovirus]                                     | 1100000001               |

TABLE 4-continued

| hMPV NCBI Accession Numbers (Amino Acid   | l Sequences)             |
|---|--------------------------|
| Virus   | GenBank Accession        |
| fusion glycoprotein [Human metapneumovirus]   | AIY25727.1               |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | ABM67072.1<br>AEZ52361.1 |
| fusion protein [Human metapneumovirus]  | AAS22093.1               |
| fusion glycoprotein [Human metapneumovirus]   | AGH27049.1               |
| fusion protein [Human metapneumovirus] fusion glycoprotein [Human metapneumovirus]                  | AAK62968.2<br>ACJ53556.1 |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53620.1               |
| fusion protein [Human metapneumovirus] F [Human metapneumovirus] [Human metapneumovirus]            | ABQ58820.1<br>AEK26886.1 |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53619.1               |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53555.1               |
| fusion [Human metapneumovirus] fusion protein [Human metapneumovirus]                               | AGL74057.1<br>ABD27850.1 |
| fusion protein [Human metapneumovirus]  | AEZ52349.1               |
| fusion protein [Human metapneumovirus]  | ABD27848.1               |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | ABD27846.1<br>ABQ66021.1 |
| fusion protein [Human metapneumovirus]  | AFM57710.1               |
| fusion protein [Human metapneumovirus]  | AFM57709.1               |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | ABH05968.1<br>AEZ52350.1 |
| fusion protein [Human metapneumovirus]  | AFM57712.1               |
| fusion protein [Human metapneumovirus]  | AEZ52364.1               |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | AAN52912.1<br>AEZ52363.1 |
| fusion [Human metapneumovirus]  | AGL74059.1               |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53583.1<br>AEZ52356.1 |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | AEZ52353.1<br>AEZ52353.1 |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53581.1               |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53578.1<br>AAS22117.1 |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | BAN75965.1               |
| fusion protein [Human metapneumovirus]  | AGF92105.1               |
| fusion protein [Human metapneumovirus] fusion protein [Human metapneumovirus]                       | AAS22077.1<br>AAN52909.1 |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53586.1               |
| fusion protein [Human metapneumovirus]  | AAQ90145.1               |
| fusion glycoprotein [Human metapneumovirus] fusion [Human metapneumovirus]                          | AGT75042.1<br>AGL74058.1 |
| fusion protein [Human metapneumovirus]  | AEL87263.1               |
| fusion glycoprotein [Human metapneumovirus]   | AGH27057.1               |
| fusion glycoprotein [Human metapneumovirus] F [Human metapneumovirus] [Human metapneumovirus]       | AHV79491.1<br>AEK26906.1 |
| fusion glycoprotein [Human metapneumovirus]   | ACJ53580.1               |
| fusion protein [Human metapneumovirus]  | AEZ52354.1               |
| fusion protein [Human metapneumovirus] G [Human metapneumovirus] [Human metapneumovirus]            | AAN52914.1<br>AEK26901.1 |
| glycoprotein [Human metapneumovirus]  | AFI56738.1               |
| glycoprotein [Human metapneumovirus]  | AFI56739.1<br>AFI56745.1 |
| glycoprotein [Human metapneumovirus] G protein [Human metapneumovirus]                              | AAQ62718.1               |
| G protein [Human metapneumovirus]   | AAQ62719.1               |
| attachment glycoprotein G [Human metapneumovirus] G protein [Human metapneumovirus]                 | AGH27104.1<br>AAQ62729.1 |
| G protein [Human metapneumovirus]   | AAQ62728.1               |
| glycoprotein [Human metapneumovirus]  | AFI56753.1               |
| glycoprotein [Human metapneumovirus] glycoprotein [Human metapneumovirus]                           | AFI56746.1<br>AFI56750.1 |
| glycoprotein [Human metapneumovirus]  | AFI56747.1               |
| G protein [Human metapneumovirus] glycoprotein [Human metapneumovirus]                              | AAQ62721.1<br>AAT46573.1 |
| glycoprotein [Human metapneumovirus]  | AFI56748.1               |
| glycoprotein [Human metapneumovirus]  | AFI56736.1               |
| glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus]              | AFI56749.1<br>AGH27131.1 |
| attachment glycoprotein G [Human metapneumovirus]   | AHV79558.1               |
| glycoprotein [Human metapneumovirus]  | AFI56740.1               |
| glycoprotein [Human metapneumovirus]<br>glycoprotein [Human metapneumovirus]                        | AFI56741.1<br>AFI56744.1 |
| attachment glycoprotein G [Human metapneumovirus]   | AHV79790.1               |
| attachment glycoprotein G [Human metapneumovirus]   | AGH27122.1               |
| attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] | AHV79763.1<br>AGZ48849.1 |
| glycoprotein [Human metapneumovirus]  | AFI56743.1               |
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| attachment surface glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGI74162.1 attachment glycoprotein G [Human metapneumovirus] AGI74162.1 attachment glycoprotein G [Human metapneumovirus] AGI74162.1 attachment glycoprotein G [Human metapneumovirus] AGI74095.1 attachment glycoprotein [Human metapneumovirus] AGI74095.1 attachment glycoprotein [Human metapneumovirus] AGI74095.1 attachment glycoprotein [Human metapneumovirus] AGI74095.1 AGI74091.1 AGI74091.1   |   |            |
| attachment surface glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] adGW43070.1 glycoprotein [Human metapneumovirus] adGW43070.1 glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] adGW6409.1 attachment glycoprotein G [Human metapneumovirus] adGJ74223.1 attachment glycoprotein [Human metapneumovirus] adAS22129.1 attachment glycoprotein G [Human metapneumovirus] adGJ74048.1 G protein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] adGJ74162.1 attachment glycoprotein G [Human metapneumovirus] adAQ62726.1 attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] adAQ62726.1 attachment glycoprotein [Human metapneumovir |   |            |
| attachment glycoprotein G [Human metapneumovirus] AHV79447.1 attachment surface glycoprotein [Human metapneumovirus] AHV9447.1 attachment surface glycoprotein [Human metapneumovirus] ABQ44523.1 attachment protein [Human metapneumovirus] ABQ44523.1 attachment surface glycoprotein [Human metapneumovirus] BAH59622.1 attachment surface glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGU8409.1 attachment glycoprotein G [Human metapneumovirus] AGU8409.1 attachment glycoprotein [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AAG74048.1 G protein [Human metapneumovirus] ABC46384.1 attachment protein [Human metapneumovirus] ABC44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH74051.1 attachment glycoprotein G [Human metapneumovirus] AGH779531.1 G protein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1 attachment metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1   |   |            |
| attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGJ74091.1 attachment glycoprotein G [Human metapneumovirus] Attachment surface glycoprotein [Human metapneumovirus] AGW43056.1 attachment protein [Human metapneumovirus] AGW43056.1 attachment protein [Human metapneumovirus] ABQ44523.1 attachment surface glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AAJ746585.1 attachment glycoprotein G [Human metapneumovirus] AGU68409.1 attachment glycoprotein [Human metapneumovirus] AGJ74223.1 attachment glycoprotein G [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC44525.1 glycoprotein [Human metapneumovirus] ABC44525.1 attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 G protein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AGW43058.1   |   |            |
| attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment protein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] BAH59622.1 attachment surface glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGU68409.1 attachment glycoprotein G [Human metapneumovirus] AGU74223.1 attachment glycoprotein G [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AAGJ74048.1 G protein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC4525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AFW79531.1 G protein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1  |   |            |
| attachment surface glycoprotein [Human metapneumovirus] AGW43056.1 attachment protein [Human metapneumovirus] BAH59622.1 attachment surface glycoprotein [Human metapneumovirus] BAH59622.1 attachment surface glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGW43070.1 glycoprotein [Human metapneumovirus] AGU68409.1 attachment glycoprotein G [Human metapneumovirus] AGJ74223.1 attachment glycoprotein [Human metapneumovirus] AGJ74223.1 attachment glycoprotein G [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] AAQ62725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC26384.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH74162.1 attachment glycoprotein G [Human metapneumovirus] AGH74055.1 attachment glycoprotein G [Human metapneumovirus] AGH7405.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] ploophoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79631.1   |   |            |
| attachment protein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AAT46585.1 attachment glycoprotein G [Human metapneumovirus] AGU68409.1 attachment glycoprotein G [Human metapneumovirus] AGJ74223.1 attachment glycoprotein G [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AAGJ74048.1 G protein [Human metapneumovirus] AAGG2725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAPC2726.1 attachment glycoprotein [Human metapneumovirus] AAPC2726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1  |   |            |
| attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGJ74223.1 attachment glycoprotein G [Human metapneumovirus] AGJ74223.1 attachment glycoprotein G [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC26384.1 attachment glycoprotein G [Human metapneumovirus] ABQ44525.1 attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment surface glycoprotein [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] ABC26384.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79631.1   | attachment protein [Human metapneumovirus]              |            |
| glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein [Human metapneumovirus] attachment glycoprotein [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AAGJ74048.1 G protein [Human metapneumovirus] AAQ62725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC455.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1   | attachment glycoprotein G [Human metapneumovirus]       |            |
| attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGJ74223.1 attachment glycoprotein [Human metapneumovirus] AAS22129.1 attachment glycoprotein G [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] AAQ62725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABQ44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGW2726.1 attachment glycoprotein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AAS48465.1  |   |            |
| attachment glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGJ74048.1 G protein [Human metapneumovirus] AAQ62725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABQ44525.1 attachment glycoprotein G [Human metapneumovirus] ATAG44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AHV79531.1 G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AHX79631.1 phosphoprotein [Human metapneumovirus] AHV79631.1  |   |            |
| attachment glycoprotein G [Human metapneumovirus] G protein [Human metapneumovirus] AAQ62725.1 glycoprotein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABQ44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AHV79531.1 G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AFW43058.1 P [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AFW43058.1 P [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1   | attachment glycoprotein G [Human metapneumovirus]       | AGJ74223.1 |
| G protein [Human metapneumovirus]  glycoprotein [Human metapneumovirus]  attachment protein [Human metapneumovirus]  attachment glycoprotein G [Human metapneumovirus]  attachment surface glycoprotein [Human metapneumovirus]  attachment glycoprotein G [Human metapneumovirus]  AGH74162.1  attachment glycoprotein G [Human metapneumovirus]  APL79531.1  G protein [Human metapneumovirus]  AAQ62726.1  attachment glycoprotein [Human metapneumovirus]  AAS48465.1  attachment surface glycoprotein [Human metapneumovirus]  AGW43058.1  P [Human metapneumovirus] [Human metapneumovirus]  AEK26894.1  phosphoprotein [Human metapneumovirus]  AHV79631.1  phosphoprotein [Human metapneumovirus]  AHV79901.1   |   |            |
| glycoprotein [Human metapneumovirus] attachment protein [Human metapneumovirus] ABC26384.1 attachment protein [Human metapneumovirus] ABC44525.1 attachment glycoprotein G [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGI74162.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AAC62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  |   |            |
| attachment glycoprotein G [Human metapneumovirus] attachment surface glycoprotein [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AHV79531.1 G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] P [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  | glycoprotein [Human metapneumovirus]                    | ABC26384.1 |
| attachment surface glycoprotein [Human metapneumovirus] AGW43071.1 attachment glycoprotein G [Human metapneumovirus] AGI741.62.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AHV79531.1 G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AEK26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1   |   |            |
| attachment glycoprotein G [Human metapneumovirus] AGJ74162.1 attachment glycoprotein G [Human metapneumovirus] AGH27095.1 attachment glycoprotein G [Human metapneumovirus] AHV79531.1 G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AEX26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1   |   |            |
| attachment glycoprotein G [Human metapneumovirus] G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] ABK26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  | attachment glycoprotein G [Human metapneumovirus]       |            |
| G protein [Human metapneumovirus] AAQ62726.1 attachment glycoprotein [Human metapneumovirus] AAS48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AEK26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  |   |            |
| attachment glycoprotein [Human metapneumovirus] AA\$48465.1 attachment surface glycoprotein [Human metapneumovirus] AGW43058.1 P [Human metapneumovirus] [Human metapneumovirus] AEK26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  |   |            |
| attachment surface glycoprotein [Human metapneumovirus] P [Human metapneumovirus] [Human metapneumovirus] AGW43058.1 AEK26894.1 phosphoprotein [Human metapneumovirus] AHV79631.1 phosphoprotein [Human metapneumovirus] AHV79901.1  | attachment glycoprotein [Human metapneumovirus]         |            |
| phosphoprotein [Human metapneumovirus] AHV79631.1<br>phosphoprotein [Human metapneumovirus] AHV79901.1   |   |            |
| phosphoprotein [Human metapneumovirus] AHV79901.1  |   |            |
|  |   |            |
|  | phosphoprotein [Human metapneumovirus]                  | AHV79570.1 |

237

TABLE 4-continued

| TABLE 4-continued  |  |  |
|--|--|--|
| hMPV NCBI Accession Numbers (A   |  |  |
| Virus  | GenBank Accession                      |  |
| phosphoprotein [Human metapneumovirus]   | AGJ74076.1<br>AAS22123.1               |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AAS22123.1<br>ABB16895.1               |  |
| phosphoprotein [Human metapneumovirus]   | AHV79579.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AGJ74244.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AHV79856.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ACJ70113.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AGZ48843.1<br>AHV79498.1               |  |
| phosphoprotein [Human metapneumovirus]   | AHV79480.1<br>AHV79480.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43382.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAS22107.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABB16898.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AGH27134.1<br>ABB16899.1               |  |
| phosphoprotein [Human metapneumovirus]   | AGH27098.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAN52866.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAS22083.1                             |  |
| phosphoprotein [Human metapneumovirus]   | YP_012606.1                            |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]    | AHV79973.1<br>AHV79462.1               |  |
| phosphoprotein [Human metapneumovirus]   | AGJ74042.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAV88362.1                             |  |
| P [Human metapneumovirus] [Human metapneumov                                     |  |  |
| phosphoprotein [Human metapneumovirus]   | AHV79453.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AGJ74261.1<br>AGH27116.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABB16444.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABB16445.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AHV79507.1                             |  |
| phosphoprotein [Human metapneumovirus]   | BAH59616.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | ABB16443.1<br>ABQ43388.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43389.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43395.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43385.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAP84042.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AAN52868.1<br>AAP84041.1               |  |
| phosphoprotein [Human metapneumovirus]   | AGH27080.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43387.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAS22099.1                             |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]    | ABB16896.1<br>AGJ74094.1               |  |
| phosphoprotein [Human metapneumovirus]   | AEZ68089.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABK97002.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAP13486.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AHV79444.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AHV79865.1<br>AGJ74226.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43383.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AAN52863.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AHV79775.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AEZ68094.1<br>AHV79883.1               |  |
| phosphoprotein [Human metapneumovirus]   | AEZ68092.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43390.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43386.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43391.1                             |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]    | ACS16062.1<br>AEZ68090.1               |  |
| phosphoprotein [Human metapneumovirus]   | AAK62967.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AEZ68093.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AEZ68088.1                             |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43392.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | ABQ43393.1<br>ABQ43384.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABQ43394.1<br>ABQ43394.1               |  |
| phosphoprotein [Human metapneumovirus]   | ABK96999.1                             |  |
| phosphoprotein [Human metapneumovirus]   | AHV79489.1                             |  |
| phosphoprotein [Human metapneumovirus]<br>phosphoprotein [Human metapneumovirus] | AGJ74235.1<br>AAS22075.1               |  |
| phosphoprotein [Human metapneumovirus]   | AAS22075.1<br>AAS22115.1               |  |
| phosphoprotein [Human metapneumovirus]   |  |  |
| phosphoprotem [Italian metaphetimovirus]   | AII17601.1                             |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]    | AII17601.1<br>ABK97000.1<br>AHV79561.1 |  |

239

TABLE 4-continued

| hMPV NCBI Accession Numbers (Amino Acid Sequences)                                       |                            |  |  |
|--|----------------------------|--|--|
| Virus  | GenBank Accession          |  |  |
|  |                            |  |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]            | AGT75040.1<br>AAN52864.1   |  |  |
| phosphoprotein [Human metapneumovirus]   | ABK97001.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AGT74979.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AHV79955.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AGH27055.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AAV88361.1                 |  |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]            | ABQ43397.1<br>AGJ74173.1   |  |  |
| P [Human metapneumovirus] [Human metapneumovirus]  | AEK26904.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | ACJ70104.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | ABK97003.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AGT74955.1                 |  |  |
| phosphoprotein [Human metapneumovirus] phosphoprotein [Human metapneumovirus]            | AAN52856.1<br>AAN52862.1   |  |  |
| phosphoprotein [Human metapneumovirus]   | AGJ74138.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AHV79613.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AGJ74060.1                 |  |  |
| phosphoprotein [Human metapneumovirus]   | AAQ67684.1                 |  |  |
| phosphoprotein [Human metapneumovirus] N [Human metapneumovirus] [Human metapneumovirus] | AEA02278.1<br>AEK26899.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | ACS16061.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AAS88425.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | YP_012605.1                |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79882.1                 |  |  |
| nucleoprotein [Human metapneumovirus]<br>nucleocapsid protein [Human metapneumovirus]    | AHV79774.1<br>AAN52886.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AAN32880.1<br>AAS22082.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79864.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79828.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGJ74084.1                 |  |  |
| nucleocapsid protein [Human metapneumovirus]   | AAN52888.1                 |  |  |
| N [Human metapneumovirus] [Human metapneumovirus] nucleoprotein [Human metapneumovirus]  | AIL23590.1<br>AAK62966.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79972.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79470.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79452.1<br>AGJ74243.1   |  |  |
| nucleoprotein [Human metapneumovirus]<br>nucleoprotein [Human metapneumovirus]           | AHV79533.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGJ74181.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79497.1                 |  |  |
| nucleoprotein [Human metapneumovirus]<br>nucleoprotein [Human metapneumovirus]           | AHV79702.1<br>AHV79648.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79435.1                 |  |  |
| putative nucleoprotein [Human metapneumovirus]   | AGJ74260.1                 |  |  |
| nucleocapsid protein [Human metapneumovirus]   | AAN52887.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68386.1                 |  |  |
| nucleocapsid protein [Human metapneumovirus] nucleoprotein [Human metapneumovirus]       | AAN52899.1<br>AAR17673.1   |  |  |
| nucleocapsid protein [Human metapneumovirus]   | AAN52898.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AEA02277.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AHV79612.1                 |  |  |
| nucleoprotein [Human metapneumovirus]<br>nucleoprotein [Human metapneumovirus]           | AGU68416.1<br>AGU68408.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68370.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AAQ67683.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGJ74137.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68344.1                 |  |  |
| nucleocapsid protein [Human metapneumovirus] nucleoprotein [Human metapneumovirus]       | ABK96997.1<br>AGU68413.1   |  |  |
| nucleocapsid protein [Human metapneumovirus]   | AAN52891.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68360.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68353.1                 |  |  |
| nucleocapsid protein [Human metapneumovirus] nucleoprotein [Human metapneumovirus]       | ABK96996.1<br>AAR17666.1   |  |  |
| N [Human metapneumovirus] [Human metapneumovirus]  | AEK26903.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGT75039.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGU68410.1                 |  |  |
| nucleoprotein [Human metapneumovirus]<br>nucleoprotein [Human metapneumovirus]           | AAS22074.1<br>AHV79560.1   |  |  |
| nucleoprotein [Human metapneumovirus]  | AGT74978.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AGJ74128.1                 |  |  |
| nucleoprotein [Human metapneumovirus]  | AAR17663.1                 |  |  |
| nucleoprotein [Human metapneumovirus] nucleoprotein [Human metapneumovirus]              | AAR17662.1<br>AAR17664.1   |  |  |
| nucleoprotein [Human metapneumovirus] nucleoprotein [Human metapneumovirus]              | AAR1 /664.1<br>AAR1 7657.1 |  |  |
|  |                            |  |  |

241

TABLE 4-continued

| Virus  | GenBank Accession        |
|--|--------------------------|
| nucleoprotein [Human metapneumovirus]        | AAR17659.1               |
| nucleoprotein [Human metapheumovirus]        | AAR17661.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68352.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68373.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68376.1<br>AGU68376.1 |
| nucleoprotein [Human metapneumovirus]        | AGU68370.1<br>AGU68342.1 |
| nucleoprotein [Human metapneumovirus]        | AGU68342.1<br>AGU68365.1 |
| nucleoprotein [Human metapneumovirus]        | AGU68363.1<br>AGU68363.1 |
| nucleoprotein [Human metapneumovirus]        | AGU68398.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68348.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68354.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68391.1               |
| nucleoprotein [Human metapheumovirus]        | AGU68389.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68399.1<br>AGU68399.1 |
| nucleoprotein [Human metapheumovirus]        | AGU68337.1               |
| nucleoprotein [Human metapneumovirus]        | AAR17660.1               |
| nucleoprotein [Human metapheumovirus]        | AAR17667.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68402.1               |
| nucleoprotein [Avian metapneumovirus]        | CDN30025.1               |
| nucleoprotein [Avian metapheumovirus type C] | AGZ87947.1               |
| Nucleoprotein [Avian metapneumovirus type C] | CAL25113.1               |
| nucleocapsid protein [Avian metapheumovirus] | ABO42286.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK38430.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK54155.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK34133.1<br>AAK38426.1 |
| nucleocapsid protein [Avian metapheumovirus] | AAK38425.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK38424.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK36424.1<br>AAF05909.1 |
| nucleocapsid protein [Avian metapheumovirus] | AAK38435.1               |
| nucleocapsid protein [Avian metapheumovirus] | AAK38428.1               |
| nucleocapsia protein [Avian metapheumovirus] | AAR17669.1               |
|  |                          |
| nucleocapsid protein [Avian metapneumovirus] | AAK38429.1               |
| nucleocapsid protein [Avian metapneumovirus] | AAK38427.1               |
| nucleocapsid protein [Avian metapneumovirus] | AAK38423.1               |
| nucleocapsid protein [Avian metapneumovirus] | AAK38434.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68338.1               |
| nucleoprotein [Avian metapneumovirus]        | YP_443837.1              |
| nucleoprotein [Human metapneumovirus]        | AGU68384.1               |
| nucleocapsid protein [Avian metapneumovirus] | AAK38431.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68405.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68382.1               |
| nucleoprotein [Human metapneumovirus]        | AGU68395.1               |
| nucleocapsid [Human metapneumovirus]         | AAL35389.3               |
| nucleoprotein [Human metapneumovirus]        | AEZ68064.1               |

TABLE 5

| Description   | Sequence  | SEQ<br>ID<br>NO: |
|---|---|------------------|
|   | PIV3 Nucleic Acid Sequences   |                  |
| >gb KJ672601.1 : 4990-6609 Human parainfluenza virus 3 strain HPIV3/Homo sapiens/ PER/FLA4815/2008 [fusion glycoprotein F0] | ATGCCAATTTCAATACTGTTAATTATTACAACCATGATC ATGCCATCACACTGCCAAATAGACATCACAAAACTACA GCATGTAGGTGTATTGGTCAACAGTCCCAAAAGGGATGA AGATATCACAAAACTTCGAAACAAGATATCTAATCCTGA GTCTCATACCAAAAACTTCGAAACAAGATATCTAATCCTGA GTCTCATACCAAAAATAGAAGATTCTAACTCTTGTGGTG ACCAACAGATCAAGCAATACAAGAGGTTATTGGATAGA CTGATCATTCCTTTATATGATGGACTAAGATTACAGAAG GATGTGATAGTGACTAATCAAGAATCCAATGAAAACAC TGATCCCAGAACAGAA | 9                |

244

TABLE 5 -continued

 $\begin{array}{ccc} & & & & & \\ & & & & & \\ \text{Description} & & \text{Sequence} & & \text{NO:} \end{array}$ 

ATTTACAGAATCAATAAAGGTGAGAGTTATAGATGTTGA TTTGAATGATTACTCAATAACCCTCCAAGTCAGACTCCC TTTATTGACCAGACTGCTGAACACTCAAATCTACAAAGT AGATTCCATATCATACAATATCCAAAATAGAGAATGGTA TATCCCTCTTCCCAGCCATATCATGACGAAAGGGGCATT TCTAGGTGGAGCAGATGTCAAAGAATGCATAGAAGCAT TCAGCAGTTATATATGCCCTTCTGATCCAGGATTTGTACT AAACCATGAAATGGAGAGCTGTCTATCAGGAAACATAT CCCAATGTCCAAGAACCACAGTCACATCAGACATAGTTC CTAGGTATGCATTTGTCAATGGAGGAGTGGTTGCGAATT GTATAACAACTACATGTACATGCAATGGTATCGGTAATA GAATCAACCAACCACCTGATCAAGGAGTCAAAATTATA ACACATAAAGAATGTAATACAATAGGTATCAACGGAAT GCTATTCAACACAAACAAAGAAGGAACTCTTGCATTCTA CACACCAGACGACATAACATTAAACAATTCTGTTGCACT TGATCCGATTGACATATCAATCGAGCTCAACAAGGCCAA ATCAGATCTTGAGGAATCAAAAGAATGGATAAGAAGGT CAAATCAAAAGCTAGATTCTATTGGAAGTTGGCATCAAT CTAGCACTACAATCATAGTTATTTTGATAATGATGATTA TATTGTTTATAATTAATATAACAATAATTACAATTGCAA TTAAGTATTACAGAATTCAAAAGAGAAATCGAGTGGAT CAAAATGATAAGCCGTATGTATTAACAAACAAG

ATGGAATACTGGAAGCACCACCAACCACGGAAAGGATGC

gi|612507167|gb| AHX22430.1| hemagglutininneuraminidase [Human parainfluenza virus 3]

TGGTAATGAGCTGGAGACATCCACAGCCACTCATGGCA ACAAGCTCACCAACAAGATAACATATATATTGTGGACG ATAACCCTGGTGTTATTATCAATAGTCTTCATCATAGTG  $\tt CTAACTAATTCCATCAAAAGTGAAAAGGCCCGCGAATC$  ${\tt ATTGCTACAAGACATAAATAATGAGTTTATGGAAGTTAC}$ AGAAAAGATCCAAGTGGCATCGGATAATACTAATGATC TAATACAGTCAGGAGTGAATACAAGGCTTCTTACAATTC AGAGTCATGTCCAGAATTATATACCAATATCATTGACAC AACAAATATCGGATCTTAGGAAATTCATTAGTGAAATTA CAATTAGAAATGATAATCAAGAAGTGCCACCACAAAGA ATAACACATGATGTGGGTATAAAACCTTTAAATCCAGAT GATTTCTGGAGATGCACGTCTGGTCTTCCATCTTTGATG AAAACTCCAAAAATAAGATTAATGCCGGGACCAGGATT ATTAGCTATGCCAACGACTGTTGATGGCTGTCTCAGAAC  $\tt CCCGTCCTTAGTGATAAATGATCTGATTTATGCTTACAC$ CTCAAATCTAATTACTCGAGGTTGCCAGGATATAGGGAA ATCATATCAAGTATTACAGATAGGGATAATAACTGTAAA CTCAGACTTGGTACCTGACTTAAATCCTAGGATCTCTCA TACCTTCAACATAAATGACAATAGAAAGTCATGTTCTCT AGCACTCCTAAATACAGATGTATATCAACTGTGTTCAAC CCCAAAAGTTGATGAAAGATCAGATTATGCATCATCAG GCATAGAAGATATTGTACTTGATATTGTCAATTATGATG GCTCAATCTCGACAACAAGATTTAAGAATAATAATAA GTTTTGATCAACCATATGCGGCATTATACCCATCTGTTG GACCAGGGATATACTACAAAGGCAAAATAATATTTCTC GGGTATGGAGGTCTTGAACATCCAATAAATGAGAATGC AATCTGCAACACAACTGGGTGTCCTGGGAAAACACAGA GAGACTGTAATCAAGCATCTCATAGTCCATGGTTTTCAG ATAGAAGGATGGTCAACTCTATAATTGTTGTTGACAAGG GCTTGAACTCAGTTCCAAAATTGAAGGTATGGACGATAT CTATGAGACAAAATTACTGGGGGTCAGAAGGAAGATTA CTTCTACTAGGTAACAAGATCTACATATACACAAGATCT ACAAGTTGGCACAGCAAGTTACAATTAGGAATAATTGA CATTACTGACTACAGTGATATAAGGATAAAATGGACAT GGCATAATGTGCTATCAAGACCAGGAAACAATGAATGT CCATGGGGACATTCATGTCCGGATGGATGTATAACGGG AGTATATACCGATGCATATCCACTCAATCCCACAGGAAG CATTGTATCATCTGTCATATTGGACTCACAAAAATCGAG AGTCAACCCAGTCATAACTTACTCAACAGCAACCGAAA GGGTAAACGAGCTGGCTATCCGAAACAAAACACTCTCA GCTGGGTACACAACAACAAGCTGCATTACACACTATAA CAAAGGGTATTGTTTTCATATAGTAGAAATAAATCATAA AAGCTTAAACACATTTCAACCCATGTTGTTCAAAACAGA GATTCCAAAAAGCTGCAGT

HPIV3\_HN\_Codon
Optimized

10

246

TABLE 5 -continued

SEQ ID Description Sequence NO:

CTGATCCAGAGCGGCGTGAACACCCGGCTGCTGACCATC CAGAGCCACGTGCAGAACTACATCCCCATCAGCCTGACC CAGCAGATCAGCGACCTGCGGAAGTTCATCAGCGAGAT CACCATCCGGAACGACAACCAGGAAGTGCCCCCCAGA GAATCACCCACGACGTGGGCATCAAGCCCCTGAACCCC GACGATTTCTGGCGGTGTACAAGCGGCCTGCCCAGCCTG ATGAAGACCCCCAAGATCCGGCTGATGCCTGGCCCTGG ACTGCTGGCCATGCCTACCACAGTGGATGGCTGTGTGCG GACCCCCAGCCTCGTGATCAACGATCTGATCTACGCCTA CACCAGCAACCTGATCACCCGGGGCTGCCAGGATATCG GCAAGAGCTACCAGGTGCTGCAGATCGGCATCATCACC GTGAACTCCGACCTGGTGCCCGACCTGAACCCTCGGATC AGCCACACCTTCAACATCAACGACAACAGAAAGAGCTG CAGCCTGGCTCTGCTGAACACCGACGTGTACCAGCTGTG CAGCACCCCAAGGTGGACGAGAGAGAGCGACTACGCCA GCAGCGCATCGAGGATATCGTGCTGGACATCGTGAAC TACGACGCAGCATCAGCACCACCCGGTTCAAGAACAA CAACATCAGCTTCGACCAGCCCTACGCCGCCCTGTACCC TTCTGTGGGCCCTGGCATCTACTACAAGGGCAAGATCAT CTTCCTGGGCTACGGCGGCCTGGAACACCCCATCAACGA GAACGCCATCTGCAACACCACCGGCTGCCCTGGCAAGA CCCAGAGAGACTGCAATCAGGCCAGCCACAGCCCCTGG TTCAGCGACCGCAGAATGGTCAACTCTATCATCGTGGTG GACAAGGGCCTGAACAGCGTGCCCAAGCTGAAAGTGTG GACAATCAGCATGCGCCAGAACTACTGGGGCAGCGAGG GCAGACTTCTGCTGCTGGGAAACAAGATCTACATCTACA CCCGGTCCACCAGCTGGCACAGCAAACTGCAGCTGGGA ATCATCGACATCACCGACTACAGCGACATCCGGATCAA GTGGACCTGGCACAACGTGCTGAGCAGACCCGGCAACA ATGAGTGCCCTTGGGGCCACAGCTGCCCCGATGGATGTA TCACCGGCGTGTACACCGACGCCTACCCCCTGAATCCTA  $\tt CCGGCTCCATCGTGTCCAGCGTGATCCTGGACAGCCAGA$ AAAGCAGAGTGAACCCCGTGATCACATACAGCACCGCC ACCGAGAGAGTGAACGAACTGGCCATCAGAAACAAGAC CCTGAGCGCCGGCTACACCACCACAAGCTGCATCACAC ACTACAACAAGGGCTACTGCTTCCACATCGTGGAAATCA ACCACAAGTCCCTGAACACCTTCCAGCCCATGCTGTTCA AGACCGAGATCCCCAAGAGCTGCTCC

HPIV3\_F\_Codon Optimized

ATGCCCATCAGCATCCTGCTGATCATCACCACAATGATC ATGGCCAGCCACTGCCAGATCGACATCACCAAGCTGCA GCACGTGGGCGTGCTCGTGAACAGCCCCAAGGGCATGA AGATCAGCCAGAACTTCGAGACACGCTACCTGATCCTGA GCCTGATCCCCAAGATCGAGGACAGCAACAGCTGCGGC GACCAGCAGATCAAGCAGTACAAGCGGCTGCTGGACAG ACTGATCATCCCCCTGTACGACGGCCTGCGGCTGCAGAA AGACGTGATCGTGACCAACCAGGAAAGCAACGAGAACA  $\tt CCGACCCCGGACCGAGAGATTCTTCGGCGGCGTGATCG$ GCACAATCGCCCTGGGAGTGGCCACAAGCGCCCAGATT ACAGCCGCTGTGGCCCTGGTGGAAGCCAAGCAGGCCAG AAGCGACATCGAGAAGCTGAAAGAGGCCATCCGGGACA CCAACAAGGCCGTGCAGAGCGTGCAGTCCAGCGTGGGC AATCTGATCGTGGCCATCAAGTCCGTGCAGGACTACGTG AACAAAGAAATCGTGCCCTCTATCGCCCGGCTGGGCTGT GAAGCTGCCGGACTGCAGCTGGGCATTGCCCTGACACA GCACTACAGCGAGCTGACCAACATCTTCGGCGACAACA TCGGCAGCCTGCAGGAAAAGGGCATTAAGCTGCAGGGA ATCGCCAGCCTGTACCGCACCAACATCACCGAGATCTTC ACCACCAGCACCGTGGATAAGTACGACATCTACGACCT GCTGTTCACCGAGAGCATCAAAGTGCGCGTGATCGACGT GGACCTGAACGACTACAGCATCACCCTGCAAGTGCGGC TGCCCCTGCTGACCAGACTGCTGAACACCCAGATCTACA AGGTGGACAGCATCTCCTACAACATCCAGAACCGCGAG TGGTACATCCCTCTGCCCAGCCACATTATGACCAAGGGC GCCTTTCTGGGCGGAGCCGACGTGAAAGAGTGCATCGA GGCCTTCAGCAGCTACATCTGCCCCAGCGACCCTGGCTT CGTGCTGAACCACGAGATGGAAAGCTGCCTGAGCGGCA ACATCAGCCAGTGCCCCAGAACCACCGTGACCTCCGAC ATCGTGCCCAGATACGCCTTCGTGAATGGCGGCGTGGTG GCCAACTGCATCACCACCACCTGTACCTGCAACGGCATC GGCAACCGGATCAACCAGCCTCCCGATCAGGGCGTGAA GATTATCACCCACAAAGAGTGTAACACCATCGGCATCA ACGGCATGCTGTTCAATACCAACAAGAGGGCACCCTG GCCTTCTACACCCCGACGATATCACCCTGAACAACTCC GTGGCTCTGGACCCCATCGACATCTCCATCGAGCTGAAC AAGGCCAAGAGCGACCTGGAAGAGTCCAAAGAGTGGAT

247

TABLE 5 -continued

|             |  | SEQ |
|-------------|--|-----|
| Description | Sequence                                 | NO: |
|             | CCGGCGGAGCAACCAGAAGCTGGACTCTATCGGCAGCT   |     |
|             | GGCACCAGAGCACCACCATCATCGTGATCCTGATTA     |     |
|             | TGATGATTATCCTGTTCATCATCAACATTACCATCATCAC |     |
|             | TATCGCCATTAAGTACTACCGGATCCAGAAACGGAACC   |     |
|             | GGGTGGACCAGAATGACAAGCCCTACGTGCTGACAAAC   |     |
|             | AAG                                      |     |
|             | PIV3 mRNA Sequences                      |     |

>gb|KJ672601.1|: 4990-6609 Human parainfluenza virus 3 strain HPIV3/Homo sapiens/ PER/FLA4815/ 2008 [fusion glycoprotein F0] AUGCCAAUUUCAAUACUGUUAAUUAUUACAACCAUGA UCAUGGCAUCACACUGCCAAAUAGACAUCACAAAACU ACAGCAUGUAGGUGUAUUGGUCAACAGUCCCAAAGGG AUGAAGAUAUCACAAAACUUCGAAACAAGAUAUCUAA UCCUGAGUCUCAUACCAAAAAUAGAAGAUUCUAACUC UUGUGGUGACCAACAGAUCAAGCAAUACAAGAGGUUA UUGGAUAGACUGAUCAUUCCUUUAUAUGAUGGACUAA GAUUACAGAAGGAUGUGAUAGUGACUAAUCAAGAAUC CAAUGAAAACACUGAUCCCAGAACAGAACGAUUCUUU GGAGGGGUAAUUGGAACUAUUGCUCUAGGAGUAGCAA CCUCAGCACAAAUUACAGCAGCAGUUGCUCUGGUUGA AGCCAAGCAGGCAAGAUCAGACAUUGAAAAACUCAAG GAAGCAAUCAGGGACACAAAUAAAGCAGUGCAGUCAG UUCAGAGCUCUGUAGGAAAUUUGAUAGUAGCAAUUAA AUCAGUCCAGGAUUAUGUCAACAAAGAAAUCGUGCCA UCGAUUGCGAGACUAGGUUGUGAAGCAGCAGGACUUC AGUUAGGGAUUGCAUUAACACAGCAUUACUCAGAAUU AACAAAUAUUUGGUGAUAACAUAGGAUCGUUACAA GAAAAAGGAAUAAAAUUACAAGGUAUAGCAUCAUUAU ACCGUACAAAUAUCACAGAAAUAUUCACAACAUCAAC AGUUGACAAAUAUGAUAUUUAUGAUCUAUUAUUUACA GAAUCAAUAAAGGUGAGAGUUAUAGAUGUUGAUUUGA AUGAUUACUCAAUAACCCUCCAAGUCAGACUCCCUUU AUUGACCAGACUGCUGAACACUCAAAUCUACAAAGUA GAUUCCAUAUCAUACAAUAUCCAAAAUAGAGAAUGGU AUAUCCCUCUUCCCAGCCAUAUCAUGACGAAAGGGGC AUUUCUAGGUGGAGCAGAUGUCAAAGAAUGCAUAGAA GCAUUCAGCAGUUAUAUAUGCCCUUCUGAUCCAGGAU UUGUACUAAACCAUGAAAUGGAGAGCUGUCUAUCAGG AAACAUAUCCCAAUGUCCAAGAACCACAGUCACAUCA GACAUAGUUCCUAGGUAUGCAUUUGUCAAUGGAGGAG UGGUUGCGAAUUGUAUAACAACUACAUGUACAUGCAA UGGUAUCGGUAAUAGAAUCAACCAACCACCUGAUCAA GGAGUCAAAAUUAUAACACAUAAAGAAUGUAAUACAA UAGGUAUCAACGGAAUGCUAUUCAACACAAACAAAGA AGGAACUCUUGCAUUCUACACACCAGACGACAUAACA UUAAACAAUUCUGUUGCACUUGAUCCGAUUGACAUAU CAAUCGAGCUCAACAAGGCCAAAUCAGAUCUUGAGGA AUCAAAAGAAUGGAUAAGAAGGUCAAAUCAAAAGCUA GAUUCUAUUGGAAGUUGGCAUCAAUCUAGCACUACAA UCAUAGUUAUUUUGAUAAUGAUGAUUAUAUUGUUUAU AAUUAAUAUAACAAUAAUUACAAUUGCAAUUAAGUAU UACAGAAUUCAAAAGAGAAAUCGAGUGGAUCAAAAUG AUAAGCCGUAUGUAUUAACAAACAAG

gi|612507167|gb|
AHX22430.1|
hemagglutininneuraminidase
[Human
parainfluenza virus
3]

AUGGAAUACUGGAAGCACCACCACCACGGAAAGGAUG CUGGUAAUGAGCUGGAGACAUCCACAGCCACUCAUGG CAACAAGCUCACCAACAAGAUAACAUAUAUAUUGUGG ACGAUAACCCUGGUGUUAUUAUCAAUAGUCUUCAUCA UAGUGCUAACUAAUUCCAUCAAAAGUGAAAAGGCCCG CGAAUCAUUGCUACAAGACAUAAAUAAUGAGUUUAUG GAAGUUACAGAAAAGAUCCAAGUGGCAUCGGAUAAUA CUAAUGAUCUAAUACAGUCAGGAGUGAAUACAAGGCU UCUUACAAUUCAGAGUCAUGUCCAGAAUUAUAUACCA AUAUCAUUGACACAACAAAUAUCGGAUCUUAGGAAAU UCAUUAGUGAAAUUACAAUUAGAAAUGAUAAUCAAGA AGUGCCACCACAAGAAUAACACAUGAUGUGGGUAUA AAACCUUUAAAUCCAGAUGAUUUCUGGAGAUGCACGU CUGGUCUUCCAUCUUUGAUGAAAACUCCAAAAAUAAG AUUAAUGCCGGGACCAGGAUUAUUAGCUAUGCCAACG ACUGUUGAUGGCUGUGUCAGAACCCCGUCCUUAGUGA UAAAUGAUCUGAUUUAUGCUUACACCUCAAAUCUAAU UACUCGAGGUUGCCAGGAUAUAGGGAAAUCAUAUCAA GUAUUACAGAUAGGGAUAAUAACUGUAAACUCAGACU UGGUACCUGACUUAAAUCCUAGGAUCUCUCAUACCUU CAACAUAAAUGACAAUAGAAAGUCAUGUUCUCUAGCA CUCCUAAAUACAGAUGUAUAUCAACUGUGUUCAACCC

61

|                          | TABLE 5 -continued   |                  |
|--------------------------|--|------------------|
| Description              | Sequence   | SEQ<br>ID<br>NO: |
|                          | CAAAAGUUGAUGAAAGAUCAGAUUAUGCAUCAUCAGG CAUAGAAGAUAUUGUACUUGAUAUUGUCAAUUAUGAU GGCUCAAUCUCGACAACAAGAUUUAAGAAUAAUAAUA UAAGUUUUGAUCAACAAGAUUUAAGAAUAAUAAUA UAAGUUUUGACCAGGGAUUAUACCCAUC UGUUGGACCAGGGGUUUGAACACAAGGCAAAAUAAUA UUUCUCGGGUAUGGACACAACUGGGUGUCCUGGGAA AACACAGAGAGACUGUGAACAUCUCAAUAAUG UGGUUUUCAGAUAGAAGGAUCUCAUAAUG UUGUUUCAGAUAGAAGGAUGUCAACUCUAUAAUUG UUGUUGACAAGGGCUUGAACUCCAUAAUGGAG GGUAUGGACGAUAUCUAAACUCUAUAAUG UGGUUUCAGAUAGAAGGAUCACUCUAUAAUG UAGAAAGGACGUUCAACACUCUAUAAUG ACAAAUGAACAGAGUUCAACAAUUACAGGGG UCAGAAGGACAUAUCUUCACAGGGAACAAAUUACUGGGGG UCAGAAGAAGAUUACUCAAGUUGCACAACAAGUU ACAAUUAGAAAAAUUACUCAAGUUGCACAACAAGUU ACAAUUAGGAACAAGAUCUACAAGUUGGCACACACAGAGUU ACAAUUAGGAACAACAGACACAGUCCACAGGAGUACACAGAGACACACAC |                  |
| HPIV3_HN_Codon Optimized | AUGGAAUACUGGAAGCACACCAACCACGCAACGACGCAACACGCCCCGGCAACGACG  | 63               |
| HPIV3_F_Codon            | CAAGACCGAGAUCCCCAAGAGCUGCUCC  AUGCCCAUCAGCAUCCUGCUGAUCAUCACCACAAUGAU   | 64               |

Optimized mRNA sequence

CAUGGCCAGCCACUGCCAGAUCGACAUCACCAAGCUGC AGCACGUGGGCGUGCUCGUGAACAGCCCCAAGGGCAU

Description

SEO

252

#### TABLE 5 -continued

ID Sequence NO: GAAGAUCAGCCAGAACUUCGAGACACGCUACCUGAUC CUGAGCCUGAUCCCCAAGAUCGAGGACAGCAACAGCU GCGGCGACCAGCAGAUCAAGCAGUACAAGCGGCUGCU GGACAGACUGAUCAUCCCCCUGUACGACGGCCUGCGGC UGCAGAAAGACGUGAUCGUGACCAACCAGGAAAGCAA CGAGAACACCGACCCCGGACCGAGAGAUUCUUCGGCG GCGUGAUCGGCACAAUCGCCCUGGGAGUGGCCACAAG CGCCCAGAUUACAGCCGCUGUGGCCCUGGUGGAAGCCA AGCAGGCCAGAAGCGACAUCGAGAAGCUGAAAGAGGC CAUCCGGGACACCAACAAGGCCGUGCAGAGCGUGCAG UCCAGCGUGGGCAAUCUGAUCGUGGCCAUCAAGUCCG UGCAGGACUACGUGAACAAAGAAAUCGUGCCCUCUAU CGCCCGGCUGGGCUGUGAAGCUGCCGGACUGCAGCUG GGCAUUGCCCUGACACAGCACUACAGCGAGCUGACCAA CAUCUUCGGCGACAACAUCGGCAGCCUGCAGGAAAAG GGCAUUAAGCUGCAGGGAAUCGCCAGCCUGUACCGCA CCAACAUCACCGAGAUCUUCACCACCAGCACCGUGGAU AAGUACGACAUCUACGACCUGCUGUUCACCGAGAGCA UCAAAGUGCGCGUGAUCGACGUGGACCUGAACGACUA CAGCAUCACCUGCAAGUGCGGCUGCCCCUGCUGACCA GACUGCUGAACACCCAGAUCUACAAGGUGGACAGCAU CUCCUACAACAUCCAGAACCGCGAGUGGUACAUCCCUC UGCCCAGCCACAUUAUGACCAAGGGCGCCUUUCUGGGC GGAGCCGACGUGAAAGAGUGCAUCGAGGCCUUCAGCA GCUACAUCUGCCCCAGCGACCCUGGCUUCGUGCUGAAC CACGAGAUGGAAAGCUGCCUGAGCGGCAACAUCAGCC AGUGCCCAGAACCACCGUGACCUCCGACAUCGUGCCC AGAUACGCCUUCGUGAAUGGCGGCGUGGUGGCCAACU GCAUCACCACCUGUACCUGCAACGGCAUCGGCAAC CGGAUCAACCAGCCUCCCGAUCAGGGCGUGAAGAUUA UCACCCACAAAGAGUGUAACACCAUCGGCAUCAACGGC AUGCUGUUCAAUACCAACAAAGAGGGCACCCUGGCCU UCUACACCCCGACGAUAUCACCCUGAACAACUCCGUG GCUCUGGACCCCAUCGACAUCUCCAUCGAGCUGAACAA GGCCAAGAGCGACCUGGAAGAGUCCAAAGAGUGGAUC CGGCGGAGCAACCAGAAGCUGGACUCUAUCGGCAGCU GGCACCAGAGCACCACCAUCAUCGUGAUCCUGAUU AUGAUGAUUAUCCUGUUCAUCAUCAACAUUACCAUCA UCACUAUCGCCAUUAAGUACUACCGGAUCCAGAAACG GAACCGGGUGGACCAGAAUGACAAGCCCUACGUGCUG ACAAACAAG

# TABLE 6

|   | PIV3 Amino Acid Sequences   |               |
|---|---|---------------|
| Description   | Sequence  | SEQ ID<br>NO: |
| >gi 612507166 gb <br>AHX22429.1 <br>fusion<br>glycoprotein<br>F0 [Human<br>parainfluenza<br>virus 3]      | MPISILLIITTMIMASHCQIDITKLQHVGVLVNSPKGMKISQ NFETRYLILSLIPKIEDSNSCGDQQIKQYKRLLDRLIIPLYDG LRLQKDVIVTNQESNENTDPRTERFFGGVIGTIALGVATSA QITAAVALVEAKQARSDIEKLKEAIRDTNKAVQSVQSSVG NLIVAIKSVQDYVNKEIVPSIARLGCEAAGLQLGIALTQHYS ELTNIFGDNIGSLQEKGIKLQGIASLYRTNITEIFTTSTVDKY DIYDLLFTESIKVRVIDVDLNDYSITLQVRLPLLTRLLNTQIY KVDSISYNIQNREWYIPLPSHIMTKGAFLGGADVKECIEAFS SYICPSDPGFVLNHEMESCLSGNISQCPRTTVTSDIVPRYAF VNGGVVANCITTTCTCNGIGNRINQPPDQGVKIITHKECNTI GINGMLFNTNKEGTLAFYTPDDITLNNSVALDPIDISIELNK AKSDLEESKEWIRRSNQKLDSIGSWHQSSTTIIVILIMMIILFI INITIITIAIKYYRIQKRNRVDQNDKPYVLTNK | 13            |
| gi 612507167 gb <br>AHX22430.1 <br>hemagglutinin-<br>neuraminidase<br>[Human<br>parainfluenza<br>virus 3] | MEYWKHTNHGKDAGNELETSTATHGNKLTNKITYILWTIT LVLLSIVFIIVLTNSIKSEKARESLLQDINNEFMEVTEKIQVA SDNTNDLIQSGVNTRLLTIQSHVQNYIPISLTQQISDLRKFIS EITIRNDNQEVPPQRITHDVGIKPLNPDDFWRCTSGLPSLMK TPKIRLMPGPGLLAMPTTVDGCVRTPSLVINDLIYAYTSNLI TRGCQDIGKSYQVLQIGIITVNSDLVPDLNPRISHTFNINDN RKSCSLALLNTDVYQLCSTPKVDERSDYASSGIEDIVLDIV NYDGSISTTRFKNNNISFDQPYAALYPSVGPGIYYKGKIIFL GYGGLEHPINENAICNTTGCPGKTQRDCNQASHSPWFSDR   | 14            |

# PIV3 Amino Acid Sequences SEQ ID Description Sequence SEQ ID RMVNSIIVVDKGLNSVPKLKVWTISMRQNYWGSEGRLLLL GNKIYIYTRSTSWHSKLQLGIIDITDYSDIRIKWTWHNVLSR PGNNECPWGHSCPDGCITGVYTDAYPLNPTGSIVSSVILDS QKSRVNPVITYSTATERVNELAIRNKTLSAGYTTTSCITHY NKGYCFHIVEINHKSLNTFQPMLFKTEIPKSCS

# TABLE 7

| Description  |   |
|--|---|
|  | GenBank Accession   |
| HPIV3/Homo sapiens/PER/FLA4815/2008  | KJ672601.1 :<br>4990-6609<br>AHX22429   |
| hemagglutinin-neuraminidase [Human parainfluenza virus 3]<br>HPIV3/Homo sapiens/PER/FLA4815/2008   | (Fusion protein)<br>KJ672601.1 :<br>6724-8442<br>AHX22430                         |
| Recombinant PIV3/PIV1 virus fusion glycoprotein (F) and hemagglutinin (HN) genes, complete cds; and RNA dependent RNA polymerase (L) gene, partial cds.  Recombinant PIV3/PIV1 virus fusion glycoprotein (F) and hemagglutinin (HN) genes, complete cds; and RNA dependent RNA polymerase (L) gene, partial cds. | (HN protein) AF016281 AAC23947 (hemagglutinin) AF016281 AAC23947 (fusion protein) |
| hemagglutinin-neuraminidase [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3]  | BAO32044.1<br>BAO32051.1<br>NP_599251.1<br>ABZ85670.1<br>AGZ8666.1                |
| C protein [Human parainfluenza virus 3]  | AAB48686.1<br>AHX22115.1<br>AGW51066.1<br>AGW51162.1<br>AGT75252.1<br>AGT75188.1  |
| C protein [Human parainfluenza virus 3]  | AGW51218.1<br>AGW51074.1<br>AGT75323.1<br>AGT75307.1<br>AHX22131.1                |
| C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3]  | AGW51243.1<br>AGT75180.1<br>AGT75212.1<br>AGW51186.1<br>AHX22075.1                |
| C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3]  | AHX22163.1<br>AGT75196.1<br>AHX22491.1<br>AHX22139.1<br>AGW51138.1                |
| C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] RecName: Full = Protein C; AltName: Full = VP18 protein  | AGW51114.1<br>AGT75220.1<br>AHX22251.1<br>P06165.1<br>AHX22187.1                  |
| C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3]  | AGT75228.1<br>AHX22179.1<br>AHX22427.1<br>AGW51210.1<br>BAA00922.1                |
| C protein [Human parainfluenza virus 3]  | AHX22315.1<br>AGW51259.1<br>AHX22435.1<br>AHX22123.1                              |
| C protein [Human parainfluenza virus 3] unnamed protein product [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3] C protein [Human parainfluenza virus 3]  | AHX22299.1<br>AGW51267.1<br>CAA28430.1<br>AGW51178.1<br>AHX22411.1<br>P06164.1    |

TABLE 7-continued

| PIV3 NCBI Accession Numbers (Nucleic Acid and Amin   | o Acid Sequences)         |
|--|---------------------------|
|  |                           |
| Description  | GenBank Accession         |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | NP_067149.1<br>AAB48685.1 |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22498.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22490.1<br>AGT75259.1  |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51137.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51145.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75298.1<br>AGW51113.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75203.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75163.1<br>AHX22506.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51129.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22194.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75211.1<br>AHX22258.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51121.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75282.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22146.1<br>AHX22138.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22322.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22370.1<br>AHX22098.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22130.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22418.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22114.1<br>AHX22410.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75306.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22170.1<br>AHX22266.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22200.1<br>AHX22090.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75195.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22226.1<br>AHX22178.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22122.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22186.1<br>AHX22066.1  |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22522.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51225.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | BAN29032.1<br>ABZ85669.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22426.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Simian Agent 10]             | AHX22058.1<br>ADR00400.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22250.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22434.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22298.1<br>AHX22442.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22074.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51153.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51241.1<br>AHX22210.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51105.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75251.1<br>AHX22362.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22474.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51217.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AIG60038.1<br>AHX22378.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51057.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75187.1<br>AGW51233.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22482.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51161.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AHX22306.1<br>AHX22162.1  |
| phosphoprotein [Human parainfluenza virus 3]   | ACJ70087.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22466.1<br>AHX22346.1  |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51089.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51073.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51185.1<br>AGW51065.1  |
| phosphoprotein [Human parainfluenza virus 3]   | ABY47603.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51049.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AHX22330.1                |

TABLE 7-continued

| DIV/2 MODI A   | - A -: 1 G                |
|--|---------------------------|
| PIV3 NCBI Accession Numbers (Nucleic Acid and Amin   | io Acid Sequences)        |
| Description  | GenBank Accession         |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51250.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75227.1<br>AGW51282.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51209.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51193.1<br>AGT75322.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75322.1<br>AGT75219.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51258.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51041.1<br>ACD99698.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51266.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGT75179.1<br>AHX22282.1  |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51169.1                |
| phosphoprotein [Human parainfluenza virus 3]   | AGW51274.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AGW51201.1<br>AGW51177.1  |
| RecName: Full = Phosphoprotein; Short = Protein P  | P06162.1                  |
| P protein [Human parainfluenza virus 3]  | AAA66818.1                |
| phosphoprotein [Human parainfluenza virus 3]<br>phosphoprotein [Human parainfluenza virus 3] | AAA46866.1<br>BAA00031.1  |
| polymerase-associated nucleocapsid phosphoprotein  | RRNZP5                    |
| (version 2) - parainfluenza virus type 3<br>[Human parainfluenza virus 3]                    |                           |
| phosphoprotein [Human parainfluenza virus 3]   | AGT75171.1                |
| phosphoprotein [Human parainfluenza virus 3]   | BAA00921.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | NP_599250.1<br>AHX22377.1 |
| D protein [Human parainfluenza virus 3]  | AHX22121.1                |
| D protein [Human parainfluenza virus 3]  | AGT75297.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGW51136.1<br>AGW51242.1  |
| D protein [Human parainfluenza virus 3]  | AGW51112.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AHX22497.1<br>AHX22145.1  |
| D protein [Human parainfluenza virus 3]  | AGT75202.1                |
| D protein [Human parainfluenza virus 3]  | AHX22385.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGW51216.1<br>AGT75281.1  |
| D protein [Human parainfluenza virus 3]  | AGT75194.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AHX22521.1<br>AGW51120.1  |
| D protein [Human parainfluenza virus 3]  | AGW31120.1<br>AGT75313.1  |
| D protein [Human parainfluenza virus 3]  | AHX22249.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AHX22097.1<br>AGW51144.1  |
| D protein [Human parainfluenza virus 3]  | AHX22089.1                |
| D protein [Human parainfluenza virus 3]  | AHX22225.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AHX22137.1<br>AHX22065.1  |
| D protein [Human parainfluenza virus 3]  | AGW51224.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGT75210.1<br>AHX22393.1  |
| D protein [Human parainfluenza virus 3]  | AGT75258.1                |
| D protein [Human parainfluenza virus 3]  | AHX22345.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGT75250.1<br>AHX22113.1  |
| D protein [Human parainfluenza virus 3]  | AGW51232.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AHX22057.1<br>AHX22209.1  |
| D protein [Human parainfluenza virus 3]  D protein [Human parainfluenza virus 3]             | AGW51056.1                |
| D protein [Human parainfluenza virus 3]  | AHX22161.1                |
| D protein [Simian Agent 10] D protein [Human parainfluenza virus 3]                          | ADR00402.1<br>AHX22361.1  |
| D protein [Human parainfluenza virus 3]  | AGW51281.1                |
| D protein [Human parainfluenza virus 3]  | AGW51184.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGW51160.1<br>AHX22465.1  |
| D protein [Human parainfluenza virus 3]  | AHX22329.1                |
| D protein [Human parainfluenza virus 3]  | AGW51064.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGW51040.1<br>AGT75226.1  |
| D protein [Human parainfluenza virus 3]  | AHX22425.1                |
| D protein [Human parainfluenza virus 3]  | AHX22305.1                |
| D protein [Human parainfluenza virus 3] D protein [Human parainfluenza virus 3]              | AGW51249.1<br>AHX22481.1  |
|  |                           |

TABLE 7-continued

| PIV3 NCBI Accession Numbers (Nucleic Acid a | nd Amino Acid Sequences) |
|---|--------------------------|
| Description                                 | GenBank Accession        |
| D protein [Human parainfluenza virus 3]     | AHX22281.1               |
| D protein [Human parainfluenza virus 3]     | AGW51048.1               |
| D protein [Human parainfluenza virus 3]     | AHX22297.1               |
| D protein [Human parainfluenza virus 3]     | AGW51088.1               |
| D protein [Human parainfluenza virus 3]     | AGT75305.1               |
| D protein [Human parainfluenza virus 3]     | AHX22185.1               |
| D protein [Human parainfluenza virus 3]     | AGW51104.1               |
| D protein [Human parainfluenza virus 3]     | AHX22081.1               |
| D protein [Human parainfluenza virus 3]     | AGW51192.1               |
| D protein [Human parainfluenza virus 3]     | AHX22489.1               |
| D protein [Human parainfluenza virus 3]     | AHX22441.1               |
| D protein [Human parainfluenza virus 3]     | AHX22409.1               |
| D protein [Human parainfluenza virus 3]     | AHX22369.1               |
| D protein [Human parainfluenza virus 3]     | AHX22321.1               |
| D protein [Human parainfluenza virus 3]     | AHX22073.1               |
| D protein [Human parainfluenza virus 3]     | AGW51152.1               |
| D protein [Human parainfluenza virus 3]     | AGW51072.1               |
| D protein [Human parainfluenza virus 3]     | AGT75321.1               |
| D protein [Human parainfluenza virus 3]     | AHX22257.1               |
| D protein [Human parainfluenza virus 3]     | AHX22129.1               |
| D protein [Human parainfluenza virus 3]     | AHX22417.1               |
| D protein [Human parainfluenza virus 3]     | AGT75218.1               |
| D protein [Human parainfluenza virus 3]     | AHX22265.1               |
| D protein [Human parainfluenza virus 3]     | AGT75178.1               |
| D protein [Human parainfluenza virus 3]     | AHX22433.1               |
| D protein [Human parainfluenza virus 3]     | AGW51273.1               |
| D protein [Human parainfluenza virus 3]     | AGW51208.1               |
| D protein [Human parainfluenza virus 3]     | AGT75170.1               |
| D protein [Human parainfluenza virus 3]     | AGT75162.1               |
| D protein [Human parainfluenza virus 3]     | AGW51257.1               |
| D protein [Human parainfluenza virus 3]     | AGW51200.1               |
| D protein [Human parainfluenza virus 3]     | AGW51176.1               |
| D protein [Human parainfluenza virus 3]     | AGT75186.1               |
| D protein [Human parainfluenza virus 3]     | AGW51265.1               |
| D protein [Human parainfluenza virus 3]     | AGW51168.1               |
|   |                          |

35

TABLE 9 TABLE 9

| TABBE 0                         |                          |               | IN ADDEE 9                                 |       |                                     |           |       |           |
|---------------------------------|--------------------------|---------------|--|-------|-------------------------------------|-----------|-------|-----------|
| Signal Peptides                 |                          |               | hMPV/PIV Cotton Rat Challenge Study Design |       |                                     |           |       |           |
| Description                     | Sequence                 | SEQ<br>ID NO: | 40   | Group | n Test Article                      | [conc]/μg | Route | Challenge |
| Harting of many                 | MEMBAGITELL              | 15            | -  | 1     | 5 Placebo                           | n/a       | IM    | hMPV/A2   |
| HuIgG <sub>k</sub> signal       | METPAQLLFLLL<br>LWLPDTTG | 15            |  | 2     | 5 hMPV vaccine mRNA                 | 30        | IM    | hMPV/A2   |
| peptide                         | LWLPDIIG                 |               |  | 3     | 5 hMPV vaccine mRNA                 | 15        | IM    | hMPV/A2   |
| IqE heavy chain                 | MDWTWILFLVAA             | 16            |  | 4     | 5 hMPV vaccine mRNA                 | 10        | IM    | hMPV/A2   |
| epsilon -1 signal peptide       | ATRVHS                   | 10            | 45   | 5     | 5 hMPV/PIV3 vaccine<br>mRNA (15/15) | 30        | IM    | hMPV/A2   |
| popozac                         |                          |               |  | 6     | 5 FI-hMPV                           | n/a       | IM    | hMPV/A2   |
| Japanese                        | MLGSNSGQRVVF             | 17            |  | 7     | 5 Placebo                           | n/a       | IM    | PIV3      |
| encephalitis PRM                | TILLLLVAPAYS             |               |  | 8     | 5 PIV3 vaccine mRNA                 | 30        | IM    | PIV3      |
| signal sequence                 |                          |               |  | 9     | 5 PIV3 vaccine mRNA                 | 15        | IM    | PIV3      |
|                                 |                          |               | 50   | 10    | 5 PIV3 vaccine mRNA                 | 10        | IM    | PIV3      |
| VSVg protein<br>signal sequence | MKCLLYLAFLFI<br>GVNCA    | 18            |  | 11    | 5 hMPV/PIV3 vaccine<br>mRNA (15/15) | 30        | IM    | PIV3      |
| Japanese<br>encephalitis JEV    | MWLVSLAIVTAC<br>AGA      | 19            |  | 12    | 5 FI-PIV3                           | n/a       | IM    | PIV3      |
| signal sequence                 |                          |               | 55   |       | 60                                  |           |       |           |

TABLE 10

|        |                       | SEQ ID |
|--------|-----------------------|--------|
| Strain | Nucleic Acid Sequence | NO:    |

Betacoronavirus Nucleic Acid Sequence

gb|KJ156934.1|: 21405-25466 Middle East respiratory  $\label{eq:total} \begin{minipage}{llll} ATGATACACTCAGTGTTTCTTGTTAACACC\\ TACAGAAAGTTACGTTGATGTAGGCCCAGATTCTGTTAAG\\ TCTGCTTGTATTGAGGTTGATATACAACAGACCTTCTTTGA\\ \end{minipage}$ 

SEO ID Strain Nucleic Acid Sequence NO:

Riyadh 14 2013, spike protein (nucleotide)

syndrome coronavirus TAAAACTTGGCCTAGGCCAATTGATGTTTCTAAGGCTGAC GGTATTATATACCCTCAAGGCCGTACATATTCTAACATAA CTATCACTTATCAAGGTCTTTTTCCCTATCAGGGAGACCAT GGTGATATGTTTTACTCTGCAGGACATGCTACAGGCA CAACTCCACAAAAGTTGTTTGTAGCTAACTATTCTCAGGA CGTCAAACAGTTTGCTAATGGGTTTGTCGTCCGTATAGGA GCAGCTGCCAATTCCACTGGCACTGTTATTATTAGCCCATC TACCAGCGCTACTATACGAAAAATTTACCCTGCTTTTATGC  $\tt TGGGTTCTTCAGTTGGTAATTTCTCAGATGGTAAAATGGG$ CCGCTTCTTCAATCATACTCTAGTTCTTTTGCCCGATGGAT GTGGCACTTTACTTAGAGCTTTTTATTGTATTCTAGAGCCT CGCTCTGGAAATCATTGTCCTGCTGGCAATTCCTATACTTC TTTTGCCACTTATCACACTCCTGCAACAGATTGTTCTGATG GCAATTACAATCGTAATGCCAGTCTGAACTCTTTTAAGGA GTATTTTAATTTACGTAACTGCACCTTTATGTACACTTATA ACATTACCGAAGATGAGATTTTAGAGTGGTTTGGCATTAC ACAAACTGCTCAAGGTGTTCACCTCTTCTCATCTCGGTATG TTGATTTGTACGGCGGCAATATGTTTCAATTTGCCACCTTG CCTGTTTATGATACTATTAAGTATTATTCTATCATTCCTCA CAGTATTCGTTCTATCCAAAGTGATAGAAAAGCTTGGGCT GCCTTCTACGTATATAAACTTCAACCGTTAACTTTCCTGTT GGATTTTTCTGTTGATGGTTATATACGCAGAGCTATAGACT GTGGTTTTAATGATTTGTCACAACTCCACTGCTCATATGAA TCCTTCGATGTTGAATCTGGAGTTTATTCAGTTTCGTCTTT CGAAGCAAAACCTTCTGGCTCAGTTGTGGAACAGGCTGAA GGTGTTGAATGTGATTTTTCACCTCTTCTGTCTGGCACACC TCCTCAGGTTTATAATTTCAAGCGTTTGGTTTTTACCAATT GCAATTATAATCTTACCAAATTGCTTTCACTTTTTTCTGTG AATGATTTTACTTGTAGTCAAATATCTCCAGCAGCAATTGC TAGCAACTGTTATTCTTCACTGATTTTTGGATTATTTTTCAT ACCCACTTAGTATGAAATCCGATCTCAGTGTTAGTTCTGCT GGTCCAATATCCCAGTTTAATTATAAACAGTCCTTTTCTAA  ${\tt TCCCACATGTTTGATCTTAGCGACTGTTCCTCATAACCTTA}$ CTACTATTACTAAGCCTCTTAAGTACAGCTATATTAACAA GTGCTCTCGTCTTCTTTCTGATGATCGTACTGAAGTACCTC AGTTAGTGAACGCTAATCAATACTCACCCTGTGTATCCATT GTCCCATCCACTGTGTGGGAAGACGGTGATTATTATAGGA AACAACTATCTCCACTTGAAGGTGGTGGCTGGCTTGTTGC TAGTGGCTCAACTGTTGCCATGACTGAGCAATTACAGATG GGCTTTGGTATTACAGTTCAATATGGTACAGACACCAATA GTGTTTGCCCCAAGCTTGAATTTGCTAATGACACAAAAAT  $\tt TGCCTCTCAATTAGGCAATTGCGTGGAATATTCCCTCTATG$ GTGTTTCGGGCCGTGGTGTTTTTCAGAATTGCACAGCTGTA  $\tt GGTGTTCGACAGCAGCGCTTTGTTTATGATGCGTACCAGA$ ATTTAGTTGGCTATTATTCTGATGATGGCAACTACTACTGT CTGCGTGCTTGTGTTAGTGTTCCTGTTTCTGTCATCTATGA  ${\tt TAAAGAAACTAAAACCCACGCTACTCTATTTGGTAGTGTT}$ GCATGTGAACACATTTCTTCTACCATGTCTCAATACTCCCG TTCTACGCGATCAATGCTTAAACGGCGAGATTCTACATAT GGCCCCCTTCAGACACCTGTTGGTTGTGTCCTAGGACTTGT TAATTCCTCTTTGTTCGTAGAGGACTGCAAGTTGCCTCTCG GTCAATCTCTGTGCTCTTCCTGACACACCTAGTACTCTC ACACCTCGCAGTGTGCGCTCTGTGCCAGGTGAAATGCGCT TGGCATCCATTGCTTTTAATCATCCCATTCAGGTTGATCAA CTTAATAGTAGTTATTTTAAATTAAGTATACCCACTAATTT TTCCTTTGGTGTGACTCAGGAGTACATTCAGACAACCATTC AGAAAGTTACTGTTGATTGTAAACAGTACGTTTGCAATGG TTTCCAGAAGTGTGAGCAATTACTGCGCGAGTATGGCCAG TTTTGTTCCAAAATAAACCAGGCTCTCCATGGTGCCAATTT ACGCCAGGATGATTCTGTACGTAATTTGTTTGCGAGCGTG AAAAGCTCTCAATCATCTCCTATCATACCAGGTTTTGGAG GTGACTTTAATTTGACACTTCTAGAACCTGTTTCTATATCT ACTGGCAGTCGTAGTGCACGTAGTGCTATTGAGGATTTGC TATTTGACAAAGTCACTATAGCTGATCCTGGTTATATGCA AGGTTACGATGATTGTATGCAGCAAGGTCCAGCATCAGCT CGTGATCTTATTTGTGCTCAATATGTGGCTGGTTATAAAGT ATTACCTCCTCTTATGGATGTTAATATGGAAGCCGCGTATA CTTCATCTTTGCTTGGCAGCATAGCAGGTGTTGGCTGGACT GCTGGCTTATCCTCCTTTGCTGCTATTCCATTTGCACAGAG TATYTTTATAGGTTAAACGGTGTTGGCATTACTCAACAG GTTCTTTCAGAGAACCAAAAGCTTATTGCCAATAAGTTTA ATCAGGCTCTGGGAGCTATGCAAACAGGCTTCACTACAAC TAATGAAGCTTTTCGGAAGGTTCAGGATGCTGTGAACAAC AATGCACAGGCTCTATCCAAATTAGCTAGCGAGCTATCTA ATACTTTTGGTGCTATTTCCGCCTCTATTGGAGACATCATA CAACGTCTTGATGTTCTCGAACAGGACGCCCAAATAGACA GACTTATTAATGGCCGTTTGACAACACTAAATGCTTTTGTT

TABLE 10-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GCACAGCAGCTTGTTCGTTCCGAATCAGCTGCTCTTTCCGC TCAATTGGCTAAAGATAAAGTCAATGAGTGTGTCAAGGCA CAATCCAAGCGTTCTGGATTTTGCGGTCAAGGCACACATA TAGTGTCCTTTGTTGTAAATGCCCCTAATGGCCTTTACTTT ATGCATGTTGGTTATTACCCTAGCAACCACATTGAGGTTGT TTCTGCTTATGGTCTTTGCGATGCAGCTAACCCTACTAATT GTATAGCCCCTGTTAATGGCTACTTTATTAAAACTAATAAC ACTAGGATTGTTGATGAGTGGTCATATACTGGCTCGTCCTT CTATGCACCTGAGCCCATCACCTCTCTTAATACTAAGTATG TTGCACCACAGGTGACATACCAAAACATTTCTACTAACCT CCCTCCTCTCTCGGCAATTCCACCGGGATTGACTTCC AAGATGAGTTGGATGAGTTTTTCAAAAATGTTAGCACCAG TATACCTAATTTTGGTTCTCTAACACAGATTAATACTACAT TACTCGATCTTACCTACGAGATGTTGTCTCTTCAACAAGTT GTTAAAGCCCTTAATGAGTCTTACATAGACCTTAAAGAGC TTGGCAATTATACTTATTACAACAAATGGCCGTGGTACAT TTGGCTTGGTTTCATTGCTGGGCTTGTTGCCTTAGCTCTAT GCGTCTTCTTCATACTGTGCTGCACTGGTTGTGGCACAAAC TGTATGGGAAAACTTAAGTGTAATCGTTGTTGTGATAGAT ACGAGGAATACGACCTCGAGCCGCATAAGGTTCATGTTCA CTAA

MERS S FL SPIKE 2cEMC/2012 (XBaI change (T to G)) (nucleotide) ATGATACACTCAGTGTTTCTACTGATGTTCTTGTTAACACC TACAGAAAGTTACGTTGATGTAGGGCCAGATTCTGTTAAG TCTGCTTGTATTGAGGTTGATATACAACAGACTTTCTTTGA TAAAACTTGGCCTAGGCCAATTGATGTTTCTAAGGCTGAC GGTATTATATACCCTCAAGGCCGTACATATTCTAACATAA CTATCACTTATCAAGGTCTTTTTCCCTATCAGGGAGACCAT GGTGATATGTTTACTCTGCAGGACATGCTACAGGCA CAACTCCACAAAAGTTGTTTGTAGCTAACTATTCTCAGGA  $\tt CGTCAAACAGTTTGCTAATGGGTTTGTCGTCCGTATAGGA$ GCAGCTGCCAATTCCACTGGCACTGTTATTATTAGCCCATC  ${\tt TACCAGCGCTACTATACGAAAAATTTACCCTGCTTTTATGC}$  $\tt TGGGTTCTTCAGTTGGTAATTTCTCAGATGGTAAAATGGG$  $\tt CCGCTTCTTCAATCATACTCTAGTTCTTTTGCCCGATGGAT$  $\tt GTGGCACTTTACTTAGAGCTTTTTATTGTATTCTGGAGCCT$ CGCTCTGGAAATCATTGTCCTGCTGGCAATTCCTATACTTC  ${\tt TTTTGCCACTTATCACACTCCTGCAACAGATTGTTCTGATG}$ GCAATTACAATCGTAATGCCAGTCTGAACTCTTTTAAGGA GTATTTTAATTTACGTAACTGCACCTTTATGTACACTTATA ACATTACCGAAGATGAGATTTTAGAGTGGTTTGGCATTAC ACAAACTGCTCAAGGTGTTCACCTCTTCTCATCTCGGTATG TTGATTTGTACGGCGGCAATATGTTTCAATTTGCCACCTTG CCTGTTTATGATACTATTAAGTATTATTCTATCATTCCTCA CAGTATTCGTTCTATCCAAAGTGATAGAAAAGCTTGGGCT GCCTTCTACGTATATAAACTTCAACCGTTAACTTTCCTGTT  $\tt GGATTTTTCTGTTGATGGTTATATACGCAGAGCTATAGACT$ GTGGTTTTAATGATTTGTCACAACTCCACTGCTCATATGAA  ${\tt TCCTTCGATGTTGAATCTGGAGTTTATTCAGTTTCGTCTTT}$  $\tt CGAAGCAAAACCTTCTGGCTCAGTTGTGGAACAGGCTGAA$ GGTGTTGAATGTGATTTTTCACCTCTTCTGTCTGGCACACC TCCTCAGGTTTATAATTTCAAGCGTTTGGTTTTTACCAATT GCAATTATAATCTTACCAAATTGCTTTCACTTTTTTCTGTG AATGATTTTACTTGTAGTCAAATATCTCCAGCAGCAATTGC TAGCAACTGTTATTCTTCACTGATTTTGGATTACTTTTCAT ACCCACTTAGTATGAAATCCGATCTCAGTGTTAGTTCTGCT GGTCCAATATCCCAGTTTAATTATAAACAGTCCTTTTCTAA TCCCACATGTTTGATTTTAGCGACTGTTCCTCATAACCTTA CTACTATTACTAAGCCTCTTAAGTACAGCTATATTAACAA GTGCTCTCGTCTTCTTCTGATGATCGTACTGAAGTACCTC AGTTAGTGAACGCTAATCAATACTCACCCTGTGTATCCATT GTCCCATCCACTGTGTGGGAAGACGGTGATTATTATAGGA AACAACTATCTCCACTTGAAGGTGGTGGCTGGCTTGTTGC TAGTGGCTCAACTGTTGCCATGACTGAGCAATTACAGATG GGCTTTGGTATTACAGTTCAATATGGTACAGACACCAATA GTGTTTGCCCCAAGCTTGAATTTGCTAATGACACAAAAAT TGCCTCTCAATTAGGCAATTGCGTGGAATATTCCCTCTATG GTGTTTCGGGCCGTGGTGTTTTTCAGAATTGCACAGCTGTA GGTGTTCGACAGCAGCGCTTTGTTTATGATGCGTACCAGA ATTTAGTTGGCTATTATTCTGATGATGGCAACTACTACTGT TTGCGTGCTTGTGTTAGTGTTCCTGTTTCTGTCATCTATGAT AAAGAAACTAAAACCCACGCTACTCTATTTGGTAGTGTTG CATGTGAACACATTTCTTCTACCATGTCTCAATACTCCCGT TCTACGCGATCAATGCTTAAACGGCGAGATTCTACATATG  ${\tt GCCCCTTCAGACACCTGTTGGTTGTGTCCTAGGACTTGTT}$ AATTCCTCTTTGTTCGTAGAGGACTGCAAGTTGCCTCTTGG TCAATCTCTGTGCTCTTCCTGACACACCTAGTACTCTCA

SEQ ID Strain Nucleic Acid Sequence NO:

CACCTCGCAGTGTGCGCTCTGTTCCAGGTGAAATGCGCTT GGCATCCATTGCTTTTAATCATCCTATTCAGGTTGATCAAC TTAATAGTAGTTATTTTAAATTAAGTATACCCACTAATTTT TCCTTTGGTGTGACTCAGGAGTACATTCAGACAACCATTC AGAAAGTTACTGTTGATTGTAAACAGTACGTTTGCAATGG TTTCCAGAAGTGTGAGCAATTACTGCGCGAGTATGGCCAG  ${\tt TTTTGTTCCAAAATAAACCAGGCTCTCCATGGTGCCAATTT}$ ACGCCAGGATGATTCTGTACGTAATTTGTTTGCGAGCGTG AAAAGCTCTCAATCATCTCCTATCATACCAGGTTTTGGAG GTGACTTTAATTTGACACTTCTGGAACCTGTTTCTATATCT ACTGGCAGTCGTAGTGCACGTAGTGCTATTGAGGATTTGC TATTTGACAAAGTCACTATAGCTGATCCTGGTTATATGCA AGGTTACGATGATTGCATGCAGCAAGGTCCAGCATCAGCT CGTGATCTTATTTGTGCTCAATATGTGGCTGGTTACAAAGT ATTACCTCCTCTTATGGATGTTAATATGGAAGCCGCGTATA CTTCATCTTTGCTTGGCAGCATAGCAGGTGTTGGCTGGACT GCTGGCTTATCCTCCTTTGCTGCTATTCCATTTGCACAGAG TATCTTTATAGGTTAAACGGTGTTGGCATTACTCAACAGG TTCTTTCAGAGAACCAAAAGCTTATTGCCAATAAGTTTAA TCAGGCTCTGGGAGCTATGCAAACAGGCTTCACTACAACT AATGAAGCTTTTCAGAAGGTTCAGGATGCTGTGAACAACA ATGCACAGGCTCTATCCAAATTAGCTAGCGAGCTATCTAA TACTTTTGGTGCTATTTCCGCCTCTATTGGAGACATCATAC AACGTCTTGATGTTCTCGAACAGGACGCCCAAATAGACAG ACTTATTAATGGCCGTTTGACAACACTAAATGCTTTTGTTG CACAGCAGCTTGTTCGTTCCGAATCAGCTGCTCTTTCCGCT CAATTGGCTAAAGATAAAGTCAATGAGTGTGTCAAGGCAC AATCCAAGCGTTCTGGATTTTGCGGTCAAGGCACACATAT AGTGTCCTTTGTTGTAAATGCCCCTAATGGCCTTTACTTCA TGCATGTTGGTTATTACCCTAGCAACCACATTGAGGTTGTT  ${\tt TCTGCTTATGGTCTTTGCGATGCAGCTAACCCTACTAATTG}$ TATAGCCCCTGTTAATGGCTACTTTATTAAAACTAATAACA  $\tt CTAGGATTGTTGATGAGTGGTCATATACTGGCTCGTCCTTC$ TATGCACCTGAGCCCATTACCTCCCTTAATACTAAGTATGT  ${\tt TGCACCACAGGTGACATACCAAAACATTTCTACTAACCTC}$  $\tt CCTCCTCTTCTCGGCAATTCCACCGGGATTGACTTCCA$ AGATGAGTTGGATGAGTTTTTCAAAAATGTTAGCACCAGT ATACCTAATTTTGGTTCCCTAACACAGATTAATACTACATT ACTCGATCTTACCTACGAGATGTTGTCTCTTCAACAAGTTG TTAAAGCCCTTAATGAGTCTTACATAGACCTTAAAGAGCT TGGCAATTATACTTATTACAACAAATGGCCGTGGTACATT  $\tt TGGCTTGGTTTCATTGCTGGGCTTGTTGCCTTAGCTCTATG$ CGTCTTCTTCATACTGTGCTGCACTGGTTGTGGCACAAACT CGAGGAATACGACCTCGAGCCGCATAAGGTTCATGTTCAC

Novel\_MERS\_S2\_subunit\_trimeric vaccine (nucleotide) ATGATCCACTCCGTGTTCCTCCTCATGTTCCTGTTGACCCC  ${\tt CACTGAGTCAGACTGCAAGCTCCCGCTGGGACAGTCCCTG}$ TGTGCGCTGCCTGACACTCCTAGCACTCTGACCCCACGCTC  $\tt CGTGCGGTCGGTGCCTGGCGAAATGCGGCTGGCCTCCATC$ GCCTTCAATCACCCAATCCAAGTGGATCAGCTGAATAGCT CGTATTTCAAGCTGTCCATCCCCACGAACTTCTCGTTCGGG GTCACCCAGGAGTACATCCAGACCACAATTCAGAAGGTCA CCGTCGATTGCAAGCAATACGTGTGCAACGGCTTCCAGAA GTGCGAGCAGCTGCTGAGAGAATACGGGCAGTTTTGCAGC AAGATCAACCAGGCGCTGCATGGAGCTAACTTGCGCCAGG ACGACTCCGTGCGCAACCTCTTTGCCTCTGTGAAGTCATCC CAGTCCTCCCCAATCATCCCGGGGTTTCGGAGGGGACTTCA ACCTGACCCTCCTGGAGCCCGTGTCGATCAGCACCGGTAG CAGATCGGCGCTCAGCCATTGAAGATCTTCTGTTCGAC AAGGTCACCATCGCCGATCCGGGCTACATGCAGGGATACG ACGACTGTATGCAGCAGGGACCAGCCTCCGCGAGGGACCT CATCTGCGCGCAATACGTGGCCGGGTACAAAGTGCTGCCT CCTCTGATGGATGTGAACATGGAGGCCGCTTATACTTCGT CCCTGCTCGGCTCTATCGCCGGCGTGGGGTGGACCGCCGG CCTGTCCTCCTTCGCCGCTATCCCCTTTGCACAATCCATTT TCTACCGGCTCAACGGCGTGGGCATTACTCAACAAGTCCT GTCGGAGAACCAGAAGTTGATCGCAAACAAGTTCAATCA GGCCCTGGGGGCCATGCAGACTGGATTCACTACGACTAAC GAAGCGTTCCAGAAGGTCCAGGACGCTGTGAACAACAAC GCCCAGGCGCTCTCAAAGCTGGCCTCCGAACTCAGCAACA CCTTCGGAGCCATCAGCGCATCGATCGGTGACATAATTCA GCGGCTGGACGTGCTGGAGCAGGACGCCCAGATCGACCG  ${\tt CCTCATCAACGGACGGCTGACCACCTTGAATGCCTTCGTG}$ GCACAACAGCTGGTCCGGAGCGAATCAGCGGCACTTTCCG CCCAACTCGCCAAGGACAAAGTCAACGAATGCGTGAAGG

SEQ ID Strain Nucleic Acid Sequence NO:

CCCAGTCCAAGAGGTCCGGTTTCTGCGGTCAAGGAACCCA TATTGTGTCCTTCGTCGTGAACGCCCCAACGGTCTGTACT TTATGCACGTCGGCTACTACCCGAGCAATCATATCGAAGT GGTGTCCGCCTACGGCCTGTGCGATGCCGCTAACCCCACT AACTGTATTGCCCCTGTGAACGGATATTTTATTAAGACCA ACAACACCCGCATTGTGGACGAATGGTCATACACCGGTTC GTCCTTCTACGCGCCCGAGCCCATCACTTCACTGAACACC AAATACGTGGCTCCGCAAGTGACCTACCAGAACATCTCCA CCAATTTGCCGCCGCCGCTGCTCGGAAACAGCACCGGAAT TGATTTCCAAGATGAACTGGACGAATTCTTCAAGAACGTG TCCACTTCCATTCCCAACTTCGGAAGCCTGACACAGATCA ACACCACCCTTCTCGACCTGACCTACGAGATGCTGAGCCT TCAACAAGTGGTCAAGGCCCTGAACGAGAGCTACATCGAC CTGAAGGAGCTGGGCAACTATACCTACTACAACAAGTGGC CGGACAAGATTGAGGAGATTCTGTCGAAAATCTACCACAT TGAAAACGAGATCGCCAGAATCAAGAAGCTTATCGGCGA AGCC

MERS\_SO\_Fulllength Spike protein (nucleotide, codon optimized) ATGGAAACCCCTGCCCAGCTGCTGTTCCTGCTGCTGCTGTG GCTGCCTGATACCACCGGCAGCTATGTGGACGTGGGCCCC GATAGCGTGAAGTCCGCCTGTATCGAAGTGGACATCCAGC AGACCTTTTTCGACAAGACCTGGCCCAGACCCATCGACGT GTCCAAGGCCGACGCATCATCTATCCACAAGGCCGGACC TACAGCAACATCACCATTACCTACCAGGGCCTGTTCCCAT ATCAAGGCGACCACGGCGATATGTACGTGTACTCTGCCGG CCACGCCACCGGCACCACACCCCAGAAACTGTTCGTGGCC AACTACAGCCAGGACGTGAAGCAGTTCGCCAACGGCTTCG TCGTGCGGATTGGCGCCGCTGCCAATAGCACCGGCACAGT GATCATCAGCCCCAGCACCAGCGCCACCATCCGGAAGATC TACCCCGCCTTCATGCTGGGCAGCTCCGTGGGCAATTTCA  $\tt GCGACGCCAGATGGGCCGGTTCTTCAACCACACCCTGGT$ GCTGCTGCCCGATGGCTGTGGCACACTGCTGAGAGCCTTC TACTGCATCCTGGAACCCAGAAGCGGCAACCACTGCCCTG CCGGCAATAGCTACACCAGCTTCGCCACCTACCACACCC CGCCACCGATTGCTCCGACGGCAACTACAACCGGAACGCC AGCCTGAACAGCTTCAAAGAGTACTTCAACCTGCGGAACT GCACCTTCATGTACACCTACAATATCACCGAGGACGAGAT CCTGGAATGGTTCGGCATCACCCAGACCGCCCAGGGCGTG CACCTGTTCAGCAGCAGATACGTGGACCTGTACGGCGGCA ACATGTTCCAGTTTGCCACCCTGCCCGTGTACGACACCATC AAGTACTACAGCATCATCCCCCACAGCATCCGGTCCATCC AGAGCGACAGAAAAGCCTGGGCCGCCTTCTACGTGTACAA GCTGCAGCCCTGACCTTCCTGCTGGACTTCAGCGTGGAC  $\tt GGCTACATCAGACGGGCCATCGACTGCGGCTTCAACGACC$ TGAGCCAGCTGCACTGCTCCTACGAGAGCTTCGACGTGGA AAGCGGCGTGTACAGCGTGTCCAGCTTCGAGGCCAAGCCT  ${\tt AGCGGCAGCGTGGTGGAACAGGCTGAGGGCGTGGAATGC}$ GACTTCAGCCCTCTGCTGAGCGGCACCCCTCCCCAGGTGT ACAACTTCAAGCGGCTGGTGTTCACCAACTGCAATTACAA CCTGACCAAGCTGCTGAGCCTGTTCTCCGTGAACGACTTC ACCTGTAGCCAGATCAGCCCTGCCGCCATTGCCAGCAACT GCTACAGCAGCCTGATCCTGGACTACTTCAGCTACCCCCT GAGCATGAAGTCCGATCTGAGCGTGTCCTCCGCCGGACCC ATCAGCCAGTTCAACTACAAGCAGAGCTTCAGCAACCCTA CCTGCCTGATTCTGGCCACCGTGCCCCACAATCTGACCAC CATCACCAAGCCCCTGAAGTACAGCTACATCAACAAGTGC AGCAGACTGCTGTCCGACGACCGGACCGAAGTGCCCCAGC TCGTGAACGCCAACCAGTACAGCCCCTGCGTGTCCATCGT GCCCAGCACCGTGTGGGAGGACGGCGACTACTACAGAAA GCAGCTGAGCCCCCTGGAAGGCGGCGGATGGCTGGTGGCT TCTGGAAGCACAGTGGCCATGACCGAGCAGCTGCAGATG GGCTTTGGCATCACCGTGCAGTACGGCACCGACACCAACA GCGTGTGCCCCAAGCTGGAATTCGCCAATGACACCAAGAT CGCCAGCCAGCTGGGAAACTGCGTGGAATACTCCCTGTAT GGCGTGTCCGGACGGGGCGTGTTCCAGAATTGCACAGCAG TGGGAGTGCGGCAGCAGAGATTCGTGTACGATGCCTACCA GAACCTCGTGGGCTACTACAGCGACGACGGCAATTACTAC TGCCTGCGGGCCTGTGTGTCCGTGCCCGTGTCCGTGATCTA CGACAAAGACAAAGACCCACGCCACACTGTTCGGCTCC GTGGCCTGCGAGCACATCAGCTCCACCATGAGCCAGTACT CCCGCTCCACCCGGTCCATGCTGAAGCGGAGAGATAGCAC CTACGGCCCCTGCAGACACCTGTGGGATGTGTGCTGGGC CTCGTGAACAGCTCCCTGTTTGTGGAAGATTGCAAGCTGC CCCTGGGCCAGAGCCTGTGTGCCCTGCCAGATACCCCTAG CACCCTGACCCCTAGAAGCGTGCGCTCTGTGCCCGGCGAA ATGCGGCTGGCCTCTATCGCCTTCAATCACCCCATCCAGGT GGACCAGCTGAACTCCAGCTACTTCAAGCTGAGCATCCCC

SEQ ID Strain Nucleic Acid Sequence NO:

ACCAACTTCAGCTTCGGCGTGACCCAGGAGTACATCCAGA CCACAATCCAGAAAGTGACCGTGGACTGCAAGCAGTACGT GTGCAACGGCTTTCAGAAGTGCGAACAGCTGCTGCGCGAG TACGGCCAGTTCTGCAGCAAGATCAACCAGGCCCTGCACG GCGCCAACCTGAGACAGGATGACAGCGTGCGGAACCTGTT CGCCAGCGTGAAAAGCAGCCAGTCCAGCCCCATCATCCCT GGCTTCGGCGGCGACTTTAACCTGACCCTGCTGGAACCTG TGTCCATCAGCACCGGCTCCAGAAGCGCCAGATCCGCCAT CGAGGACCTGCTGTTCGACAAAGTGACCATTGCCGACCCC GGCTACATGCAGGGCTACGACGATTGCATGCAGCAGGGCC CAGCCAGCGCCAGGGATCTGATCTGTGCCCAGTATGTGGC CGGCTACAAGGTGCTGCCCCCCCTGATGGACGTGAACATG GAAGCCGCCTACACCTCCAGCCTGCTGGGCTCTATTGCTG GCGTGGGATGGACAGCCGGCCTGTCTAGCTTTGCCGCCAT CCCTTTCGCCCAGAGCATCTTCTACCGGCTGAACGGCGTG GGCATCACACACAGGTGCTGAGCGAGAACCAGAAGCTG ATCGCCAACAAGTTTAACCAGGCACTGGGCGCCATGCAGA CCGGCTTCACCACCACCAACGAGGCCTTCAGAAAGGTGCA GGACGCCGTGAACAACAACGCCCAGGCTCTGAGCAAGCT GGCCTCCGAGCTGAGCAATACCTTCGGCGCCATCAGCGCC TCCATCGGCGACATCATCCAGCGGCTGGACGTGCTGGAAC AGGACGCCCAGATCGACCGGCTGATCAACGGCAGACTGA CCACCCTGAACGCCTTCGTGGCACAGCAGCTCGTGCGGAG CGAATCTGCCGCTCTGTCTGCTCAGCTGGCCAAGGACAAA GTGAACGAGTGCGTGAAGGCCCAGTCCAAGCGGAGCGGC TTTTGTGGCCAGGGCACCCACATCGTGTCCTTCGTCGTGAA TGCCCCCAACGGCCTGTACTTTATGCACGTGGGCTATTACC CCAGCAACCACATCGAGGTGGTGTCCGCCTATGGCCTGTG  $\tt CGACGCCGCCAATCCTACCAACTGTATCGCCCCCGTGAAC$ GGCTACTTCATCAAGACCAACACCCCGGATCGTGGACG AGTGGTCCTACACAGGCAGCAGCTTCTACGCCCCCGAGCC CATCACCTCCCTGAACACCAAATACGTGGCCCCCCAAGTG ACATACCAGAACATCTCCACCAACCTGCCCCCTCCACTGC TGGGAAATTCCACCGGCATCGACTTCCAGGACGAGCTGGA  $\tt CGAGTTCTTCAAGAACGTGTCCACCTCCATCCCCAACTTCG$ GCAGCCTGACCCAGATCAACACCACTCTGCTGGACCTGAC CTACGAGATGCTGTCCCTGCAACAGGTCGTGAAAGCCCTG AACGAGAGCTACATCGACCTGAAAGAGCTGGGGAACTAC ACCTACTACAACAAGTGGCCTTGGTACATTTGGCTGGGCT TTATCGCCGGCCTGGTGGCCCTGTGCGTGTTCTTC ATCCTGTGCTGCACCGGCTGCGGCACCAATTGCATGGGCA AGCTGAAATGCAACCGGTGCTGCGACAGATACGAGGAAT ACGACCTGGAACCTCACAAAGTGCATGTGCAC

#### Betacoronavirus mRNA Sequences

gb|KJ156934.1|: 21405-25466 Middle East respiratory syndrome coronavirus isolate Riyadh\_14\_2013, spike protein (nucleotide) AUGAUACACUCAGUGUUUCUACUGAUGUUCUUGUUAAC ACCUACAGAAAGUUACGUUGAUGUAGGGCCAGAUUCUG UUAAGUCUGCUUGUAUUGAGGUUGAUAUACAACAGACC UUCUUUGAUAAAACUUGGCCUAGGCCAAUUGAUGUUUC UAAGGCUGACGGUAUUAUAUACCCUCAAGGCCGUACAU AUUCUAACAUAACUAUCACUUAUCAAGGUCUUUUUCCCU AUCAGGGAGACCAUGGUGAUAUGUAUGUUUACUCUGCA GGACAUGCUACAGGCACAACUCCACAAAAGUUGUUUGU AGCUAACUAUUCUCAGGACGUCAAACAGUUUGCUAAUG GGUUUGUCGUCCGUAUAGGAGCAGCUGCCAAUUCCACUG GCACUGUUAUUAUUAGCCCAUCUACCAGCGCUACUAUAC GAAAAAUUUACCCUGCUUUUAUGCUGGGUUCUUCAGUU GGUAAUUUCUCAGAUGGUAAAAUGGGCCGCUUCUUCAA UCAUACUCUAGUUCUUUUGCCCGAUGGAUGUGGCACUU UACUUAGAGCUUUUUAUUGUAUUCUAGAGCCUCGCUCU GGAAAUCAUUGUCCUGCUGGCAAUUCCUAUACUUCUUU UGCCACUUAUCACACUCCUGCAACAGAUUGUUCUGAUGG CAAUUACAAUCGUAAUGCCAGUCUGAACUCUUUUAAGG AGUAUUUUAAUUUACGUAACUGCACCUUUAUGUACACU UAUAACAUUACCGAAGAUGAGAUUUUUAGAGUGGUUUGG CAUUACACAAACUGCUCAAGGUGUUCACCUCUUCUCAUC UCGGUAUGUUGAUUUGUACGGCGGCAAUAUGUUUCAAU UUGCCACCUUGCCUGUUUAUGAUACUAUUAAGUAUUAU UCUAUCAUUCCUCACAGUAUUCGUUCUAUCCAAAGUGAU AGAAAAGCUUGGGCUGCCUUCUACGUAUAUAAACUUCA ACCGUUAACUUUCCUGUUGGAUUUUUUCUGUUGAUGGUU AUAUACGCAGAGCUAUAGACUGUGGUUUUAAUGAUUUG UCACAACUCCACUGCUCAUAUGAAUCCUUCGAUGUUGAA UCUGGAGUUUAUUCAGUUUCGUCUUUCGAAGCAAAACC UUCUGGCUCAGUUGUGGAACAGGCUGAAGGUGUUGAAU GUGAUUUUUCACCUCUUCUGUCUGGCACACCUCCUCAGG

Strain Nucleic Acid Sequence SEQ ID NO:

UUUAUAAUUUCAAGCGUUUGGUUUUUACCAAUUGCAAU UAUAAUCUUACCAAAUUGCUUUCACUUUUUUCUGUGAA UGAUUUUACUUGUAGUCAAAUAUCUCCAGCAGCAAUUG CUAGCAACUGUUAUUCUUCACUGAUUUUUGGAUUAUUUU UCAUACCCACUUAGUAUGAAAUCCGAUCUCAGUGUUAG UUCUGCUGGUCCAAUAUCCCAGUUUAAUUAUAAACAGU CCUUUUCUAAUCCCACAUGUUUGAUCUUAGCGACUGUUC CUCAUAACCUUACUACUAUUACUAAGCCUCUUAAGUACA GCUAUAUUAACAAGUGCUCUCGUCUUCUUUCUGAUGAU CGUACUGAAGUACCUCAGUUAGUGAACGCUAAUCAAUA CUCACCCUGUGUAUCCAUUGUCCCAUCCACUGUGUGGGA AGACGGUGAUUAUUAUAGGAAACAACUAUCUCCACUUG AAGGUGGUGGCUGGCUGUUGCUAGUGGCUCAACUGUU GCCAUGACUGAGCAAUUACAGAUGGGCUUUGGUAUUAC AGUUCAAUAUGGUACAGACACCAAUAGUGUUUGCCCCA AGCUUGAAUUUGCUAAUGACACAAAAAUUGCCUCUCAA UUAGGCAAUUGCGUGGAAUAUUCCCUCUAUGGUGUUUC GGGCCGUGGUGUUUUUCAGAAUUGCACAGCUGUAGGUG UUCGACAGCAGCGCUUUGUUUAUGAUGCGUACCAGAAU UUAGUUGGCUAUUAUUCUGAUGAUGGCAACUACUACUG UCUGCGUGCUUGUGUUAGUGUUCCUGUUUCUGUCAUCU AUGAUAAAGAAACUAAAACCCACGCUACUCUAUUUGGU AGUGUUGCAUGUGAACACAUUUCUUCUACCAUGUCUCA AUACUCCCGUUCUACGCGAUCAAUGCUUAAACGGCGAGA UUCUACAUAUGGCCCCCUUCAGACACCUGUUGGUUGUGU CCUAGGACUUGUUAAUUCCUCUUUGUUCGUAGAGGACU GCAAGUUGCCUCUCGGUCAAUCUCUCUGUGCUCUUCCUG ACACACCUAGUACUCUCACACCUCGCAGUGUGCGCUCUG UGCCAGGUGAAAUGCGCUUGGCAUCCAUUGCUUUUAAU CAUCCCAUUCAGGUUGAUCAACUUAAUAGUAGUUAUUU UAAAUUAAGUAUACCCACUAAUUUUUCCUUUGGUGUGA CUCAGGAGUACAUUCAGACAACCAUUCAGAAAGUUACU GUUGAUUGUAAACAGUACGUUUGCAAUGGUUUCCAGAA GUGUGAGCAAUUACUGCGCGAGUAUGGCCAGUUUUGUU CCAAAAUAAACCAGGCUCUCCAUGGUGCCAAUUUACGCC AGGAUGAUUCUGUACGUAAUUUGUUUGCGAGCGUGAAA AGCUCUCAAUCAUCUCCUAUCAUACCAGGUUUUGGAGGU GACUUUAAUUUGACACUUCUAGAACCUGUUUCUAUAUC UACUGGCAGUCGUAGUGCACGUAGUGCUAUUGAGGAUU UGCUAUUUGACAAAGUCACUAUAGCUGAUCCUGGUUAU AUGCAAGGUUACGAUGAUUGUAUGCAGCAAGGUCCAGC AUCAGCUCGUGAUCUUAUUUGUGCUCAAUAUGUGGCUG GUUAUAAAGUAUUACCUCCUCUUAUGGAUGUUAAUAUG GAAGCCGCGUAUACUUCAUCUUUGCUUGGCAGCAUAGCA GGUGUUGGCUGGACUGCUGGCUUAUCCUCCUUUGCUGCU AUUCCAUUUGCACAGAGUAUYUUUUAUAGGUUAAACGG UGUUGGCAUUACUCAACAGGUUCUUUCAGAGAACCAAA AGCUUAUUGCCAAUAAGUUUAAUCAGGCUCUGGGAGCU AUGCAAACAGGCUUCACUACAACUAAUGAAGCUUUUCG GAAGGUUCAGGAUGCUGUGAACAACAAUGCACAGGCUC UAUCCAAAUUAGCUAGCGAGCUAUCUAAUACUUUUGGU GCUAUUUCCGCCUCUAUUGGAGACAUCAUACAACGUCUU GAUGUUCUCGAACAGGACGCCCAAAUAGACAGACUUAU UAAUGGCCGUUUGACAACACUAAAUGCUUUUGUUGCAC AGCAGCUUGUUCGUUCCGAAUCAGCUGCUCUUUCCGCUC AAUUGGCUAAAGAUAAAGUCAAUGAGUGUGUCAAGGCA CAAUCCAAGCGUUCUGGAUUUUGCGGUCAAGGCACACAU AUAGUGUCCUUUGUUGUAAAUGCCCCUAAUGGCCUUUA CUUUAUGCAUGUUGGUUAUUACCCUAGCAACCACAUUG AGGUUGUUUCUGCUUAUGGUCUUUGCGAUGCAGCUAAC CCUACUAAUUGUAUAGCCCCUGUUAAUGGCUACUUUAU UAAAACUAAUAACACUAGGAUUGUUGAUGAGGGUCAU AUACUGGCUCGUCCUUCUAUGCACCUGAGCCCAUCACCU CUCUUAAUACUAAGUAUGUUGCACCACAGGUGACAUACC AAAACAUUUCUACUAACCUCCCUCCUCCUCUUCUCGGCA AUUCCACCGGGAUUGACUUCCAAGAUGAGUUGGAUGAG UUUUUCAAAAAUGUUAGCACCAGUAUACCUAAUUUUGG UUCUCUAACACAGAUUAAUACUACAUUACUCGAUCUUAC CUACGAGAUGUUGUCUCUUCAACAAGUUGUUAAAGCCC UUAAUGAGUCUUACAUAGACCUUAAAGAGCUUGGCAAU UAUACUUAUUACAACAAAUGGCCGUGGUACAUUUGGCU UGGUUUCAUUGCUGGGCUUGUUGCCUUAGCUCUAUGCG UCUUCUUCAUACUGUGCUGCACUGGUUGUGGCACAAACU GUAUGGGAAAACUUAAGUGUAAUCGUUGUUGUGAUAGA UACGAGGAAUACGACCUCGAGCCGCAUAAGGUUCAUGU UCACUAA

| Strain                     | Nucleic Acid Sequence  | SEQ ID<br>NO: |
|----------------------------|--|---------------|
| MERS S FL                  | AUGAUACACUCAGUGUUUCUACUGAUGUUCUUGUUAAC   | 66            |
| SPIKE                      | ACCUACAGAAAGUUACGUUGAUGUAGGGCCAGAUUCUG<br>UUAAGUCUGCUUGUAUUGAGGUUGAUAUACAACAGACU   |               |
| 2cEMC/2012<br>(XBaI change | UUCUUUGAUAAAACUUGGCCUAGGCCAAUUGAUGUUUC   |               |
| (U to G))                  | UAAGGCUGACGGUAUUAUAUACCCUCAAGGCCGUACAU   |               |
| (nucleotide)               | AUUCUAACAUAACUAUCACUUAUCAAGGUCUUUUUCCCU  |               |
| ,                          | AUCAGGGAGACCAUGGUGAUAUGUAUGUUUACUCUGCA   |               |
|                            | GGACAUGCUACAGGCACAACUCCACAAAAGUUGUUUGU   |               |
|                            | AGCUAACUAUUCUCAGGACGUCAAACAGUUUGCUAAUG   |               |
|                            | GGUUUGUCGUCCGUAUAGGAGCAGCUGCCAAUUCCACUG  |               |
|                            | GCACUGUUAUUAUUAGCCCAUCUACCAGCGCUACUAUAC  |               |
|                            | GAAAAAUUUACCCUGCUUUUAUGCUGGGUUCUUCAGUU<br>GGUAAUUUCUCAGAUGGUAAAAUGGGCCGCUUCUUCAA   |               |
|                            | UCAUACUCUAGUUCUUUUGCCCGAUGGAUGUGGCACUU   |               |
|                            | UACUUAGAGCUUUUUAUUGUAUUCUGGAGCCUCGCUCU   |               |
|                            | GGAAAUCAUUGUCCUGCUGGCAAUUCCUAUACUUCUUU   |               |
|                            | UGCCACUUAUCACACUCCUGCAACAGAUUGUUCUGAUGG  |               |
|                            | CAAUUACAAUCGUAAUGCCAGUCUGAACUCUUUUAAGG   |               |
|                            | AGUAUUUUAAUUUACGUAACUGCACCUUUAUGUACACU   |               |
|                            | UAUAACAUUACCGAAGAUGAGAUUUUAGAGUGGUUUGG   |               |
|                            | CAUUACACAAACUGCUCAAGGUGUUCACCUCUUCUCAUC<br>UCGGUAUGUUGAUUUGUACGGCGCAAUAUGUUUCAAU   |               |
|                            | UUGCCACCUUGCCUGUUUAUGAUACUAUUAAGUAUUAU   |               |
|                            | UCUAUCAUUCCUCACAGUAUUCGUUCUAUCCAAAGUGAU  |               |
|                            | AGAAAAGCUUGGGCUGCCUUCUACGUAUAUAAACUUCA   |               |
|                            | ACCGUUAACUUUCCUGUUGGAUUUUUCUGUUGAUGGUU   |               |
|                            | AUAUACGCAGAGCUAUAGACUGUGGUUUUAAUGAUUUG   |               |
|                            | UCACAACUCCACUGCUCAUAUGAAUCCUUCGAUGUUGAA  |               |
|                            | UCUGGAGUUUAUUCAGUUUCGUCUUUCGAAGCAAAACC   |               |
|                            | UUCUGGCUCAGUUGUGGAACAGGCUGAAGGUGUUGAAU GUGAUUUUUCACCUCUCUGGCACACCUCCUCAGG          |               |
|                            | UUUAUAAUUUCAAGCGUUUGGUUUUUACCAAUUGCAAU   |               |
|                            | UAUAAUCUUACCAAAUUGCUUUCACUUUUUUCUGUGAA   |               |
|                            | UGAUUUUACUUGUAGUCAAAUAUCUCCAGCAGCAAUUG   |               |
|                            | CUAGCAACUGUUAUUCUUCACUGAUUUUGGAUUACUUU   |               |
|                            | UCAUACCCACUUAGUAUGAAAUCCGAUCUCAGUGUUAG   |               |
|                            | UUCUGCUGGUCCAAUAUCCCAGUUUAAUUAUAAACAGU   |               |
|                            | CCUUUUCUAAUCCCACAUGUUUGAUUUUAGCGACUGUUC CUCAUAACCUUACUACUAUUACUAAGCCUCUUAAGUACA    |               |
|                            | GCUAUAUUAACAAGUGCUCUCGUCUUCUUUCUGAUGAU   |               |
|                            | CGUACUGAAGUACCUCAGUUAGUGAACGCUAAUCAAUA   |               |
|                            | CUCACCCUGUGUAUCCAUUGUCCCAUCCACUGUGUGGGA  |               |
|                            | AGACGGUGAUUAUUAUAGGAAACAACUAUCUCCACUUG   |               |
|                            | AAGGUGGUGGCUUGUUGCUAGUGGCUCAACUGUU   |               |
|                            | GCCAUGACUGAGCAAUUACAGAUGGGCUUUGGUAUUAC   |               |
|                            | AGUUCAAUAUGGUACAGACACCAAUAGUGUUUGCCCCA AGCUUGAAUUUGCUAAUGACACAAAAAUUGCCUCUCAA      |               |
|                            | UUAGGCAAUUGCGUGGAAUAUUCCCUCUAUGGUGUUUC   |               |
|                            | GGGCCGUGGUUUUUCAGAAUUGCACAGCUGUAGGUG   |               |
|                            | UUCGACAGCAGCGCUUUGUUUAUGAUGCGUACCAGAAU   |               |
|                            | UUAGUUGGCUAUUAUUCUGAUGAUGGCAACUACUACUG   |               |
|                            | UUUGCGUGCUUGUGUUAGUGUUCCUGUUUCUGUCAUCU   |               |
|                            | AUGAUAAAGAAACUAAAACCCACGCUACUCUAUUUGGU   |               |
|                            | AGUGUUGCAUGUGAACACAUUUCUUCUACCAUGUCUCA   |               |
|                            | AUACUCCCGUUCUACGCGAUCAAUGCUUAAACGGCGAGA<br>UUCUACAUAUGGCCCCCUUCAGACACCUGUUGGUUGUGU |               |
|                            | CCUAGGACUUGUUAAUUCCUCUUUGUUCGUAGAGGACU   |               |
|                            | GCAAGUUGCCUCUUGGUCAAUCUCUGUGUGCUCUUCCUG  |               |
|                            | ACACACCUAGUACUCUCACACCUCGCAGUGUGCGCUCUG  |               |
|                            | UUCCAGGUGAAAUGCGCUUGGCAUCCAUUGCUUUUAAU   |               |
|                            | CAUCCUAUUCAGGUUGAUCAACUUAAUAGUAGUUAUUU   |               |
|                            | UAAAUUAAGUAUACCCACUAAUUUUUCCUUUGGUGUGA   |               |
|                            | CUCAGGAGUACAUUCAGACAACCAUUCAGAAAGUUACU   |               |
|                            | GUUGAUUGUAAACAGUACGUUUGCAAUGGUUUCCAGAA<br>GUGUGAGCAAUUACUGCGCGAGUAUGGCCAGUUUUGUU   |               |
|                            | CCAAAAUAAACCAGGCUCUCCAUGGUGCCAAUUUACGCC  |               |
|                            | AGGAUGAUUCUGUACGUAAUUUGUUUGCGAGCGUGAAA   |               |
|                            | AGCUCUCAAUCAUCUCCUAUCAUACCAGGUUUUGGAGGU  |               |
|                            | GACUUUAAUUUGACACUUCUGGAACCUGUUUCUAUAUC   |               |
|                            | UACUGGCAGUCGUAGUGCACGUAGUGCUAUUGAGGAUU   |               |
|                            | UGCUAUUUGACAAAGUCACUAUAGCUGAUCCUGGUUAU   |               |
|                            | AUGCAAGGUUACGAUGAUUGCAUGCAGCAAGGUCCAGC   |               |
|                            | AUCAGCUCGUGAUCUUAUUUGUGCUCAAUAUGUGGCUG GUUACAAAGUAUUACCUCCUCUUAUGGAUGUUAAUAUG      |               |
|                            | GAAGCCGCGUAUACUUCAUCUUUGCUUGGCAGCAUAGCA  |               |
|                            | GGUGUUGGCUGGACUGCUUAUCCUCCUUUGCUGCU  |               |
|                            | AUUCCAUUUGCACAGAGUAUCUUUUAUAGGUUAAACGG   |               |

SEQ ID Strain Nucleic Acid Sequence NO:

UGUUGGCAUUACUCAACAGGUUCUUUCAGAGAACCAAA AGCUUAUUGCCAAUAAGUUUAAUCAGGCUCUGGGAGCU AUGCAAACAGGCUUCACUACAACUAAUGAAGCUUUUCA GAAGGUUCAGGAUGCUGUGAACAACAAUGCACAGGCUC UAUCCAAAUUAGCUAGCGAGCUAUCUAAUACUUUUGGU GCUAUUUCCGCCUCUAUUGGAGACAUCAUACAACGUCUU GAUGUUCUCGAACAGGACGCCCAAAUAGACAGACUUAU UAAUGGCCGUUUGACAACACUAAAUGCUUUUGUUGCAC AGCAGCUUGUUCGUUCCGAAUCAGCUGCUCUUUCCGCUC AAUUGGCUAAAGAUAAAGUCAAUGAGUGUGUCAAGGCA CAAUCCAAGCGUUCUGGAUUUUGCGGUCAAGGCACACAU AUAGUGUCCUUUGUUGUAAAUGCCCCUAAUGGCCUUUA CUUCAUGCAUGUUGGUUAUUACCCUAGCAACCACAUUGA GGUUGUUUCUGCUUAUGGUCUUUGCGAUGCAGCUAACC CUACUAAUUGUAUAGCCCCUGUUAAUGGCUACUUUAUU AAAACUAAUAACACUAGGAUUGUUGAUGAGUGGUCAUA UACUGGCUCGUCCUUCUAUGCACCUGAGCCCAUUACCUC CCUUAAUACUAAGUAUGUUGCACCACAGGUGACAUACCA AAACAUUUCUACUAACCUCCCUCCUCCUCUUCUCGGCAA UUCCACCGGGAUUGACUUCCAAGAUGAGUUGGAUGAGU UUUUCAAAAAUGUUAGCACCAGUAUACCUAAUUUUGGU UCCCUAACACAGAUUAAUACUACAUUACUCGAUCUUACC UACGAGAUGUUGUCUCUUCAACAAGUUGUUAAAGCCCU UAAUGAGUCUUACAUAGACCUUAAAGAGCUUGGCAAUU AUACUUAUUACAACAAAUGGCCGUGGUACAUUUGGCUU GGIIIIICAIIIIGCIIGGGCIIIIGIIIIGCCIIIIAGCIICIIAIIGCGII CUUCUUCAUACUGUGCUGCACUGGUUGUGGCACAAACUG UAUGGGAAAACUUAAGUGUAAUCGUUGUUGUGAUAGAU ACGAGGAAUACGACCUCGAGCCGCAUAAGGUUCAUGUUC ACUAA

AUGAUCCACUCCGUGUUCCUCCUCAUGUUCCUGUUGACC

Novel\_MERS\_S2\_subunit\_trimeric vaccine (nucleotide)

 $\tt CCCACUGAGUCAGACUGCAAGCUCCCGCUGGGACAGUCC$ CUGUGUGCGCUGACACUCCUAGCACUCUGACCCCA CGCUCCGUGCGGUCGGUGCCUGGCGAAAUGCGGCUGGCC UCCAUCGCCUUCAAUCACCCAAUCCAAGUGGAUCAGCUG AAUAGCUCGUAUUUCAAGCUGUCCAUCCCCACGAACUUC UCGUUCGGGGUCACCCAGGAGUACAUCCAGACCACAAUU CAGAAGGUCACCGUCGAUUGCAAGCAAUACGUGUGCAAC GGCUUCCAGAAGUGCGAGCAGCUGCUGAGAGAAUACGG GCAGUUUUGCAGCAAGAUCAACCAGGCGCUGCAUGGAGC UAACUUGCGCCAGGACGACUCCGUGCGCAACCUCUUUGC CUCUGUGAAGUCAUCCCAGUCCUCCCCAAUCAUCCCGGG AUUCGGAGGGGACUUCAACCUGACCCUCCUGGAGCCCGU GUCGAUCAGCACCGGUAGCAGAUCGGCGCGCUCAGCCAU UGAAGAUCUUCUGUUCGACAAGGUCACCAUCGCCGAUCC GGGCUACAUGCAGGAUACGACGACUGUAUGCAGCAGG GACCAGCCUCGCGAGGGACCUCAUCUGCGCGCAAUACG UGGCCGGGUACAAAGUGCUGCCUCCUCUGAUGGAUGUG AACAUGGAGGCCGCUUAUACUUCGUCCCUGCUCGGCUCU AUCGCCGGCGUGGGGUGGACCGCCGGCCUGUCCUUC GCCGCUAUCCCCUUUGCACAAUCCAUUUUCUACCGGCUC AACGGCGUGGGCAUUACUCAACAAGUCCUGUCGGAGAAC CAGAAGUUGAUCGCAAACAAGUUCAAUCAGGCCCUGGG GGCCAUGCAGACUGGAUUCACUACGACUAACGAAGCGUU CCAGAAGGUCCAGGACGCUGUGAACAACAACGCCCAGGC GCUCUCAAAGCUGGCCUCCGAACUCAGCAACACCUUCGG AGCCAUCAGCGCAUCGAUCGGUGACAUAAUUCAGCGGCU GGACGUGCUGGAGCAGGACGCCCAGAUCGACCGCCUCAU CAACGGACGGCUGACCACCUUGAAUGCCUUCGUGGCACA ACAGCUGGUCCGGAGCGAAUCAGCGGCACUUUCCGCCCA ACUCGCCAAGGACAAAGUCAACGAAUGCGUGAAGGCCCA GUCCAAGAGGUCCGGUUUCUGCGGUCAAGGAACCCAUAU UGUGUCCUUCGUCGUGAACGCCCCAACGGUCUGUACUU UAUGCACGUCGGCUACUACCCGAGCAAUCAUAUCGAAGU GGUGUCCGCCUACGGCCUGUGCGAUGCCGCUAACCCCAC UAACUGUAUUGCCCCUGUGAACGGAUAUUUUAUUAAGA CCAACAACACCCGCAUUGUGGACGAAUGGUCAUACACCG GUUCGUCCUUCUACGCGCCCGAGCCCAUCACUUCACUGA ACACCAAAUACGUGGCUCCGCAAGUGACCUACCAGAACA UCUCCACCAAUUUGCCGCCGCCGCUGCUCGGAAACAGCA CCGGAAUUGAUUUCCAAGAUGAACUGGACGAAUUCUUC AAGAACGUGUCCACUUCCAUUCCCAACUUCGGAAGCCUG ACACAGAUCAACACCACCCUUCUCGACCUGACCUACGAG AUGCUGAGCCUUCAACAAGUGGUCAAGGCCCUGAACGAG AGCUACAUCGACCUGAAGGAGCUGGGCAACUAUACCUAC UACAACAAGUGGCCGGACAAGAUUGAGGAGAUUCUGUC

| Strain                        | Nucleic Acid Sequence  | SEQ ID<br>NO: |
|-------------------------------|--|---------------|
|                               | GAAAAUCUACCACAUUGAAAACGAGAUCGCCAGAAUCA<br>AGAAGCUUAUCGGCGAAGCC                     |               |
| MERS_S0_Full-<br>length Spike | AUGGAAACCCCUGCCCAGCUGCUGUUCCUGCUGCUGCUG<br>UGGCUGCCUGAUACCACCGGCAGCUAUGUGGACGUGGGC | 68            |
| protein                       | CCCGAUAGCGUGAAGUCCGCCUGUAUCGAAGUGGACAUC  |               |
| (nucleotide,                  | CAGCAGACCUUUUUCGACAAGACCUGGCCCAGACCCAUC  |               |
| codon                         | GACGUGUCCAAGGCCGACGGCAUCAUCUAUCCACAAGGC<br>CGGACCUACAGCAACAUCACCAUUACCUACCAGGGCCUG |               |
| optimized)                    | UUCCCAUAUCAGCGACCACGGCGAUAUGUACCUGGGCCUG UUCCCAUAUCAAGGCGACCACGGCGAUAUGUACGUGUAC   |               |
|                               | UCUGCCGGCCACGCCACCGGCACCCCCAGAAACUG  |               |
|                               | UUCGUGGCCAACUACAGCCAGGACGUGAAGCAGUUCGCC  |               |
|                               | AACGGCUUCGUCGUGCGAUUGGCGCCGCUGCCAAUAGC   |               |
|                               | ACCGGCACAGUGAUCAUCAGCCCCAGCACCAGCGCCACC  |               |
|                               | AUCCGGAAGAUCUACCCCGCCUUCAUGCUGGGCAGCUCC<br>GUGGGCAAUUUCAGCGACGGCAAGAUGGGCCGGUUCUU  |               |
|                               | CAACCACACCUGGUGCUGCUGCCGAUGGCUGUGGCAC  |               |
|                               | ACUGCUGAGAGCCUUCUACUGCAUCCUGGAACCCAGAAG  |               |
|                               | CGGCAACCACUGCCCUGCCGGCAAUAGCUACACCAGCUU  |               |
|                               | CGCCACCUACCACACCCGCCACCGAUUGCUCCGACGG<br>CAACUACAACCGGAACGCCAGCCUGAACAGCUUCAAAGA   |               |
|                               | GUACUUCAACCUGCGGAACUGCACCUUCAUGUACACCUA  |               |
|                               | CAAUAUCACCGAGGACGAGAUCCUGGAAUGGUUCGGCA   |               |
|                               | UCACCCAGACCGCCCAGGGCGUGCACCUGUUCAGCAGCA  |               |
|                               | GAUACGUGGACCUGUACGGCGCAACAUGUUCCAGUUU  |               |
|                               | GCCACCCUGCCCGUGUACGACACCAUCAAGUACUACAGC AUCAUCCCCCACAGCAUCCGGUCCAUCCAGAGCGACAGA    |               |
|                               | AAAGCCUGGGCCGCCUUCUACGUGUACAAGCUGCAGCCC  |               |
|                               | CUGACCUUCCUGCUGGACUUCAGCGUGGACGGCUACAUC  |               |
|                               | AGACGGGCCAUCGACUGCGGCUUCAACGACCUGAGCCAG  |               |
|                               | CUGCACUGCUCCUACGAGAGCUUCGACGUGGAAAGCGGC  |               |
|                               | GUGUACAGCGUGUCCAGCUUCGAGGCCAAGCCUAGCGGC<br>AGCGUGGUGGAACAGGCUGAGGGCGUGGAAUGCGACUU  |               |
|                               | CAGCCCUCUGCUGAGCGCCCCCCCCCCCAGGUGUACAA   |               |
|                               | CUUCAAGCGGCUGGUGUUCACCAACUGCAAUUACAACCU  |               |
|                               | GACCAAGCUGCUGAGCCUGUUCUCCGUGAACGACUUCAC  |               |
|                               | CUGUAGCCAGAUCAGCCCUGCCGCCAUUGCCAGCAACUG<br>CUACAGCAGCCUGAUCCUGGACUACUUCAGCUACCCCCU |               |
|                               | GAGCAUGAAGUCCGAUCUGAGCGUGUCCUCCGCCGGACC  |               |
|                               | CAUCAGCCAGUUCAACUACAAGCAGAGCUUCAGCAACCC  |               |
|                               | UACCUGCCUGAUUCUGGCCACCGUGCCCCACAAUCUGAC  |               |
|                               | CACCAUCACCAAGCCCCUGAAGUACAGCUACAUCAACAA<br>GUGCAGCAGACUGCUGUCCGACGGACCGGAC         |               |
|                               | CCAGCUCGUGAACGCCAACCAGUACAGCCCCUGCGUGUC  |               |
|                               | CAUCGUGCCCAGCACCGUGUGGGAGGACGGCGACUACUA  |               |
|                               | CAGAAAGCAGCUGAGCCCCCUGGAAGGCGGCGGAUGGCU  |               |
|                               | GGUGGCUUCUGGAAGCACAGUGGCCAUGACCGAGCAGCU<br>GCAGAUGGGCUUUGGCAUCACCGUGCAGUACGGCACCGA |               |
|                               | CACCAACAGCGUGUGCCCCAAGCUGGAAUUCGCCAAUGA  |               |
|                               | CACCAAGAUCGCCAGCCAGCUGGGAAACUGCGUGGAAUA  |               |
|                               | CUCCCUGUAUGGCGUGUCCGGACGGGGCGUGUUCCAGAA  |               |
|                               | UUGCACAGCAGUGGGAGUGCGGCAGCAGAGAUUCGUGU<br>ACGAUGCCUACCAGAACCUCGUGGGCUACUACAGCGACG  |               |
|                               | ACGGCAAUUACUACUGCCUGCGGGCCUGUGUGUCCGUGC  |               |
|                               | CCGUGUCCGUGAUCUACGACAAAGAGACAAAGACCCACG  |               |
|                               | CCACACUGUUCGGCUCCGUGGCCUGCGAGCACAUCAGCU  |               |
|                               | CCACCAUGAGCCAGUACUCCCGCUCCACCCGGUCCAUGC UGAAGCGGAGAGAUAGCACCUACGGCCCCUGCAGACAC     |               |
|                               | CUGUGGGAUGUGUGCUGGGCCUCGUGAACAGCUCCCUGU  |               |
|                               | UUGUGGAAGAUUGCAAGCUGCCCCUGGGCCAGAGCCUGU  |               |
|                               | GUGCCCUGCCAGAUACCCCUAGCACCCUGACCCCUAGAA  |               |
|                               | GCGUGCGCUCUGUGCCCGGCGAAAUGCGGCUGGCCUCUA<br>UCGCCUUCAAUCACCCCAUCCAGGUGGACCAGCUGAACU |               |
|                               | CCAGCUACUACAGCUGAGCAUCCCACCAACUUCAGCU  |               |
|                               | UCGGCGUGACCCAGGAGUACAUCCAGACCACAAUCCAGA  |               |
|                               | AAGUGACCGUGGACUGCAAGCAGUACGUGUGCAACGGC   |               |
|                               | UUUCAGAAGUGCGAACAGCUGCUGCGCGAGUACGGCCAG  |               |
|                               | UUCUGCAGCAAGAUCAACCAGGCCCUGCACGGCGCCAAC<br>CUGAGACAGGAUGACAGCGUGCGGAACCUGUUCGCCAGC |               |
|                               | GUGAAAAGCAGCCAGUCCAGCCCAUCAUCCCUGGCUUC   |               |
|                               | GGCGGCGACUUUAACCUGACCCUGCUGGAACCUGUGUCC  |               |
|                               | AUCAGCACCGGCUCCAGAAGCGCCAGAUCCGCCAUCGAG  |               |
|                               | GACCUGCUGUUCGACAAAGUGACCAUUGCCGACCCCGGC<br>UACAUGCAGGGCUACGACGAUUGCAUGCAGCAGGGCCCA |               |
|                               | GCCAGCGCCAGGAUCUGAUCUGUGCCCAGUAUGUGGCC   |               |
|                               | GGCUACAAGGUGCUGCCCCCCUGAUGGACGUGAACAUG   |               |
|                               | GAAGCCGCCUACACCUCCAGCCUGCUGGGCUCUAUUGCU  |               |

| Strain | Nucleic Acid Sequence                   | SEQ ID<br>NO: |
|--------|---|---------------|
|        | GGCGUGGGAUGGACAGCCGGCCUGUCUAGCUUUGCCGCC |               |
|        | AUCCCUUUCGCCCAGAGCAUCUUCUACCGGCUGAACGGC |               |
|        | GUGGGCAUCACACAACAGGUGCUGAGCGAGAACCAGAA  |               |
|        | GCUGAUCGCCAACAAGUUUAACCAGGCACUGGGCGCCAU |               |
|        | GCAGACCGGCUUCACCACCAACGAGGCCUUCAGAAA    |               |
|        | GGUGCAGGACGCCGUGAACAACAACGCCCAGGCUCUGAG |               |
|        | CAAGCUGGCCUCCGAGCUGAGCAAUACCUUCGGCGCCAU |               |
|        | CAGCGCCUCCAUCGGCGACAUCAUCCAGCGGCUGGACGU |               |
|        | GCUGGAACAGGACGCCCAGAUCGACCGGCUGAUCAACGG |               |
|        | CAGACUGACCACCCUGAACGCCUUCGUGGCACAGCAGCU |               |
|        | CGUGCGGAGCGAAUCUGCCGCUCUGUCUGCUCAGCUGGC |               |
|        | CAAGGACAAAGUGAACGAGUGCGUGAAGGCCCAGUCCA  |               |
|        | AGCGGAGCGGCUUUUGUGGCCAGGGCACCCACAUCGUGU |               |
|        | CCUUCGUCGUGAAUGCCCCCAACGGCCUGUACUUUAUGC |               |
|        | ACGUGGGCUAUUACCCCAGCAACCACAUCGAGGUGGUGU |               |
|        | CCGCCUAUGGCCUGUGCGACGCCCAAUCCUACCAACU   |               |
|        | GUAUCGCCCCGUGAACGGCUACUUCAUCAAGACCAACA  |               |
|        | ACACCCGGAUCGUGGACGAGUGGUCCUACACAGGCAGCA |               |
|        | GCUUCUACGCCCCGAGCCCAUCACCUCCCUGAACACCA  |               |
|        | AAUACGUGGCCCCCAAGUGACAUACCAGAACAUCUCCA  |               |
|        | CCAACCUGCCCCUCCACUGCUGGGAAAUUCCACCGGCA  |               |
|        | UCGACUUCCAGGACGAGCUGGACGAGUUCUUCAAGAACG |               |
|        | UGUCCACCUCCAUCCCCAACUUCGGCAGCCUGACCCAGA |               |
|        | UCAACACCACUCUGCUGGACCUGACCUACGAGAUGCUGU |               |
|        | CCCUGCAACAGGUCGUGAAAGCCCUGAACGAGAGCUACA |               |
|        | UCGACCUGAAAGAGCUGGGGAACUACACCUACUACAACA |               |
|        | AGUGGCCUUGGUACAUUUGGCUGGGCUUUAUCGCCGGCC |               |
|        | UGGUGGCCCUGGCCCUGUGCGUGUUCUUCAUCCUGUGCU |               |
|        | GCACCGGCUGCGGCACCAAUUGCAUGGGCAAGCUGAAAU |               |
|        | GCAACCGGUGCUGCGACAGAUACGAGGAAUACGACCUGG |               |
|        | AACCUCACAAAGUGCAUGUGCAC                 |               |
|        | JAJDUDUAJDUDAAAJAJUJJAA                 |               |

# TABLE 11

| TADLE II  |  |               |  |  |
|---|--|---------------|--|--|
| В   | etacoronavirus Amino Acid Sequences  |               |  |  |
| Strain  | Amino Acid Sequence  | SEQ ID<br>NO: |  |  |
| gb KJ156934.1 : 21405-25466 Middle East respiratory syndrome coronavirus isolate Riyadh_14_2013, spike protein (amino acid) | MIHSVFLLMFLLTPTESYVDVGPDSVKSACIEVDIQQTFFDK TWPRPIDVSKADGIIYPQGRTYSNITITYQGLFPYQGDHGDM YVYSAGHATGTTPQKLFVANYSQDVKQFANGFVVRIGAAANS TGTVIISPSTSATIRKIYPAFMLGSSVGNFSDGKMGRFFNHT LVLLPDGCGTLLRAFYCILEPRSGNHCPAGNSYTSFATTHTP ATDCSDGNYNRNASLNSFKEYFNLRNCTFMYTYNITEDEILE WFGITQTAQGVHLFSSRYVDLYGGNMFQFATLPVYDTIKYYS IIPHSIRSIQSDRKAWAAAFYVYKLQPLTFLLDFSVDGYIRRA IDCGFNDLSQLHCSYESFDVESGVYSVSSFEAKPSGSVVEQA EGVECDFSPLLSGTPPQVYNFKRLVFTNCNYNLTKLLSLFSV NDFtCSQISPAAIASNCYSSLILDYFSYPLSMKSDLSVSSAG PISQFNYKQSFSNPTCLILATVPHNLTTITKPLKYSYINKCS RLLSDDRTEVPQLVNANQYSPCVSIVPSTVWEDGDYYRKQLS PLEGGGWLVASGSTVANTEQLQMGFGITVQYGTDTNSVCPKL EFANDTKIASQLGNCVEYSLYGVSGRGVFQNCTAVGVRQQRF VYDAYQNLVGYYSDDGMYYCLRACVSVPVSVIYDKETKTHAT LFGSVACEHISSTMSQYSRSTRSMLKRRDSTYGPLQTPVGCV LGLVNSSLFVEDCKLPLGQSLCALPDTPSTLTPRSVRSVPGE MRLASIAFNHPIQVDQLNSSYFKLSIPTNFSFGVTQEYIQTT IQKVTVDCKQYVCNGFQKCEQLLREYGQFCSKINQALHGANL RQDDSVRNLFASVKSSQSPIIPGFGGDFNLTLLEPVSISTG SRSARSAIEDLLFDKVTIADPGYMQGYDDCMQQGPASARDLI CAQYVAGYKULPPLMDVNMEAAYTSSLLGSIAGVGWTAGLSS FAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFNQALGAM QTGFTTTNEAFTKVQDAVNNNAQALSKLASELSNTFGAISAS IGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQLVRSESA ALSAQLAKDKVNECVKAQSKRSGFCGGGTHIVSFVVNAPNGL YFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNGYFIKN NTRIVDEWSYTGSSFYAPEPITSLNTKYVAPQVTYQNISTNL PPPLLGNSTGIDFQDELDEFFRNVSTSIPNFGSLTQINTTLL DLTYEMLSLQVVKALNESYIDLKELGNYTYYNKWPWYIWLG FIAGLVALACVFFILCCTGCGTNCMGKLKCNRCCDRYEEYD LEPHKVHVH | 24            |  |  |

| TABLE 11-continued  Betacoronavirus Amino Acid Sequences       |   |    |  |  |
|--|---|----|--|--|
|  |   |    |  |  |
| MERS S FL SPIKE 2cEMC/2012 (XBaI change (T to G)) (amino acid) | MIHSVFLLMFLLTPTESYVDVGPDSVKSACIEVDIQQTFFDK TWPRPIDVSKADGIIYPQGRTYSNITITYQGLFPYQGDHGDM YVYSAGHATGTTPQKLFVANYSQDVKQFANGFVVRIGAAANS TGTVIISPSTSATIRKIYPAFMLGSSVGMFSDGKMGRFFNHT LVLLPDGCGTLLRAFYCILEPRSGNHCPAGNSYTSFATYHTP ATDCSDGNYNRNASLNSFKEYFNLRNCTFMYTYNITEDEILE WFGITQTAQGVHLFSSRYVDLYGGMMFQFATLPVYDTIKYYS IIPHSIRSIQSDRKAWAAFYVYKLQPLTFLLDFSVDGYIRRA IDCGFNDLSQLHCSYESFDVESGVYSVSSFEAKPSGSVVEQA EGVECDFSPLLSGTPPQVYNFKRLVFTNCNYNLTKLLSLFSV NDFTCSQISPAAIASNCYSSLILDYFSYPLSMKSDLSVSSAG PISQFNYKQSFSNPTCLILATVPHNLTTITKPLKYSYINKCS RLLSDDRTEVPQLVNANQYSPCVSIVPSTVWEDGDYYRKQLS PLEGGGWLVASGSTVAMTEQLQMGFGITVQYGTDTNSVCPKL EFANDTKIASQLGNCVEYSLYGYSGRGVFQNCTAYGVRQQFP VYDAYQNLVGYYSDDGNYYCLRACVSVPVSVIYDKETKTHAT LFGSVACEHISSTMSQYSRSTRSMLKRRDSTYGPLQTPVGCV LGLVNSSLFVEDCKLPLGQSLCALPDTPSTLTPRSVRSVPGE MRLASIAFNHPIQVDQLNSSYFKLSIPTNFSFGVTQEYIQTT IQKVTVDCKQVVCNGFGKCEQLLREYGQFCSKINQALHGANL RQDDSVRNLFASVKSSQSSPIIPGFGGDFNLTLLEPVSISTG SRSARSAIEDLLFDKVTIADPGTMQGYDDCMQQGPASARDLI CAQYVAGYKVLPPLMDVNMEAAYTSSLLGSIAGVGWTAGLSS FAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFNQALGAM QTGFTTTNEAFQKVQDAVNNNAQALSKLASELSNTFGAISAS IGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQLVRSESA ALSAQLAKDKVNECVKAQSKRSGFCQGTHIVSFVVNAPNGL YFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNGYFIKTN NTRIVDEWSYTGSSFYAPEPITSLNTKYVAPQVTYQNISTNL PPPLLGNSTGIDFQDELDEFFKNVSTSIPNFGSLTQINTTLL PPPLLGNSTGIDFQDELDEFFKNVSTSIPNFGSLTQINTTLL DLTYYEMLSLQQVVKALMESYIDLKELGNTYYNKFWFYINLG FIAGLVALALCVFFILCCTGCGTNCMGKLKCNRCCDRYEEYD LEPHKVHVH | 25 |  |  |
| Novel_MERS_S2_sub-<br>unit_trimeric<br>vaccine (amino<br>acid) | MIHSVFLLMFLLTPTESDCKLPLGQSLCALPDTPSTLTPRSV RSVPGEMRLASIAFNHPIQVDQLNSSYFKLSIPTNFSFGVTQ EYIQTTIQKVTVDCKQYVCNGFQKCEQLLREYGQFCSKINQA LHGANLRQDDSVRNLFASVKSSQSSPIIPGFGGDFNLTLLEP VSISTGSRSARSAIEDLLFDKVTIADPGYMQGYDDCMQQGPA SARDLICAQYVAGYKVLPPLMDVNMEAAYTSSLLGSIAGVGW TAGLSSFAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFN QALGAMQTGFTTTNEAFQKVQDAVNNNAQALSKLASELSNTF GAISASIGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQL VRSESAALSAQLAKDKVNECVKAQSKRSGFCGQGTHIVSFVV NAPNGLYFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNG YFIKTNNTRIVDEWSYTGSSFYAPEPITSLNTKYVAPQVTYQ NISTNLPPPLLGNSTGIDFQDELDEFFRNVSTSIPNFGSLTQ INTTLLDLTYEMLSLQQVVKALNESYIDLKELGNYTYYNKWP DKIEETLSKIYHIENEIARIKKLIGEA   | 26 |  |  |
| Isolate A1-<br>Hasa_1_2013<br>(NCBI accession<br>#AGN70962)    | MIHSVFLLMFLLTPTESYVDVGPDSVKSACIEVDIQQTFFDK TWPRPIDVSKADGIIYPQGTFYSNITITTYQGLFFYQGDHGDM YVYSAGHATGTTPQKLFVANYSQDVKQFANGFVVRIGAAANS TGTVIISPSTSATIRKIYPAFMLGSSVGMFSDGKMGRFFNHT LVLLPDGCGTLLRAFYCILEPRSGNHCPAGNSYTSFATYHTP ATDCSDGNYNRNASLNSFKEYFMLRNCTFMYTYNITEDEILE WFGITQTAQGVHLFSSRYVDLYGGMMFQFATLPVYDTIKYYS IIPHSIRSIQSDRKAWAAFYVYKLQPLTFLLDFSVDGYIRRA IDCGFNDLSQLHCSYESFDVESGVYSVSSFEAKPSGSVVEQA EGVECDFSPLLSGTPPQVYNFKRLVFTNCNYNLTKLLSLFSV NDFTCSQISPAAIASNCYSSLILDYFSYPLSMKSDLSVSSAG PISQFNYKQSFSNPTCLILATVPHNLTTITKPLKYSYINKCS RLLSDDRTEVPQLVNANQYSPCVSIVPSTVWEDGDYYRKQLS PLEGGGWLVASGSTVAMTEQLQMGFGITVQYGTDTNSVCPKL EFANDTKIASQLGNCVEYSLYGVSGRGVFQNCTAVGVRQQFF VYDAYQNLVGYYSDDGNYYCLRACVSVPVSVIYDKETKTHAT LFGSVACEHISSTMSQYSRSTRSMLKRRDSTYGPLQTPVGCV LGLVNSSLFVEDCKLPLGQSLCALPDTPSTLTPRSVRSVPGE MRLASIAFNHPIQVDQLNSSYFKLSIPTNFSFGVTQEYIQTT IQKVTVDCKQYVCNGFQKCEQLLREYGGFCSKINQALHGANL RQDDSVRNLFASVKSSQSSPIIPGFGGDFNLTLLEPVSISTG SRSARSAIEDLLFDKVTIADPGYMQGYDDCMQQGPASARDLI  | 27 |  |  |

CAQYVAGYKVLPPLMDVNMEAAYTSSLLGSIAGVGWTAGLSS FAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFNQALGAM

|  | ZADIE 11 gentinued   |       |
|--|--|-------|
|  | TABLE 11-continued  Betacoronavirus Amino Acid Sequences   |       |
| Strain   | Amino Acid Sequence  | SEQ I |
|  | QTGFTTTNEAFRKVQDAVNNNAQALSKLASELSNTFGAISAS IGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQLVRSESA ALSAQLAKDKVNECVKAQSKRSGFCGQGTHIVSFVVNAPNGL YFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNGYFIKTN NTRIVDEWSYTGSSFYAPEPITSLNTKYVAPHVTYQNISTNL PPPLLGNSTGIDFQDELDEFFKNVSTSIPNFGSLTQINTTLL DLTYEMLSLQQVVKALNESYIDLKELGNYTYYNKWFWYIWLG FIAGLVALALCVFFILCCTGCGTNCMGKLKCNRCCDRYEEYD LEPHKVHVH  |       |
| Middle East<br>respiratory<br>syndrome<br>coronavirus S<br>protein<br>UniProtKB-<br>R9UQ53                             | MIHSVFLLMFLLTPTESYVDVGPDSVKSACIEVDIQQTFFDK TWPRPIDVSKADGIIYPQGRTYSNITITYQGLFPYQGDHGDM YVYSAGHATGTTPQKLFVANYSQDVKQFANGFVVRIGAAANS TGTVIISPSTSATIRKIYPAFMLGSSVGNFSDGKMGRFFNHT LVLLPDGCGTLLRAFYCILEPRSGNHCPAGNSYTSFATYHTP ATDCSDGNYNRNASLNSFKEYFNLRNCTFMYTYNITEDEILE WFGITQTAQGVHLFSSRYVDLYGGNMFQFATLPVYDTIKYYS IIPHSIRSIQSDRKAWAAFYVYKLQPLTFLLDFSVDGYIRRA IDCGFNDLSQLHCSYESFDVESGVYSVSSFEAKPSGSVVEQA EGVECDFSPLLSGTPPQVYNFKRLVFTNCNYNLTKLLSLFSV NDFTCSQISPAAIASNCYSSLILDYFSYPLSMKSDLSVSSAG PISQFNYKQSFSNPTCLILATVPHNLTTITKPLKYSYINKCS RLLSDDRTEVPQLVNANQYSPCVSIVYSTVWEDGDYYRKQLS PLEGGGWLVASGSTVAMTEQLQMGFGITVQYGTDTNSVCPKL EFANDTKIASQLGNCVEYSLYGVSGRGVFQNCTAVGVRQQRF VYDAYQNLVGYYSDDGNYYCLRACVSVPVSVIYDKETKTHAT LFGSVACEHISSTMSQYSRSTRSMLKRRDSTYGPLQTPVGCV LGLVNSSLFVEDCKLPLQGSLCALPDTPSTLTPRSVRSVPGE MRLASIAFNHPIQVDQLNSSYFKLSIPTNFSFGVTQEYIQTT IQKVTVDCKQYVCNGFQKCEQLLREYGQFCSKINQALHGANL RQDDSVRNLFASVKSSQSSPIPGFGGDFNLTLLEPVSISTG SRSARSAIEDLLFDKVTIADPGYMQGYDDCMQQGPASARDLI CAQYVAGYKVLPPLMDVNMEAAYTSSLLGSIAGVGWTAGLSS FAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFNQALGAM QTGFTTTNEAFRKVQDAVNNNAQALSKLASELSNTFGAISAS IGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQLVRSESA ALSAQLAKDKVNECVKAQSKRSGFCGQGTHIVSFVVNAPNGL YFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNGYFIKTN NTRIVDEWSYTGSSFYAPEBITSLNTKYVAPHVTYQNISTNL PPPLLGNSTGIDFQDELDEFFKNVSTSIPNFGSLTQINTTLL DLTYEMLSLQQVVKALNESYIDLKELGNYTYYNKØPWYIWLG FIAGLVALALCVFFILCCTGCGTNCMGKLKCNRCCDRYEEYD LEPHKVHVH | 28    |
| Human SARS coronavirus (SARS-CoV) (Severe acute respiratory syndrome coronavirus) Spike glycoprotein UniProtKB- P59594 | MFIFLLFLTLTSGSDLDRCTTFDDVQAPNYTQHTSSMRGVYY PDEIFRSDTLYLTQDLFLPFYSNVTGFHTINHTFGNPVIPFK DGIYFAATEKSNVVRGWVFGSTMNNKSQSVIIINNSTNVVIR ACNFELCDNPFFAVSKPMGTQTHTMIFDNAFNCTFEYISDAF SLDVSEKSGNFKHLREFVFKNKDGFLYVYKGYQPIDVVRDLP SGFNTLKPIFKLPLGINITNFRAILTAFSPAQDIWGTSAAAY FVGYLKPTTFMLKYDENGTITDAVDCSQNPLAELKCSVKSFE IDKGIYQTSNFRVVPSGDVVRFPNITNLCPFGEVFNATKFPS VYAWERKKISNCVADYSVYLNSTFFSTFKCYGVSATKLNDLC FSNVYADSFVVKGDDVRQIAPGQTGVIADYNYKLPDDFMGCV LÄWNTRNIDATSTGNYNYKYRYLRHGKLRPFERDISNVPFSP DGKPCTPPALNCYWPLNDYGFYTTTGIGYQPYRVVVLSFELL NAPATVCGFKLSTDLIKNQCVNFNFNGLTGTGVLTPSSKRFQ PFQQFGRDVSDFTDSVRDPKTSEILDISPCSFGGVSVITPGT NASSEVAVLYQDVNCTDVSTAIHADQLTPAWRIYSTGNNVFQ TQAGCLIGAEHVDTSYECDIPIGAGICASYHTVSLLRSTSQK SIVAYTMSLGADSSIAYSNNTIAIPTNFSISITTEVMPVSMA KTSVDCNMYICGDSTECANLLLQYGSFCTQLNRALSGIAAEQ DRNTREVFAQVKQMYKTPTLKYFGGFNFSQILPDPLKPTKRS FIBDLLFNKVTLADAGFMKQYGECLGDINARDLICAQKFNGL TVLPPLLTDDMIAAYTAALVSGTATAGWTFGAGAALQIPFAM QMAYRFNGIGVTQNVLYENQKQIANQFNKAISQIQESLTTTS TALGKLQDVVNQNAQALNTLVKQLSSNFGAISSVLNDILSRL DKVEAEVQIDRLITGRLQSLQTYVTQQLIRAAEIRASANLAA TKMSECVLGQSKRVDFCGKGYHLMSPPQAAPHGVVFLHVTVV PSQERNFTTAPAICHEGKAYFPREGVFVFNGTSWFITQRNFF SPQIITTDNTFVSGNCDVVIGIINNTVYDPLQPELDSFKEEL DKYFKNHTSPDVDLGDISGINASVVNIQKEIDRLNEVAKNLL ESLIDLQELGKYEQYIKWPWYVULGFIAGLIAIVMVTILLCC MTSCCSCLKGACSCGSCCKFDEDDSEPVLKGVKLHYT   | 29    |

286

| TABLE II-CONCINUED | ${	t TABLE}$ | 11-continued |  |
|--------------------|--------------|--------------|--|
|--------------------|--------------|--------------|--|

Betacoronavirus Amino Acid Sequences

SEQ ID
Amino Acid Sequence NO:

Human coronavirus OC43 (HCoV-OC43) Spike glycoprotein UniProtKB-P36334

Strain

MFLILLISLPTAFAVIGDLKCTSDNINDKDTGPPPISTDTVD VTNGLGTYYVLDRVYLNTTLFLNGYYPTSGSTYRNMALKGSV LLSRLWFKPPFLSDFINGIFAKVKNTKVIKDRVMYSEFPAIT IGSTFVNTSYSVVVQPRTINSTQDGDNKLQGLLEVSVCQYNM CEYPQTICHPNLGNHRKELWHLDTGVVSCLYKRNFTYDVNAD YLYFHFYQEGGTFYAYFTDTGVVTKFLFNVYLGMALSHYYVM PLTCNSKLTLEYWVTPLTSRQYLLAFNQDGIIFNAEDCMSDF MSEIKCKTQSIAPPTGVYELNGYTVQPIADVYRRKPNLPNCN IEAWLNDKSVPSPLNWERKTFSNCNFNMSSLMSFIQADSFTC NNIDAAKIYGMCFSSITIDKFAIPNGRKVDLQLGNLGYLQSF NYRIDTTATSCQLYYNLPAANVSVSRFNPSTWNKRFGFIEDS VFKPRPAGVLTNHDVVYAQHCFKAPKNFCPCKLNGSCVGSGP GKNNGIGTCPAGTNYLTCDNLCTPDPITFTGTYKCPQTKSLV GIGEHCSGLAVKSDYCGGNSCTCRPQAFLGWSADSCLQGDKC NIFANFILHDVNSGLTCSTDLOKANTDIILGVCVNYDLYGIL GOGIFVEVNATYYNSWONLLYDSNGNLYGFRDYIINRTFMIR SCYSGRVSAAFHANSSEPALLFRNIKCNYVFNNSLTROLOPI NYFDSYLGCVVNAYNSTAISVOTCDLTVGSGYCVDYSKNRRS RGAITTGYRFTNFEPFTVNSVNDSLEPVGGLYEIOIPSEFTI GNMVEFIOTSSPKVTIDCAAFVCGDYAACKSOLVEYGSFCDN INAILTEVNELLDTTQLQVANSLMNGVTLSTKLKDGVNFNVD DINFSPVLGCLGSECSKASSRSAIEDLLFDKVKLSDVGFVEA YNNCTGGAEIRDLICVQSYKGIKVLPPLLSENQISGYTLAAT SASLFPPWTAAAGVPFYLNVQYRINGLGVTMDVLSQNQKLIA NAFNNALYAIQEGFDATNSALVKIQAVVNANAEALNNLLQQL SNRFGAISASLQEILSRLDALEAEAQIDRLINGRLTALNAYV  ${\tt SQQLSDSTLVKFSAAQAMEKVNECVKSQSSRINFCGNGNHII}$  ${\tt SLVQNAPYGLYFIHFSYVPTKYVTARVSPGLCIAGDRGIAPK}$  ${\tt SGYFVNVNNTWMYTGSGYYYPEPITENNVVVMSTCAVNYTKA}$ PYVMLNTSIPNLPDFKEELDQWFKNQTSVAPDLSLDYINVTF LDLQVEMNRLQEAIKVLNQSYINLKDIGTYEYYVKWPWYVWL LICLAGVAMLVLLFFICCCTGCGTSCFKKCGGCCDDYTGYQE LVIKTSHDD

Human coronavirus HKU1 (isolate N5) (HCoV-HKU1) Spike glycoprotein UniProtKB-OOZME7  ${\tt MFLIIFILPTTLAVIGDFNCTNSFINDYNKTIPRISEDVVDV}$ SLGLGTYYVLNRVYLNTTLLFTGYFPKSGANFRDLALKGSIY LSTLWYKPPFLSDFNNGIFSKVKNTKLYVNNTLYSEFSTIVI GSVFVNTSYTIVVQPHNGILEITACQYTMCEYPHTVCKSKGS IRNESWHIDSSEPLCLFKKNFTYNVSADWLYFHFYQERGVFY AYYADVGMPTTFLFSLYLGTILSHYYVMPLTCNAISSNTDNE TLEYWVTPLSRRQYLLNFDEHGVITNAVDCSSSFLSEIQCKT QSFAPNTGVYDLSGFTVKPVATVYRRIPNLPDCDIDNWLNNV SVPSPLNWERRIFSNCNFNLSTLLRLVHVDSFSCNNLDKSKI FGSCFNSITVDKFAIPNRRRDDLQLGSSGFLQSSNYKIDISS SSCQLYYSLPLVNVTINNFNPSSWNRRYGFGSFNLSSYDVVY SDHCFSVNSDFCPCADPSVVNSCAKSKPPSAICPAGTKYRHC DLDTTLYVKNWCRCSCLPDPISTYSPNTCPQKKVVVGIGEHC PGLGINEEKCGTQLNHSSCFCSPDAFLGWSFDSCISNNRCNI FSNFIFNGINSGTTCSNDLLYSNTEISTGVCVNYDLYGITGQ GIFKEVSAAYYNNWQNLLYDSNGNIIGFKDFLTNKTYTILPC YSGRVSAAFYQNSSSPALLYRNLKCSYVLNNISFISQPFYFD SYLGCVLNAVNLTSYSVSSCDLRMGSGFCIDYALPSSRRKRR GISSPYRFVTFEPFNVSFVNDSVETVGGLFEIQIPTNFTIAG HEEFIOTSSPKVTIDCSAFVCSNYAACHDLLSEYGTFCDNIN SILNEVNDLLDITQLQVANALMQGVTLSSNLNTNLHSDVDNI DFKSLLGCLGSQCGSSSRSLLEDLLFNKVKLSDVGFVEAYNN CTGGSEIRDLLCVQSFNGIKVLPPILSETQISGYTTAATVAA MFPPWSAAAGVPFSLNVQYRINGLGVTMDVLNKNQKLIANAF NKALLSIONGFTATNSALAKIOSVVNANAOALNSLLOOLFNK FGAISSSLQEILSRLDNLEAQVQIDRLINGRLTALNAYVSQQ LSDITLIKAGASRAIEKVNECVKSOSPRINFCGNGNHILSLV ONAPYGLLFIHFSYKPTSFKTVLVSPGLCLSGDRGIAPKOGY FIKONDSWMFTGSSYYYPEPISDKNVVFMNSCSVNFTKAPFI YLNNSIPNI.SDFEAELSLWEKNHTSIAPNI.TENSHINATELD LYYEMNVIQESIKSLNSSFINLKEIGTYEMYVKWPWYIWLLI VILFIIFLMILFFICCCTGCGSACFSKCHNCCDEYGGHNDFV IKASHDD

Novel\_SARS\_S2

MFIFLLFLTLTSGSDLDRALSGIAAEQDRNTREVFAQVKQMY
KTPTLKYFGGFNFSQILPDPLKPTKRSFIEDLLFNKVTLADA
GFMKQYGECLGDINARDLICAQKFNGLTVLPPLLTDDMIAAY
TAALVSGTATAGWTFGAGAALQIPFAMQMAYRFNGIGVTQNV
LYENQKQIANQFNKAISQIQESLTTTSTALGKLQDVVNQNAQ
ALNTLVKQLSSNFGAISSVLMDILSRLDKVEAEVQIDRLTTG
RLQSLQTYVTQQLIRAAEIRASANLAATKMSECVLGQSKRVD

31

TABLE 11-continued

| Bet                    | acoronavirus Amino Acid Sequences   |               |
|------------------------|---|---------------|
| Strain                 | Amino Acid Sequence   | SEQ ID<br>NO: |
|                        | FCGKGYHLMSFPQAAPHGVVFLHVTYVPSQERNFTTAPAICH<br>EGKAYFPREGVFVFNGTSWFITQRNFFSPQIITTDNTFVSGN<br>CDVVIGIINNTVYDPLQPELDSFKEELDKYFKNHTSPDVDLG<br>DISGINASVVNIQKEIDRLNEVAKNLNESLIDLQELGKYEQY<br>IKWPWYWLGFIAGLIAIVMVTILLCCMTSCCSCLKGACSCG<br>SCCKFDEDDSEPVLKGVKLHYT   |               |
| Novel_MERS_S2          | MIHSVFLLMFLLTPTESDCKLPLGQSLCALPDTPSTLTPRSV RSVPGEMRLAS1AFNHPTQVDQLMSSYFKLSIPTMFSFGVTQ EYIQTTIQKVTVDCKQYVCNGFQKCEQLLREYGQFCSKINQA LHGANLRQDDSVRNLFASVKSSQSSPIIPGFGGDFNLTLLEP VSISTGSRSARSAIEDLLFDKVTIADPGYMQGYDDCMQQGPA SARDLICAQYVAGYKVLPPLMDVNMEAAYTSSLLGSIAGVGW TAGLSSFAAIPFAQSIFYRLNGVGITQQVLSENQKLIANKFN QALGAMQTGFTTTNEAFQKVQDAVNNNAQALSKLASELSNTF GAISASIGDIIQRLDVLEQDAQIDRLINGRLTTLNAFVAQQL VRSESAALSAQLAKDKVNECVKAQSKRSGFCGQGTHIVSFVV NAPNGLYFMHVGYYPSNHIEVVSAYGLCDAANPTNCIAPVNG YFIKTNNTRIVDEWSYTGSSFYAPEPITSLNTKYVAPQVTYQ NISTNLPPPLLCRSTGIDFQDELDEFFKNVSTSIPNFGSLTQ INTTLLDLTYEMLSLQQVVKALNESYIDLKELGNYTYYNKWP | 33            |
| Novel_Trimeric_SARS_S2 | MFIFLLFLTLTSGSDLDRALSGIAAEQDRNTREVFAQVKQMY KTPTLKYFGGFNFSQILPDPLKPTKRSFIEDLLFNKVTLADA GFMKQYGECLGDINARDLICAQKFNGLTVLPPLLTDDMIAAY TAALVSGTATAGMTFGAGAALQIPFAMQMAYRFNGIGVTQNV LYENQKQIANQFNKAISQIQESLTTTSTALGKLQDVVNQNAQ ALNTLVKQLSSNFGAISSVLNDILSRLDKVEAEVQIDRLITG RLQSLQTTVTQQLIRAAEIRASANLAATKMSECVLGQSKRVD FCGKGYHLMSFPQAAPHGVVFLHVTYVPSQERNFTTAPAICH EGKAYFPREGVFVFNGTSWFITQRNFFSPQIITTDNTFVSGN CDVVIGIINNTVYDPLQPELDSFKEELDKYFKNHTSPDVDLG DISGINASVVNIQKEIDRLMEVAKNLNESLIDLQELGKYBQY IKWPWYVWLGFIAGLIAIVMVTILLCCMTSCCSCLKGACSCG SCCKFDEDDSEPVLKGVKLHYT  | 34            |

TABLE 12

| F                    | ull-length S      | pike Glycoprotein | Amino Acid Se | quences (Homo sapiens strains)  |
|----------------------|-------------------|-------------------|---------------|---|
| GenBank<br>Accession | Country           | Collection Date   | Release Date  | Virus Name  |
| AFY13307             | United<br>Kingdom | 2012 Sep. 11      | 2012 Dec. 5   | Betacoronavirus England 1, complete genome  |
| AFS88936             | 12mgaviii         | 2012 Jun. 13      | 2012 Sep. 27  | Human betacoronavirus 2c<br>EMC/2012, complete genome   |
| AGG22542             | United<br>Kingdom | 2012 Sep. 19      | 2013 Feb. 27  | Human betacoronavirus 2c England-<br>Qatar/2012, complete genome  |
| AHY21469             | Jordan            | 2012              | 2014 May 4    | Human betacoronavirus 2c Jordan-<br>N3/2012 isolate MG167, complete<br>genome   |
| AGH58717             | Jordan            | 2012 April        | 2013 Mar. 25  | Human betacoronavirus 2c Jordan-<br>N3/2012, complete genome  |
| AGV08444             | Saudi<br>Arabia   | 2013 May 7        | 2013 Sep. 17  | Middle East respiratory syndrome<br>coronavirus isolate Al-<br>Hasa_12_2013, complete genome                                  |
| AGV08546             | Saudi<br>Arabia   | 2013 May 11       | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Al-  |
| AGV08535             | Saudi<br>Arabia   | 2013 May 12       | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Al-  |
| AGV08558             | Saudi<br>Arabia   | 2013 May 15       | 2013 Sep. 17  | Hasa_16_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Al-                                  |
| AGV08573             | Saudi<br>Arabia   | 2013 May 23       | 2013 Sep. 17  | Hasa_17_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Al-<br>Hasa_18_2013, complete genome |
| AGV08480             | Saudi<br>Arabia   | 2013 May 23       | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Al-Hasa_19_2013, complete genome   |

TABLE 12-continued

| F                    | ull-length S      | pike Glycoprotein | Amino Acid Se | quences (Homo sapiens strains)  |
|----------------------|-------------------|-------------------|---------------|---|
| GenBank<br>Accession | Country           | Collection Date   | Release Date  | Virus Name  |
| AGN70962             | Saudi<br>Arabia   | 2013 May 9        | 2013 Jun. 10  | Middle East respiratory syndrome<br>coronavirus isolate Al-<br>Hasa_1_2013, complete genome   |
| AGV08492             | Saudi<br>Arabia   | 2013 May 30       | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Al-Hasa_21_2013, complete genome   |
| AHI48517             | Saudi<br>Arabia   | 2013 May 2        | 2014 Feb. 6   | Middle East respiratory syndrome coronavirus isolate Al-  |
| AGN70951             | Saudi<br>Arabia   | 2013 Apr. 21      | 2013 Jun. 10  | Hasa_25_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Al-  |
| AGN70973             | Saudi<br>Arabia   | 2013 Apr. 22      | 2013 Jun. 10  | Hasa_2_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Al-   |
| AGN70929             | Saudi<br>Arabia   | 2013 May 1        | 2013 Jun. 10  | Hasa_3_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Al-   |
| <b>A</b> GV08408     | Saudi<br>Arabia   | 2012 Jun. 19      | 2013 Sep. 17  | Hasa_4_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Bisha_1_2012,   |
| AGV08467             | Saudi<br>Arabia   | 2013 May 13       | 2013 Sep. 17  | complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate  |
| AID50418             | United<br>Kingdom | 2013 Feb. 10      | 2014 Jun. 18  | Buraidah_1_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate England/2/2013,   |
| AJD81451             | United<br>Kingdom | 2013 Feb. 10      | 2015 Jan. 18  | complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate England/3/2013,  |
| <b>A</b> JD81440     | United<br>Kingdom | 2013 Feb. 13      | 2015 Jan. 18  | complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate England/4/2013,  |
| AHB33326             | France            | 2013 May 7        | 2013 Dec. 7   | complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate FRA/UAE,   |
| <b>A</b> IZ48760     | USA               | 2014 June         | 2014 Dec. 14  | complete genome Middle East respiratory syndrome coronavirus isolate Florida/USA- 2_Saudi Arabia_2014, complete                                       |
| <b>A</b> GV08455     | Saudi<br>Arabia   | 2013 Jun. 4       | 2013 Sep. 17  | genome<br>Middle East respiratory syndrome<br>coronavirus isolate Hafr-Al-  |
| AHI48561             | Saudi<br>Arabia   | 2013 Aug. 5       | 2014 Feb. 6   | Batin_1_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Hafr-Al-   |
| <b>A</b> HI48539     | Saudi<br>Arabia   | 2013 Aug. 28      | 2014 Feb. 6   | Batin_2_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Hafr-Al-   |
| AIZ74417             | France            | 2013 Apr. 26      | 2015 Mar. 10  | Batin_6_2013, complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Hu-France<br>(UAE) - FRA1_1627-<br>2013_BAL_Sanger, complete |
| <b>A</b> IZ74433     | France            | 2013 May 7        | 2015 Mar. 10  | genome Middle East respiratory syndrome coronavirus isolate Hu-France - FRA2_130569-2013_IS_HTS,  |
| AIZ74439             | France            | 2013 May 7        | 2015 Mar. 10  | complete genome Middle East respiratory syndrome coronavirus isolate Hu-France - FRA2_130569-2013_InSpu_Sanger,                                       |
| AIZ74450             | France            | 2013 May 7        | 2015 Mar. 10  | complete genome<br>Middle East respiratory syndrome<br>coronavirus isolate Hu-France -<br>FRA2_130569-2013_Isolate_Sanger,                            |
| AKK52602             | Saudi<br>Arabia   | 2015 Feb. 10      | 2015 Jun. 8   | complete genome Middle East respiratory syndrome coronavirus isolate Hu/Riyadh_KSA_2959_2015,   |
| AKK52612             | Saudi<br>Arabia   | 2015 Mar. 1       | 2015 Jun. 8   | complete genome Middle East respiratory syndrome coronavirus isolate Hu/Riyadh_KSA_4050_2015, complete genome   |

291

TABLE 12-continued

| F         | full-length S   |                 | Amino Acid Se | quences (Homo sapiens strains)  |  |  |
|-----------|-----------------|-----------------|---------------|---|--|--|
|           | GenBank         |                 |               |   |  |  |
| Accession | Country         | Collection Date | Release Date  | Virus Name  |  |  |
| AHN10812  | Saudi<br>Arabia | 2013 Nov. 6     | 2014 Mar. 24  | Middle East respiratory syndrome coronavirus isolate Jeddah_1_2013, complete genome                                       |  |  |
| AID55071  | Saudi<br>Arabia | 2014 Apr. 21    | 2014 Nov. 12  | Middle East respiratory syndrome<br>coronavirus isolate<br>Jeddah_C10306/KSA/2014-04-20,                                  |  |  |
| AID55066  | Saudi<br>Arabia | 2014            | 2014 Nov. 12  | complete genome Middle East respiratory syndrome coronavirus isolate Jeddah_C7149/KSA/2014-04-05, complete genome         |  |  |
| AID55067  | Saudi<br>Arabia | 2014            | 2014 Nov. 12  | Middle East respiratory syndrome<br>coronavirus isolate<br>Jeddah_C7569/KSA/2014-04-03,                                   |  |  |
| AID55068  | Saudi<br>Arabia | 2014 Apr. 7     | 2014 Nov. 12  | complete genome Middle East respiratory syndrome coronavirus isolate Jeddah_C7770/KSA/2014-04-07, complete genome         |  |  |
| AID55069  | Saudi<br>Arabia | 2014 Apr. 12    | 2014 Nov. 12  | Middle East respiratory syndrome coronavirus isolate Jeddah_C8826/KSA/2014-04-12, complete genome                         |  |  |
| AID55070  | Saudi<br>Arabia | 2014 Apr. 14    | 2014 Nov. 12  | Middle East respiratory syndrome coronavirus isolate Jeddah_C9055/KSA/2014-04-14, complete genome                         |  |  |
| AHE78108  | Saudi<br>Arabia | 2013 Nov. 5     | 2014 May 1    | Middle East respiratory syndrome<br>coronavirus isolate MERS-CoV-<br>Jeddah-human-1, complete genome                      |  |  |
| AKL59401  | South<br>Korea  | 2015 May 20     | 2015 Jun. 9   | Middle East respiratory syndrome<br>coronavirus isolate MERS-<br>CoV/KOR/KNIH/002_05_2015,                                |  |  |
| ALD51904  | Thailand        | 2015 Jun. 17    | 2015 Jul. 7   | complete genome Middle East respiratory syndrome coronavirus isolate MERS- CoV/THA/CU/17_06_2015,                         |  |  |
| AID55072  | Saudi<br>Arabia | 2014 Apr. 15    | 2014 Nov. 12  | complete genome Middle East respiratory syndrome coronavirus isolate Makkah_C9355/KSA/Makkah/2014- 04-15, complete genome |  |  |
| AHC74088  | Qatar           | 2013 Oct. 13    | 2013 Dec. 23  | Middle East respiratory syndrome coronavirus isolate Qatar3, complete   |  |  |
| AHC74098  | Qatar           | 2013 Oct. 17    | 2013 Dec. 23  | genome Middle East respiratory syndrome coronavirus isolate Qatar4, complete genome                                       |  |  |
| AHI48572  | Saudi<br>Arabia | 2013 Aug. 15    | 2014 Feb. 6   | Middle East respiratory syndrome coronavirus isolate Riyadh_14_2013, complete genome                                      |  |  |
| AGV08379  | Saudi<br>Arabia | 2012 Oct. 23    | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Riyadh_1_2012, complete genome                                       |  |  |
| AID55073  | Saudi<br>Arabia | 2014 Apr. 22    | 2014 Nov. 12  | Middle East respiratory syndrome coronavirus isolate Riyadh_2014KSA_683/KSA/2014, complete genome                         |  |  |
| AGV08584  | Saudi<br>Arabia | 2012 Oct. 30    | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Riyadh_2_2012, complete genome                                       |  |  |
| AGV08390  | Saudi<br>Arabia | 2013 Feb. 5     | 2013 Sep. 17  | Middle East respiratory syndrome coronavirus isolate Riyadh_3_2013, complete genome                                       |  |  |
| AHI48605  | Saudi<br>Arabia | 2013 Mar. 1     | 2014 Feb. 6   | Middle East respiratory syndrome coronavirus isolate Riyadh_4_2013, complete genome                                       |  |  |
| AHI48583  | Saudi<br>Arabia | 2013 Jul. 2     | 2014 Feb. 6   | Middle East respiratory syndrome coronavirus isolate Riyadh_5_2013, complete genome                                       |  |  |
| AHI48528  | Saudi<br>Arabia | 2013 Jul. 17    | 2014 Feb. 6   | Middle East respiratory syndrome coronavirus isolate Riyadh_9_2013, complete genome                                       |  |  |

TABLE 12-continued

| F                    | Full-length Spike Glycoprotein Amino Acid Sequences (Homo sapiens strains) |                 |              |  |  |
|----------------------|--|-----------------|--------------|--|--|
| GenBank<br>Accession | Country  | Collection Date | Release Date | Virus Name   |  |
| AHI48594             | Saudi<br>Arabia  | 2013 Jun. 12    | 2014 Feb. 6  | Middle East respiratory syndrome coronavirus isolate Taif_1_2013, complete genome                    |  |
| AHI48550             | Saudi<br>Arabia  | 2013 Jun. 12    | 2014 Feb. 6  | Middle East respiratory syndrome coronavirus isolate Wadi-Ad-Dawasir_1_2013, complete genome         |  |
| AIY60558             | United<br>Arab<br>Emirates   | 2014 Mar. 7     | 2014 Dec. 6  | Middle East respiratory syndrome coronavirus strain Abu Dhabi/Gayathi_UAE_2_2014, complete genome    |  |
| AIY60538             | United<br>Arab<br>Emirates   | 2014 Apr. 10    | 2014 Dec. 6  | Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_16_2014, complete genome           |  |
| AIY60528             | United<br>Arab<br>Emirates   | 2014 Apr. 10    | 2014 Dec. 6  | Middle East respiratory syndrome<br>coronavirus strain Abu<br>Dhabi_UAE_18_2014, complete            |  |
| AIY60588             | United<br>Arab<br>Emirates   | 2014 Apr. 13    | 2014 Dec. 6  | genome Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_26_2014, complete           |  |
| AIY60548             | United<br>Arab<br>Emirates   | 2014 Apr. 19    | 2014 Dec. 6  | genome Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_30_2014, complete           |  |
| AIY60568             | United<br>Arab<br>Emirates   | 2014 Apr. 17    | 2014 Dec. 6  | genome Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_33_2014, complete           |  |
| AIY60518             | United<br>Arab<br>Emirates   | 2014 Apr. 7     | 2014 Dec. 6  | genome Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_8_2014, complete            |  |
| AIY60578             | United<br>Arab<br>Emirates   | 2013 Nov. 15    | 2014 Dec. 6  | genome Middle East respiratory syndrome coronavirus strain Abu Dhabi_UAE_9_2013, complete            |  |
| AKJ80137             | China  | 2015 May 27     | 2015 Jun. 5  | genome<br>Middle East respiratory syndrome<br>coronavirus strain ChinaGD01,<br>complete genome       |  |
| AHZ64057             | USA  | 2014 May 10     | 2014 May 14  | Middle East respiratory syndrome coronavirus strain Florida/USA-2_Saudi Arabia_2014, complete genome |  |
| AKM76229             | Oman   | 2013 Oct. 28    | 2015 Jun. 23 | Middle East respiratory syndrome<br>coronavirus strain<br>Hu/Oman_2285_2013, complete                |  |
| AKM76239             | Oman   | 2013 Dec. 28    | 2015 Jun. 23 | genome Middle East respiratory syndrome coronavirus strain Hu/Oman_2874_2013, complete genome        |  |
| AKI29284             | Saudi<br>Arabia  | 2015 Jan. 6     | 2015 May 27  | Middle East respiratory syndrome<br>coronavirus strain Hu/Riyadh-KSA-<br>2049/2015, complete genome  |  |
| AKI29265             | Saudi<br>Arabia  | 2015 Jan. 21    | 2015 May 27  | Middle East respiratory syndrome coronavirus strain Hu/Riyadh-KSA-2343/2015, complete genome         |  |
| AKI29255             | Saudi<br>Arabia  | 2015 Jan. 21    | 2015 May 27  | Middle East respiratory syndrome coronavirus strain Hu/Riyadh-KSA-2345/2015, complete genome         |  |
| AKI29275             | Saudi<br>Arabia  | 2015 Jan. 26    | 2015 May 27  | Middle East respiratory syndrome coronavirus strain Hu/Riyadh-KSA-2466/2015, complete genome         |  |
| AKK52582             | Saudi<br>Arabia  | 2015 Feb. 10    | 2015 Jun. 8  | Middle East respiratory syndrome coronavirus strain Hu/Riyadh_KSA_2959_2015, complete genome         |  |
| AKK52592             | Saudi<br>Arabia  | 2015 Mar. 1     | 2015 Jun. 8  | Middle East respiratory syndrome coronavirus strain Hu/Riyadh_KSA_4050_2015, complete genome         |  |

TABLE 12-continued

|                      | Full-length S              | pike Glycoprotein | Amino Acid Se | quences (Homo sapiens strains)   |
|----------------------|----------------------------|-------------------|---------------|--|
| GenBank<br>Accession | Country                    | Collection Date   | Release Date  | Virus Name   |
| AHZ58501             | USA                        | 2014 Apr. 30      | 2014 May 13   | Middle East respiratory syndrome<br>coronavirus strain Indiana/USA-<br>1_Saudi Arabia_2014, complete<br>genome |
| AGN52936             | United<br>Arab<br>Emirates | 2013              | 2013 Jun. 10  | Middle East respiratory syndrome coronavirus, complete genome  |

|  | TABLE 13   |               |
|--|--|---------------|
| Description  | Sequence   | SEQ ID<br>NO: |
|  | MeV Nucleic Acid Sequences   |               |
| GC_F_MEASLES_B3.1 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 1864 | <del>-</del>   | 35            |
|  | TGCTGCAGGGGGCGTTGTAACAAAAAGGGAGAACAAG TTGGTATGTCAAGACCAGGCCTAAAGCCTGACCTTACA GGAACATCAAAATCCTATGTAAGATCGCTTTTGATGATA ATAGGCTGGAGCCTCGGTGGCCAAGCTTCTTGCCCCTT GGGCCTCCCCCCCCCTCCCCCTTCCTGCACCCGT ACCCCCGTGGTCTTTGAATAAAGTCTGAGTGGGCGGC |               |
| GC_F_MEASLES_B3.1<br>ORF Sequence, NT                                      | ATGGGTCTCAAGGTGAACGTCTCTGCCGTATTCATGGC AGTACTGTTAACTCTCCAAACACCCGCCGGTCAAATTC ATTGGGGCAATCTCTCTAAGATAGGGGTAGTAGGAATA GGAAGTGCAAGCTACAAAGTTATGACTCGTTCCAGCCA TCAATCATTAGTCATAAAATTAATGCCCAATATAACTCT                                  | 36            |

CCTCAATAACTGCACGAGGGTAGAGATTGCAGAATACA GGAGACTACTAAGAACAGTTTTGGAACCAATTAGGGAT

TABLE 13-continued

SEQ ID Description Sequence NO:

GCACTTAATGCAATGACCCAGAACATAAGGCCGGTTCA GAGCGTAGCTTCAAGTAGGAGACACAAGAGATTTGCG GGAGTAGTCCTGGCAGGTGCGGCCCTAGGTGTTGCCAC AGCTGCTCAGATAACAGCCGGCATTGCACTTCACCGGT  ${\tt CCATGCTGAACTCTCAGGCCATCGACAATCTGAGAGCG}$ AGCCTGGAAACTACTAATCAGGCAATTGAGGCAATCAG ACAAGCAGGGCAGGAGATGATATTGGCTGTTCAGGGTG TCCAAGACTACATCAATAATGAGCTGATACCGTCTATG AACCAGCTATCTTGTGATCTAATCGGTCAGAAGCTCGG GCTCAAATTGCTTAGATACTATACAGAAATCCTGTCATT ATTTGGCCCCAGCCTACGGGACCCCATATCTGCGGAGA TATCTATCCAGGCTTTGAGTTATGCACTTGGAGGAGAT ATCAATAAGGTGTTAGAAAAGCTCGGATACAGTGGAG GCGATTTACTAGGCATCTTAGAGAGCAGAGGAATAAAG GCTCGGATAACTCACGTCGACACAGAGTCCTACTTCAT AGTCCTCAGTATAGCCTATCCGACGCTGTCCGAGATTA AGGGGGTGATTGTCCACCGGCTAGAGGGGGTCTCGTAC AACATAGGCTCTCAAGAGTGGTATACCACTGTGCCCAA GTATGTTGCAACCCAAGGGTACCTTATCTCGAATTTTGA TGAGTCATCATGTACTTTCATGCCAGAGGGGACTGTGT GCAGCCAAAATGCCTTGTACCCGATGAGTCCTCTGCTC CAAGAATGCCTCCGGGGGTCCACCAAGTCCTGTGCTCG TACACTCGTATCCGGGTCTTTTTGGGAACCGGTTCATTTT ATCACAAGGGAACCTAATAGCCAATTGTGCATCAATTC TTTGTAAGTGTTACACAACAGGTACGATTATTAATCAA GACCCTGACAAGATCCTAACATACATTGCTGCCGATCG CTGCCCGGTAGTCGAGGTGAACGGCGTGACCATCCAAG TCGGGAGCAGGAGGTATCCAGACGCTGTGTACTTGCAC AGAATTGACCTCGGTCCTCCCATATCATTGGAGAGGTT GGACGTAGGGACAAATCTGGGGAATGCAATTGCCAAA TTGGAGGATGCCAAGGAATTGTTGGAATCATCGGACCA GATATTGAGAAGTATGAAAGGTTTATCGAGCACTAGCA  ${\tt TAGTCTACATCCTGATTGCAGTGTGTCTTGGAGGGTTGA}$ TAGGGATCCCCACTTTAATATGTTGCTGCAGGGGGGCGT  ${\tt TGTAACAAAAAGGGAGAACAAGTTGGTATGTCAAGAC}$ CAGGCCTAAAGCCTGACCTTACAGGAACATCAAAATCC TATGTAAGATCGCTTTGA

GC\_F\_MEASLES\_B3.1 mRNA Sequence (assumes T100 tail) mRNA Sequence Length: 1925 G\*GGGAAATAAGAGAGAAAAGAAGAAGTAAGAAGAAAT ATAAGAGCCACCATGGGTCTCAAGGTGAACGTCTCTGC CGTATTCATGGCAGTACTGTTAACTCTCCAAACACCCG  $\tt CCGGTCAAATTCATTGGGGCAATCTCTCTAAGATAGGG$ GTAGTAGGAATAGGAAGTGCAAGCTACAAAGTTATGA  $\tt CTCGTTCCAGCCATCAATCATTAGTCATAAAATTAATGC$ CCAATATAACTCTCCTCAATAACTGCACGAGGGTAGAG ATTGCAGAATACAGGAGACTACTAAGAACAGTTTTGGA ACCAATTAGGGATGCACTTAATGCAATGACCCAGAACA  ${\tt TAAGGCCGGTTCAGAGCGTAGCTTCAAGTAGGAGACAC}$ AAGAGATTTGCGGGAGTAGTCCTGGCAGGTGCGGCCCT AGGTGTTGCCACAGCTGCTCAGATAACAGCCGGCATTG CACTTCACCGGTCCATGCTGAACTCTCAGGCCATCGAC AATCTGAGAGCGAGCCTGGAAACTACTAATCAGGCAAT TGAGGCAATCAGACAAGCAGGGCAGGAGATGATATTG GCTGTTCAGGGTGTCCAAGACTACATCAATAATGAGCT GATACCGTCTATGAACCAGCTATCTTGTGATCTAATCG GTCAGAAGCTCGGGCTCAAATTGCTTAGATACTATACA GAAATCCTGTCATTATTTGGCCCCAGCCTACGGGACCC CATATCTGCGGAGATATCTATCCAGGCTTTGAGTTATGC ACTTGGAGGAGATATCAATAAGGTGTTAGAAAAGCTCG GATACAGTGGAGGCGATTTACTAGGCATCTTAGAGAGC AGAGGAATAAAGGCTCGGATAACTCACGTCGACACAG AGTCCTACTTCATAGTCCTCAGTATAGCCTATCCGACGC TGTCCGAGATTAAGGGGGTGATTGTCCACCGGCTAGAG GGGGTCTCGTACAACATAGGCTCTCAAGAGTGGTATAC CACTGTGCCCAAGTATGTTGCAACCCAAGGGTACCTTA TCTCGAATTTTGATGAGTCATCATGTACTTTCATGCCAG AGGGGACTGTGTGCAGCCAAAATGCCTTGTACCCGATG AGTCCTCTGCTCCAAGAATGCCTCCGGGGGTCCACCAA GTCCTGTGCTCGTACACTCGTATCCGGGTCTTTTGGGAA CCGGTTCATTTTATCACAAGGGAACCTAATAGCCAATT GTGCATCAATTCTTTGTAAGTGTTACACAACAGGTACG ATTATTAATCAAGACCCTGACAAGATCCTAACATACAT  $\tt TGCTGCCGATCGCTGCCCGGTAGTCGAGGTGAACGGCG$ TGACCATCCAAGTCGGGAGCAGGAGGTATCCAGACGCT GTGTACTTGCACAGAATTGACCTCGGTCCTCCCATATCA TTGGAGAGGTTGGACGTAGGGACAAATCTGGGGAATG CAATTGCCAAATTGGAGGATGCCAAGGAATTGTTGGAA

|  | TABLE 13-continued  |               |
|--|---|---------------|
| Description  | Sequence  | SEQ ID<br>NO: |
|  | TCATCGGACCAGATATTGAGAAGTATGAAAGGTTTATC GAGCACTAGCATAGTCTACATCCTGATTGCAGTGTGTC TTGGAGGGTTGATAGGGATCCCCACTTTAATATGTTGCT GCAGGGGCGTTGTAACAAAAAGGGAGAAACAAGTTGG TATGTCAAGACCAGGCCTAAAGCCTGACCTTACAGGAA CATCAAAATCCTATGTAAGATCGCTTTGATGATAATAG GCTGGAGCCTCGGTGGCCAAGCTTCTTGCCCCTTGGGC CTCCCCCAGCCCCTCCTCCCCTTCCTGCACCCGTACCC CCGTGGTCTTTGAATAAAAAAAAAA   |               |
| GC_F_MEASLES_D8 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 1864 | TCAAGCTTTTGGACCCTCGTACAGAAGCTAATACGACT CACTATAGGGAAATAAGAGAGAAAAGAAGAGTAAGAA GAAATATAAGAGCCACCATGGGTCTCAAGGTGAACGTC TCTGTCATATTCATGGCAGTACTGTTAACTCTCTAAACA CCCACCGGTCAAATCCATTGGGGCAATCTCTCTAAGAT AGGGGTGGTAGGGTAG   | 38            |
| GC_F_MEASLES_D8 ORF Sequence, NT   | ATGGGTCTCAAGGTGAACGTCTCTGTCATATTCATGGC AGTACTGTTAACTCTTCAAACACCCACCGGTCAAATCC ATTGGGGCAATCTCTCAAGATAGGGGTGGTAGGGGTA GGAAGTGCAAGCTACAAAGTTATGACTCGTTCCAGCCA TCAATCATTAGTCATAAAGTTAATGCCCAATATAACTCT CCTCAACAATTGCACGAGGGTAGGGATTGCAGAATACA GGAGACTACTGAACAACAGTTCTGGAACCAATTAGAGAT GCACTTAATGCAATGACCCAGAATATAAGACCGGTTCA GAGTGTAGCTCAAGTAGGAGACAACAAGAGATTTCGGG GAGTTGTCCTGGCAGGTGCGGCCCTAGGCGTTGCCACA GCTGCTCAAATAACAGCCGGTATTGACACTTCACCAGTC CATGCTGAACTCTCAAGCCATCGACAATCTGAGAGGA GCCTAGAAACTACTAATCAGGCAATTGAGGCAATCAGA CAAGCAGGGCAGG | 39            |

TABLE 13-continued

Description Sequence SEQ ID

CCAAGACTACATCAATAATGAGCTGATACCGTCTATGA ATCAACTATCTTGTGATTTAATCGGCCAGAAGCTAGGG CTCAAATTGCTCAGATACTATACAGAAATCCTGTCATT ATTTGGCCCCAGCTTACGGGACCCCATATCTGCGGAGA TATCTATCCAGGCTTTGAGCTATGCGCTTGGAGGAGAT ATCAATAAGGTGTTGGAAAAGCTCGGATACAGTGGAG GTGATCTACTGGGCATCTTAGAGAGCAGAGGAATAAAG GCCCGGATAACTCACGTCGACACAGAGTCCTACTTCAT TGTACTCAGTATAGCCTATCCGACGCTATCCGAGATTA AGGGGGTGATTGTCCACCGGCTAGAGGGGGTCTCGTAC AACATAGGCTCTCAAGAGTGGTATACCACTGTGCCCAA GTATGTTGCAACCCAAGGGTACCTTATCTCGAATTTTGA TGAGTCATCATGCACTTTCATGCCAGAGGGGACTGTGT GCAGCCAGAATGCCTTGTACCCGATGAGTCCTCTGCTC CAAGAATGCCTCCGGGGGTCCACTAAGTCCTGTGCTCG TACACTCGTATCCGGGTCTTTCGGGAACCGGTTCATTTT ATCACAGGGGAACCTAATAGCCAATTGTGCATCAATCC TTTGCAAGTGTTACACAACAGGAACAATCATTAATCAA GACCCTGACAAGATCCTAACATACATTGCTGCCGATCA CTGCCCGGTGGTCGAGGTGAATGGCGTGACCATCCAAG TCGGGAGCAGGAGGTATCCGGACGCTGTGTACTTGCAC AGGATTGACCTCGGTCCTCCCATATCTTTGGAGAGGTT GGACGTAGGGACAAATCTGGGGAATGCAATTGCTAAGT TGGAGGATGCCAAGGAATTGTTGGAGTCATCGGACCAG ATATTGAGGAGTATGAAAGGTTTATCGAGCACTAGTAT AGTTTACATCCTGATTGCAGTGTGTCTTGGAGGATTGAT AGGGATCCCCGCTTTAATATGTTGCTGCAGGGGGCGTT GTAACAAGAAGGGAGAACAAGTTGGTATGTCAAGACC AGGCCTAAAGCCTGATCTTACAGGAACATCAAAATCCT ATGTAAGGTCACTCTGA

G\*GGAAATAAGAGAGAAAAGAAGAGTAAGAAGAAAT

GC\_F\_MEASLES\_D8 mRNA Sequence (assumes T100 tail) Sequence Length: 1925

ATAAGAGCCACCATGGGTCTCAAGGTGAACGTCTCTGT  ${\tt CATATTCATGGCAGTACTGTTAACTCTTCAAACACCCAC}$  $\tt CGGTCAAATCCATTGGGGCAATCTCTCTAAGATAGGGG$  $\tt TGGTAGGGGTAGGAAGTGCAAGCTACAAAGTTATGACT$ CGTTCCAGCCATCAATCATTAGTCATAAAGTTAATGCC CAATATAACTCTCCTCAACAATTGCACGAGGGTAGGGA TTGCAGAATACAGGAGACTACTGAGAACAGTTCTGGAA CCAATTAGAGATGCACTTAATGCAATGACCCAGAATAT AAGACCGGTTCAGAGTGTAGCTTCAAGTAGGAGACACA AGAGATTTGCGGGAGTTGTCCTGGCAGGTGCGGCCCTA GGCGTTGCCACAGCTGCTCAAATAACAGCCGGTATTGC ACTTCACCAGTCCATGCTGAACTCTCAAGCCATCGACA ATCTGAGAGCGAGCCTAGAAACTACTAATCAGGCAATT GAGGCAATCAGACAAGCAGGGCAGGAGATGATATTGG CTGTTCAGGGTGTCCAAGACTACATCAATAATGAGCTG ATACCGTCTATGAATCAACTATCTTGTGATTTAATCGGC CAGAAGCTAGGGCTCAAATTGCTCAGATACTATACAGA AATCCTGTCATTATTTGGCCCCAGCTTACGGGACCCCAT ATCTGCGGAGATATCTATCCAGGCTTTGAGCTATGCGC TTGGAGGAGATATCAATAAGGTGTTGGAAAAGCTCGGA TACAGTGGAGGTGATCTACTGGGCATCTTAGAGAGCAG AGGAATAAAGGCCCGGATAACTCACGTCGACACAGAG TCCTACTTCATTGTACTCAGTATAGCCTATCCGACGCTA TCCGAGATTAAGGGGGTGATTGTCCACCGGCTAGAGGG GGTCTCGTACAACATAGGCTCTCAAGAGTGGTATACCA CTGTGCCCAAGTATGTTGCAACCCAAGGGTACCTTATC TCGAATTTTGATGAGTCATCATGCACTTTCATGCCAGAG GGGACTGTGTGCAGCCAGAATGCCTTGTACCCGATGAG TCCTCTGCTCCAAGAATGCCTCCGGGGGTCCACTAAGT CCTGTGCTCGTACACTCGTATCCGGGTCTTTCGGGAACC GGTTCATTTTATCACAGGGGAACCTAATAGCCAATTGT GCATCAATCCTTTGCAAGTGTTACACAACAGGAACAAT CATTAATCAAGACCCTGACAAGATCCTAACATACATTG CTGCCGATCACTGCCCGGTGGTCGAGGTGAATGGCGTG ACCATCCAAGTCGGGAGCAGGAGGTATCCGGACGCTGT GTACTTGCACAGGATTGACCTCGGTCCTCCCATATCTTT GGAGAGGTTGGACGTAGGGACAAATCTGGGGAATGCA ATTGCTAAGTTGGAGGATGCCAAGGAATTGTTGGAGTC ATCGGACCAGATATTGAGGAGTATGAAAGGTTTATCGA  $\tt GCACTAGTATAGTTTACATCCTGATTGCAGTGTGTCTTG$ GAGGATTGATAGGGATCCCCGCTTTAATATGTTGCTGC  $\tt AGGGGGCGTTGTAACAAGAAGGGAGAACAAGTTGGTA$ TGTCAAGACCAGGCCTAAAGCCTGATCTTACAGGAACA TCAAAATCCTATGTAAGGTCACTCTGATGATAATAGGC TGGAGCCTCGGTGGCCAAGCTTCTTGCCCCTTGGGCCTC

40

| Description      | Sequence   | SEQ II<br>NO: |
|------------------|--|---------------|
|                  | CCCCAGCCCTCCTCCCCTTCCTGCACCCGTACCCCCG  |               |
|                  | TGGTCTTTGAATAAAGTCTGAGTGGGCGGCAAAAAAA  |               |
|                  | AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA  |               |
|                  | AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA   |               |
|                  | AAAAAAAAAAAAAAAAATCTAG   |               |
| C_H_MEASLES_B3   | TCAAGCTTTTGGACCCTCGTACAGAAGCTAATACGACT   | 41            |
| Sequence, NT (5' | CACTATAGGGAAATAAGAGAGAAAAGAAGAGTAAGAA  |               |
| TR, ORF, 3'      | GAAATATAAGAGCCACCATGTCACCGCAACGAGACCG  |               |
| TTR)             | GATAAATGCCTTCTACAAAGATAACCCTTATCCCAAGG   |               |
| Sequence Length: | GAAGTAGGATAGTTATTAACAGAGAACATCTTATGATT   |               |
| :065             | GACAGACCCTATGTTCTGCTGGCTGTTCTGTTCGTCATG<br>TTTCTGAGCTTGATCGGATTGCTGGCAATTGCAGGCATT |               |
|                  | AGACTTCATCGGGCAGCCATCTACACCGCGGAGATCCA   |               |
|                  | TAAAAGCCTCAGTACCAATCTGGATGTGACTAACTCCA   |               |
|                  | TCGAGCATCAGGTCAAGGACGTGCTGACACCACTCTTT   |               |
|                  | AAAATCATCGGGGATGAAGTGGGCCTGAGAACACCTC  |               |
|                  | AGAGATTCACTGACCTAGTGAAATTCATCTCGGACAAG   |               |
|                  | ATTAAATTCCTTAATCCGGATAGGGAGTACGACTTCAG   |               |
|                  | AGATCTCACTTGGTGCATCAACCCGCCAGAGAGGATCA   |               |
|                  | AACTAGATTATGATCAATACTGTGCAGATGTGGCTGCT   |               |
|                  | GAAGAGCTCATGAATGCATTGGTGAACTCAACTCTACT   |               |
|                  | GGAGACCAGAACAACCACTCAGTTCCTAGCTGTCTCAA   |               |
|                  | AGGGAAACTGCTCAGGGCCCACTACAATCAGAGGTCA<br>ATTCTCAAACATGTCGCTGTCCTTGTTGGACTTGTACTT   |               |
|                  | AGGTCGAGGTTACAATGTGTCGTTGTTGGACTTGTACTT  |               |
|                  | CATCCCAGGGAATGTATGGGGGGAACCTACCTAGTTGAA  |               |
|                  | AAGCCTAATCTGAACAGCAAAGGGTCAGAGTTGTCACA   |               |
|                  | ACTGAGCATGTACCGAGTGTTTGAAGTAGGTGTGATCA   |               |
|                  | GAAACCCGGGTTTGGGGGCTCCGGTGTTCCATATGACA   |               |
|                  | AACTATTTTGAGCAACCAGTCAGTAATGGTCTCGGCAA   |               |
|                  | CTGTATGGTGGCTTTGGGGGAGCTCAAACTCGCAGCCC   |               |
|                  | TTTGTCACGGGGACGATTCTATCATAATTCCCTATCAGG  |               |
|                  | GATCAGGGAAAGGTGTCAGCTTCCAGCTCGTCAAGCTG   |               |
|                  | GGTGTCTGGAAATCCCCAACCGACATGCAATCCTGGGT<br>CCCCTTATCAACGGATGATCCAGTGGTAGACAGGCTTT   |               |
|                  | ACCTCTCATCACAGAGGTGTCATCGCTGACAATCAA   |               |
|                  | GCAAAATGGGCTGTCCCGACAACACGAACAGATGACA  |               |
|                  | AGTTGCGAATGGAGACATGCTTCCAGCAGGCGTGTAAA   |               |
|                  | GGTAAAATCCAAGCACTCTGCGAGAATCCCGAGTGGGT   |               |
|                  | ACCATTGAAGGATAACAGGATTCCTTCATACGGGGTCC   |               |
|                  | TGTCTGTTGATCTGAGTCTGACGGTTGAGCTTAAAATCA  |               |
|                  | AAATTGCTTCGGGATTCGGGCCATTGATCACACACGGC   |               |
|                  | TCAGGGATGGACCTATACAAATCCAACTGCAACAATGT   |               |
|                  | GTATTGGCTGACTATTCCGCCAATGAGAAATCTAGCCT   |               |
|                  | TAGGCGTAATCAACACATTGGAGTGGATACCGAGATTC AAGGTTAGTCCCAACCTCTTCACTGTCCCAATTAAGGA      |               |
|                  | AGCAGGCGAAGACTGCCCAACTACCTG  |               |
|                  | CGGAGGTGGACGTGATGTCAAACTCAGTTCCAACCTG  |               |
|                  | GTGATTCTACCTGGTCAAGATCTCCAATATGTTTTTGGCA   |               |
|                  | ACCTACGATACCTCCAGGGTTGAGCATGCTGTGGTTTA   |               |
|                  | TTACGTTTACAGCCCAAGCCGCTCATTTTCTTACTTTTA  |               |
|                  | TCCTTTTAGGTTGCCTATAAAGGGGGTCCCAATCGAAC   |               |
|                  | TACAAGTGGAATGCTTCACATGGGATCAAAAACTCTGG   |               |
|                  | TGCCGTCACTTCTGTGTGCTTGCGGACTCAGAATCCGGT  |               |
|                  | GGACTTATCACTCACTCTGGGATGGTGGGCATGGGAGT   |               |
|                  | CAGCTGCACAGCTACCCGGGAAGATGGAACCAATCGC  |               |
|                  | AGATAATGATAATAGGCTGGAGCCTCGGTGGCCAAGCT<br>TCTTGCCCCTTGGGCCTCCCCCAGCCCCTCCTCCCCTT   |               |
|                  | CCTGCACCCGTACCCCCGTGGTCTTTGAATAAAGTCTG   |               |
|                  | AGTGGGCGGC   |               |
| C H MEASLES B3   | ATGTCACCGCAACGAGACCGGATAAATGCCTTCTACAA   | 42            |
| RF Sequence, NT  | AGATAACCCTTATCCCAAGGGAAGTAGGATAGTTATTA   | 44            |
| Dogaciico, Mi    | ACAGAGAACATCTTATGATTGACAGACCCTATGTTCTG   |               |
|                  | CTGGCTGTTCTGTTCGTCATGTTTCTGAGCTTGATCGGA  |               |
|                  | TTGCTGGCAATTGCAGGCATTAGACTTCATCGGGCAGC   |               |
|                  | CATCTACACCGCGGAGATCCATAAAAGCCTCAGTACCA   |               |
|                  | ATCTGGATGTGACTAACTCCATCGAGCATCAGGTCAAG   |               |
|                  | GACGTGCTGACACCACTCTTTAAAATCATCGGGGATGA   |               |
|                  | AGTGGGCCTGAGAACACCTCAGAGATTCACTGACCTAG   |               |
|                  | TGAAATTCATCTCGGACAAGATTAAATTCCTTAATCCG   |               |
|                  | GATAGGGAGTACGACTTCAGAGATCTCACTTGGTGCAT   |               |
|                  | CAACCCGCCAGAGAGGATCAAACTAGATTATGATCAAT   |               |

TABLE 13-continued

SEQ ID Description Sequence NO:

CCACTACAATCAGAGGTCAATTCTCAAACATGTCGCTG TCCTTGTTGGACTTGTACTTAGGTCGAGGTTACAATGTG TCATCTATAGTCACTATGACATCCCAGGGAATGTATGG GGGAACCTACCTAGTTGAAAAGCCTAATCTGAACAGCA AAGGGTCAGAGTTGTCACAACTGAGCATGTACCGAGTG TTTGAAGTAGGTGTGATCAGAAACCCGGGTTTGGGGGC TCCGGTGTTCCATATGACAAACTATTTTGAGCAACCAG TCAGTAATGGTCTCGGCAACTGTATGGTGGCTTTGGGG GAGCTCAAACTCGCAGCCCTTTGTCACGGGGACGATTC TATCATAATTCCCTATCAGGGATCAGGGAAAGGTGTCA GCTTCCAGCTCGTCAAGCTGGGTGTCTGGAAATCCCCA ACCGACATGCAATCCTGGGTCCCCTTATCAACGGATGA TCCAGTGGTAGACAGGCTTTACCTCTCATCTCACAGAG GTGTCATCGCTGACAATCAAGCAAAATGGGCTGTCCCG ACAACACGAACAGATGACAAGTTGCGAATGGAGACAT GCTTCCAGCAGGCGTGTAAAAGGTAAAATCCAAGCACTC TGCGAGAATCCCGAGTGGGTACCATTGAAGGATAACAG GATTCCTTCATACGGGGTCCTGTCTGTTGATCTGAGTCT GACGGTTGAGCTTAAAATCAAAATTGCTTCGGGATTCG GGCCATTGATCACACACGGCTCAGGGATGGACCTATAC AAATCCAACTGCAACAATGTGTATTGGCTGACTATTCC GCCAATGAGAAATCTAGCCTTAGGCGTAATCAACACAT TGGAGTGGATACCGAGATTCAAGGTTAGTCCCAACCTC TTCACTGTCCCAATTAAGGAAGCAGGCGAAGACTGCCA TGCCCCAACATACCTACCTGCGGAGGTGGACGGTGATG TCAAACTCAGTTCCAACCTGGTGATTCTACCTGGTCAA GATCTCCAATATGTTTTGGCAACCTACGATACCTCCAG GGTTGAGCATGCTGTGGTTTATTACGTTTACAGCCCAA  $\tt GCCGCTCATTTCTTACTTTTATCCTTTTAGGTTGCCTAT$ AAAGGGGTCCCAATCGAACTACAAGTGGAATGCTTCA CATGGGATCAAAAACTCTGGTGCCGTCACTTCTGTGTG CTTGCGGACTCAGAATCCGGTGGACTTATCACTCACTCT GGGATGGTGGGCATGGGAGTCAGCTGCACAGCTACCCG GGAAGATGGAACCAATCGCAGATAA

GC\_H\_MEASLES\_B3 mRNA Sequence (assumes T100 tail) Sequence Length: 2126 G\*GGGAAATAAGAGAGAAAAGAAGAAGAAAT ATAAGAGCCACCATGTCACCGCAACGAGACCGGATAA ATGCCTTCTACAAAGATAACCCTTATCCCAAGGGAAGT AGGATAGTTATTAACAGAGAACATCTTATGATTGACAG ACCCTATGTTCTGCTGGCTGTTCTGTTCGTCATGTTTCT GAGCTTGATCGGATTGCTGGCAATTGCAGGCATTAGAC  ${\tt TTCATCGGGCAGCCATCTACACCGCGGAGATCCATAAA}$ AGCCTCAGTACCAATCTGGATGTGACTAACTCCATCGA GCATCAGGTCAAGGACGTGCTGACACCACTCTTTAAAA TCATCGGGGATGAAGTGGGCCTGAGAACACCTCAGAG ATTCACTGACCTAGTGAAATTCATCTCGGACAAGATTA AATTCCTTAATCCGGATAGGGAGTACGACTTCAGAGAT CTCACTTGGTGCATCAACCCGCCAGAGAGGGATCAAACT AGATTATGATCAATACTGTGCAGATGTGGCTGCTGAAG AGCTCATGAATGCATTGGTGAACTCAACTCTACTGGAG ACCAGAACAACCACTCAGTTCCTAGCTGTCTCAAAGGG AAACTGCTCAGGGCCCACTACAATCAGAGGTCAATTCT CAAACATGTCGCTGTCCTTGTTGGACTTGTACTTAGGTC GAGGTTACAATGTGTCATCTATAGTCACTATGACATCC CAGGGAATGTATGGGGGAACCTACCTAGTTGAAAAGCC TAATCTGAACAGCAAAGGGTCAGAGTTGTCACAACTGA GCATGTACCGAGTGTTTGAAGTAGGTGTGATCAGAAAC CCGGGTTTGGGGGCTCCGGTGTTCCATATGACAAACTA TTTTGAGCAACCAGTCAGTAATGGTCTCGGCAACTGTA TGGTGGCTTTGGGGGAGCTCAAACTCGCAGCCCTTTGT CACGGGGACGATTCTATCATAATTCCCTATCAGGGATC AGGGAAAGGTGTCAGCTTCCAGCTCGTCAAGCTGGGTG TCTGGAAATCCCCAACCGACATGCAATCCTGGGTCCCC TTATCAACGGATGATCCAGTGGTAGACAGGCTTTACCT CTCATCTCACAGAGGTGTCATCGCTGACAATCAAGCAA AATGGGCTGTCCCGACAACACGAACAGATGACAAGTTG CGAATGGAGACATGCTTCCAGCAGGCGTGTAAAGGTAA AATCCAAGCACTCTGCGAGAATCCCGAGTGGGTACCAT TGAAGGATAACAGGATTCCTTCATACGGGGTCCTGTCT GTTGATCTGAGTCTGACGGTTGAGCTTAAAATCAAAAT TGCTTCGGGATTCGGGCCATTGATCACACACGGCTCAG GGATGGACCTATACAAATCCAACTGCAACAATGTGTAT TGGCTGACTATTCCGCCAATGAGAAATCTAGCCTTAGG CGTAATCAACACATTGGAGTGGATACCGAGATTCAAGG TTAGTCCCAACCTCTTCACTGTCCCAATTAAGGAAGCA GGCGAAGACTGCCATGCCCCAACATACCTACCTGCGGA GGTGGACGGTGATGTCAAACTCAGTTCCAACCTGGTGA

|  | TABLE 13-continued   |               |
|--|--|---------------|
| Description  | Sequence   | SEQ II<br>NO: |
|  | TTCTACCTGGTCAAGATCTCCAATATGTTTTTGGCAACCT ACGATACCTCCAGGGTTGAGCATGCTGTGTTTATTAC GTTTACAGCCCAAGCCGCTCATTTTCTTACTTTTATCCT TTTAGGTTGCCTATAAAGGGGGTCCCAATCGAACTACA AGTGGAATGCTTCACATGGGATCAAAAACTCTGGTGCC GTCACTTCTGTGTGCTTGCGGACTCAGAATCCGGTGGA CTTATCACTCACTCTGGGATGGTGGGCATCAGAATCCCAGAT AATGATAATAGGCTGGGCCTCGGTGGCCCAAGCTTCTT GCCCCTTGGGCTCCCCCCAGCCCTCCTCCCCTTCCTG CACCCGTACCCCGTGGTCTTTGAATAAAGTCTGAGTG GCCGCAAAAAAAAAA  |               |
| GC_H_MEASLES_D8 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 2065 | TCAAGCTTTTGGACCCTCGTACAGAAGCTAATACGACT CACTATAGGGAAATAAGAGAGAAAAGAAGATAAGAA GAAATATAAGAGCCACCATGTCACCACAACGAGACCG GATAAATGCCTTCTACAAAAGACAACCCCCATCCTAAGG GAAGTAGGATAGTTATTAACAAGAGAAACCCCCATCCTAAGG GAAGTAGGATAGTTATTATACAAGAGAAACCCCCATCCTAAGG GAAGTAGGATAGTTTTTCTTGCTGGTTTTTATTCGTCATG TTTCTGAGCTTGATCGGGTTGCTAGCCATTGCAGGCATT AGACCTTCATCGGGCAGCCATCTACACCGCAAGAGATCCA TAAAAGCCTCAGCACCAATCTGGATGTAACTAACTCAA TCGAGCATCAGGTAAGGACACCTCTACACCACAAGAATCCAA TCGAGCATCAGGTAAGAGACGTCTGACACCACACTCTTC AAGATCATCGGTGATGAAAGTCACTCCAAGA TCAAATCCCTGACTAGAAGTCATCTCTGACAAGA TTAAATTCCTTAATCCGGACAGGGAATACGACTCCACA GAATTCACTTGATCAGGACAGGA | 44            |
| GC_H_MEASLES_D8<br>ORF Sequence, NT                                      | ATGTCACCACAACGAGACCGGATAAATGCCTTCTACAA<br>AGACAACCCCCATCCTAAGGGAAGTAGGATAGTTATTA<br>ACAGAGAACATCTTATGATTGATAGACCTTATGTTTTGC  | 45            |

ATGTCACCACAACGAGACCGGATAAATGCCTTCTACAA AGACAACCCCCATCCTAAGGGAAGTAGGATAGTTATTA ACAGGAACATCTTATGATTGATAGACCTTATGTTTTGC TGGCTGTTCTATTCGTCATGTTTCTGAGCTTGATCGGGT TGCTAGCCATTGCAGGCATTAGACTTCATCGGCAGCC

### TABLE 13-continued

SEQ ID Description Sequence NO:

ATCTACACCGCAGAGATCCATAAAAGCCTCAGCACCAA TCTGGATGTAACTAACTCAATCGAGCATCAGGTTAAGG ACGTGCTGACACCACTCTTCAAGATCATCGGTGATGAA GTGGGCTTGAGGACACCTCAGAGATTCACTGACCTAGT GAAGTTCATCTCTGACAAGATTAAATTCCTTAATCCGG ACAGGGAATACGACTTCAGAGATCTCACTTGGTGTATC AACCCGCCAGAGAGAATCAAATTGGATTATGATCAATA CTGTGCAGATGTGGCTGCTGAAGAACTCATGAATGCAT TGGTGAACTCAACTCTACTGGAGACCAGGGCAACCAAT CAGTTCCTAGCTGTCTCAAAGGGAAACTGCTCAGGGCC CACTACAATCAGAGGCCAATTCTCAAACATGTCGCTGT CCCTGTTGGACTTGTATTTAAGTCGAGGTTACAATGTGT CATCTATAGTCACTATGACATCCCAGGGAATGTACGGG GGAACTTACCTAGTGGAAAAGCCTAATCTGAGCAGCAA AGGGTCAGAGTTGTCACAACTGAGCATGCACCGAGTGT TTGAAGTAGGTGTTATCAGAAATCCGGGTTTGGGGGCT CCGGTATTCCATATGACAAACTATCTTGAGCAACCAGT CAGTAATGATTTCAGCAACTGCATGGTGGCTTTGGGGG AGCTCAAGTTCGCAGCCCTCTGTCACAGGGAAGATTCT ATCACAATTCCCTATCAGGGATCAGGGAAAGGTGTCAG CTTCCAGCTTGTCAAGCTAGGTGTCTGGAAATCCCCAA CCGACATGCAATCCTGGGTCCCCCTATCAACGGATGAT CCAGTGATAGACAGGCTTTACCTCTCATCTCACAGAGG CGTTATCGCTGACAATCAAGCAAAATGGGCTGTCCCGA CAACACGGACAGATGACAAGTTGCGAATGGAGACATG CTTCCAGCAGGCGTGTAAGGGTAAAATCCAAGCACTTT GCGAGAATCCCGAGTGGACACCATTGAAGGATAACAG GATTCCTTCATACGGGGTCTTGTCTGTTGATCTGAGTCT  ${\tt GACAGTTGAGCTTAAAATCAAAATTGTTTCAGGATTCG}$ GGCCATTGATCACACACGGTTCAGGGATGGACCTATAC AAATCCAACCACAACAATATGTATTGGCTGACTATCCC GCCAATGAAGAACCTGGCCTTAGGTGTAATCAACACAT  ${\tt TGGAGTGGATACCGAGATTCAAGGTTAGTCCCAACCTC}$ TTCACTGTTCCAATTAAGGAAGCAGGCGAGGACTGCCA  $\tt TGCCCCAACATACCTACCTGCGGAGGTGGATGGTGATG$  ${\tt TCAAACTCAGTTCCAATCTGGTGATTCTACCTGGTCAAG}$ ATCTCCAATATGTTCTGGCAACCTACGATACTTCCAGA  $\tt GTTGAACATGCTGTAGTTTATTACGTTTACAGCCCAAGC$ CGCTCATTTTCTTACTTTTATCCTTTTAGGTTGCCTGTAA GGGGGTCCCCATTGAATTACAAGTGGAATGCTTCACA TGGGACCAAAAACTCTGGTGCCGTCACTTCTGTGTGCTT GCGGACTCAGAATCTGGTGGACATATCACTCACTCTGG GATGGTGGGCATGGGAGTCAGCTGCACAGCCACTCGGG AAGATGGAACCAGCCGCAGATAG

GC\_H\_MEASLES\_D8 mRNA Sequence (assumes T100 tail) Sequence Length: 2126

G\*GGAAATAAGAGAGAAAAGAAGAGTAAGAAGAAATATAAGAGCCACCATGTCACCACAACGAGACCGGATAA ATGCCTTCTACAAAGACAACCCCCATCCTAAGGGAAGT AGGATAGTTATTAACAGAGAACATCTTATGATTGATAG ACCTTATGTTTTGCTGGCTGTTCTATTCGTCATGTTTCTG AGCTTGATCGGGTTGCTAGCCATTGCAGGCATTAGACT TCATCGGGCAGCCATCTACACCGCAGAGATCCATAAAA GCCTCAGCACCAATCTGGATGTAACTAACTCAATCGAG CATCAGGTTAAGGACGTGCTGACACCACTCTTCAAGAT CATCGGTGATGAAGTGGGCTTGAGGACACCTCAGAGAT TCACTGACCTAGTGAAGTTCATCTCTGACAAGATTAAA TTCCTTAATCCGGACAGGGAATACGACTTCAGAGATCT CACTTGGTGTATCAACCCGCCAGAGAGAATCAAATTGG ATTATGATCAATACTGTGCAGATGTGGCTGCTGAAGAA CTCATGAATGCATTGGTGAACTCAACTCTACTGGAGAC CAGGGCAACCAATCAGTTCCTAGCTGTCTCAAAGGGAA ACTGCTCAGGGCCCACTACAATCAGAGGCCAATTCTCA AACATGTCGCTGTCCCTGTTGGACTTGTATTTAAGTCGA GGTTACAATGTGTCATCTATAGTCACTATGACATCCCA GGGAATGTACGGGGGAACTTACCTAGTGGAAAAGCCT AATCTGAGCAGCAAAGGGTCAGAGTTGTCACAACTGAG CATGCACCGAGTGTTTGAAGTAGGTGTTATCAGAAATC CGGGTTTGGGGGCTCCGGTATTCCATATGACAAACTAT CTTGAGCAACCAGTCAGTAATGATTTCAGCAACTGCAT GGTGGCTTTGGGGGAGCTCAAGTTCGCAGCCCTCTGTC ACAGGGAAGATTCTATCACAATTCCCTATCAGGGATCA GGGAAAGGTGTCAGCTTCCAGCTTGTCAAGCTAGGTGT CTGGAAATCCCCAACCGACATGCAATCCTGGGTCCCCC TATCAACGGATGATCCAGTGATAGACAGGCTTTACCTC  ${\tt TCATCTCACAGAGGCGTTATCGCTGACAATCAAGCAAA}$ ATGGGCTGTCCCGACAACACGGACAGATGACAAGTTGC GAATGGAGACATGCTTCCAGCAGGCGTGTAAGGGTAA

### TABLE 13-continued

Description Sequence SEQ ID NO:

AATCCAAGCACTTTGCGAGAATCCCGAGTGGACACCAT
TGAAGGATTACAGGATTCCTCATACGGGGTCTTGTCT
GTTGATCTGAGTCTGAGTCTGACTCACACAAAAT
TGTTCAGGATTCGGCCATTGAGTCACACGGTTCAG

TGTTTCAGGATTCGGGCCATTGATCACACACGGTTCAG GGATGGACCTATACAAATCCAACCACAACAATATGTAT TGGCTGACTATCCCGCCAATGAAGAACCTGGCCTTAGG TGTAATCAACACATTGGAGTGGATACCGAGATTCAAGG TTAGTCCCAACCTCTTCACTGTTCCAATTAAGGAAGCA GGCGAGGACTGCCATGCCCAACATACCTACCTGCGGA GGTGGATGTGATGTCAAACTCAGTTCCAATCTGGTGA TTCTACCTGGTCAAGATCTCCAATATGTTCTGGCAACCT ACGATACTTCCAGAGTTGAACATGCTGTAGTTTATTAC GTTTACAGCCCAAGCCGCTCATTTTCTTACTTTTATCCT TTTAGGTTGCCTGTAAGGGGGGTCCCCATTGAATTACA AGTGGAATGCTTCACATGGGACCAAAAACTCTGGTGCC GTCACTTCTGTGTGCTTGCGGACTCAGAATCTGGTGGA CATATCACTCACTCTGGGATGGTGGGCATGGGAGTCAG CTGCACAGCCACTCGGGAAGATGGAACCAGCCGCAGA TAGTGATAATAGGCTGGAGCCTCGGTGGCCAAGCTTCT TGCCCCTTGGGCCTCCCCCAGCCCCTCCTCCCCTTCCT GCACCCGTACCCCCGTGGTCTTTGAATAAAGTCTGAGT GGGCGCCAAAAAAAAAAAAAAAAAAAAAAAAAAAA ΤΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑ

## MeV mRNA Sequences

GC\_F\_MEASLES\_B3.1 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 1864 UCAAGCUUUUGGACCCUCGUACAGAAGCUAAUACGAC UCACUAUAGGGAAAUAAGAGAGAAAAGAAGAGUAAG AAGAAAUAUAAGAGCCACCAUGGGUCUCAAGGUGAA CGUCUCUGCCGUAUUCAUGGCAGUACUGUUAACUCUC CAAACACCCGCCGGUCAAAUUCAUUGGGGCAAUCUCU CUAAGAUAGGGUAGUAGGAAUAGGAAGUGCAAGCU ACAAAGUUAUGACUCGUUCCAGCCAUCAAUCAUUAGU CAUAAAAUUAAUGCCCAAUAUAACUCUCCUCAAUAAC UGCACGAGGGUAGAGAUUGCAGAAUACAGGAGACUA CUAAGAACAGUUUUGGAACCAAUUAGGGAUGCACUU AAUGCAAUGACCCAGAACAUAAGGCCGGUUCAGAGCG UAGCUUCAAGUAGGAGACACAAGAGAUUUGCGGGAG UAGUCCUGGCAGGUGCGGCCCUAGGUGUUGCCACAGC UGCUCAGAUAACAGCCGGCAUUGCACUUCACCGGUCC AUGCUGAACUCUCAGGCCAUCGACAAUCUGAGAGCGA GCCUGGAAACUACUAAUCAGGCAAUUGAGGCAAUCAG ACAAGCAGGCAGGAGAUGAUAUUGGCUGUUCAGGG UGUCCAAGACUACAUCAAUAAUGAGCUGAUACCGUCU AUGAACCAGCUAUCUUGUGAUCUAAUCGGUCAGAAGC UCGGGCUCAAAUUGCUUAGAUACUAUACAGAAAUCCU GUCAUUAUUUGGCCCCAGCCUACGGGACCCCAUAUCU GCGGAGAUAUCUAUCCAGGCUUUGAGUUAUGCACUU GGAGGAGAUAUCAAUAAGGUGUUAGAAAAGCUCGGA UACAGUGGAGGCGAUUUACUAGGCAUCUUAGAGAGC AGAGGAAUAAAGGCUCGGAUAACUCACGUCGACACAG AGUCCUACUUCAUAGUCCUCAGUAUAGCCUAUCCGAC GCUGUCCGAGAUUAAGGGGGUGAUUGUCCACCGGCUA GAGGGGUCUCGUACAACAUAGGCUCUCAAGAGUGG UAUACCACUGUGCCCAAGUAUGUUGCAACCCAAGGGU ACCUUAUCUCGAAUUUUGAUGAGUCAUCAUGUACUU UCAUGCCAGAGGGGACUGUGUGCAGCCAAAAUGCCUU GUACCCGAUGAGUCCUCUGCUCCAAGAAUGCCUCCGG GGGUCCACCAAGUCCUGUGCUCGUACACUCGUAUCCG GGUCUUUUGGGAACCGGUUCAUUUUAUCACAAGGGA ACCUAAUAGCCAAUUGUGCAUCAAUUCUUUGUAAGU GUUACACAACAGGUACGAUUAUUAAUCAAGACCCUGA CAAGAUCCUAACAUACAUUGCUGCCGAUCGCUGCCCG GUAGUCGAGGUGAACGGCGUGACCAUCCAAGUCGGGA GCAGGAGGUAUCCAGACGCUGUGUACUUGCACAGAAU UGACCUCGGUCCUCCCAUAUCAUUGGAGAGGUUGGAC GUAGGGACAAAUCUGGGGAAUGCAAUUGCCAAAUUG GAGGAUGCCAAGGAAUUGUUGGAAUCAUCGGACCAG AUAUUGAGAAGUAUGAAAGGUUUAUCGAGCACUAGC AUAGUCUACAUCCUGAUUGCAGUGUGUCUUGGAGGG UUGAUAGGGAUCCCCACUUUAAUAUGUUGCUGCAGG GGGCGUUGUAACAAAAAGGGAGAACAAGUUGGUAUG UCAAGACCAGGCCUAAAGCCUGACCUUACAGGAACAU CAAAAUCCUAUGUAAGAUCGCUUUGAUGAUAAUAGG CUGGAGCCUCGGUGGCCAAGCUUCUUGCCCCUUGGGC

TABLE 13-continued

|  | TABLE 13-continued  |               |
|--|---|---------------|
| Description  | Sequence  | SEQ II<br>NO: |
|  | CUCCCCCAGCCCCUCCUCCCCUUCCUGCACCCGUACC<br>CCCGUGGUCUUUGAAUAAAGUCUGAGUGGGCGGC   |               |
| GC_F_MEASLES_B3.1 ORF Sequence, NT   | AUGGGUCUCAAGGUGAACGUCUCUGCCGUAUUCAUGG CAGUACUGUUAACUCUCCAAACACCCGCCGGUCAAAU UCAUUGGGGCAUCUCUCUAAGAUAGGGGUAGUAGG AAUAGGAAGUGCAAGCUACAAAGUUAUGACUCGUUC CAGCAUCAUCAUUAGUCAUAAAAUUAAUGCCCAAU AUAACUCUCUCCAAUAACCCACGAGGGUAGAGAUUG CAGAAUACAGGAGCAUCUAAGAACGUUUUGGAAC CAAUUAGGGAUCACUAAGAACGUUUUGGAAC CAAUUAGGGAUCACUAAGAACAGUUUUGGAAC CAAUUAGGGAUCACUAAGAACAGUUUUGGAAC CAAUUAGGGAUCACACACAGAGGCCCACAACAU AAGACAGUUUCAAGAGCGUACCUAAGAACAGUUUGGAAC CAAUUAGGGAUCCACACACGUGCUCAGAACACU UGCACUUCACCGGUCCAUGCUGAAACUCUCAGGCCAU GGACAUCACACGUGCUCAGAACUACUAAGACAGCCGGCAU UGCACUUCACCGGUCCAUGCUGAAACUACUAAUCAAG CAAUUGAGGCAAUCAAGACCAGGCAGACAA AAGAGAUUUGCCACGGCCAUCGAAACUACAAUA AUGACCUGUUCAGGCCAUCGAAACUACAAUA AUGAGCUGUUCAGGGCGCCCAAGACUACAAUA AUGAGCUGUUCAGGGCGCCCAAGACUACAAUA AUGAGCUGUUCAGGGCGCCCAAGACUACAAUA AUGAGCUGAUACCGUCUAUGAACCAGCUAUCUUGUGA UCUAAUCGGUCAGACACCAGACUACAUCAAUA AUGAGCUGAUACCGUCUAUGAACCAGCUAUCUUGUGA UCUAAUCAGAAAUCCUGUCAUAUUUUGGCCCCAGCC UUUGAGUUAUGCACGUCUAUAUUUUGCCCCAGCC UUUGAGUUAUGCACCUUGAGAGACUAACACAGGC UUUGAGUAACAGAAGACAGAGGAGAAUACAAAAAAGGU GUUAGAAAAUCCUGGAGAGAAAAACAACAACAAAAAAGGU GUUAGAAAAACCCAAGAGUCCUACUUCAUAACGCAAGC UUUGAGUAACAGAAGACAAGAGGAAAAAAGGCUCGAAU AACUCACGUCAACACAAGAGUCCUACUUCAUAAGGCCCAAGAU AACUCAAGAGGAACCAAAGAGGAGAAUAAAGGCUCGAAU AAGCCAACACAAGAGUCCUACUUCAUAAGGGGG UGAUUGCCACCAGGCUGACCAAGAGUCCUCAACAACAU AGGCUCUUAAGAGAGGAGAAUAACAGCUCGAAUUAC AGGCUCUUAAAGAGAGACCUUAUUCAUAAGCCCAAGAUU GUUGCAACCCAAGGGUACCCUUAUCCAACACAU AGGCUCUUAAGAGAGGAGACCUUAUCCAACACAU AGGCUCUUAAGAGAGCACCAAGAGCCUGGUCCCCAAGUAU AGUCACCAAGAGUGCUUAUCAGAACCCUGCC CCAAGAAUGCCUCCCGAGGGGACCAAGACCCAAGGUCCCCAAGUCU AAUUCUUUGUAAGGAACCCAAGGUACCAACACAGGUCCCCAAGACCCCAAGGCACCAAGACCCCAAGGCACGACGA | 70            |
| GC_F_MEASLES_B3.1 mRNA Sequence (assumes T100 tail) mRNA Sequence Length: 1925 | G*GGAAAUAAGAGAAAAAGAAGAUAAGAAGAAA UAUAAGAGCCACCAUGGGUCUCAAGGUGAACGUCUCU GCCGUAUUCAUGGGCAGUACUGUUAACUCUCCAAACAC CCGCCGGUCAAAUUCAUUGGGCAAUCUCUUCUAAGAU AGGGGUAGAAAUUCAUUGGGCAAUCUCUCUAAGAU UAUGACUCGUUCCAGCAUCAAUCAUUAGUCAUAAAA UUAAUGCCCAAUAUAACUCUCCUCAAUAACUGCACGA GGGUAGAGAUUGCAGAAUACAGAAGCUACUAAGAA UAACGCCAAUAUAACUCUCCUCAAUAACUGCACGA CAGUUUUGGAACCAAUAAGGAGCUACUAAAAA UGACCCAGAACAUAAGGAGCUACUAAAAA UGACCCAGAACAUAAGGCAGCUUCAAGAA CAGUUUUGGAACCAAUAGGGAUGCCUUAAAUCA AGUAGGAGGCCCUAGGUGUUCACGGGAGUAGUCU GGCAGGUGCGGCCUUAGGGAUUUCACCGGUCCUAG AUAACAGCCGGCAUUGCAACACUCCACCGGUCCUAGA ACUCUCAAGCCAGCAAUCUGAGAGCGAGCCUGGA AACUACUAAUCAGGCAAUCUGAGAGCAGCCUGGA GACUACCUAAUCAGGCAAUCUGAGAGCCAGCCUGA GACUACUAAUCAGGCAAUCGAGAACCAGCCUGACAAGCA GGCAGGAGAUGAUAAUCGGUCAGAGCCAGCCUCAA GACUACUAAUCAGGCAAUCGAAAACCCGUCCAAU CUAAAUGACUCAAAUACGGACAAUCUGAGAGCAACC AGCUACCUUAAAUAAUCAGCAAAUCCUGUCAUU AUUUGGCCCCAGCCUACGGGACCCCAUAUCUGCGGCG CAAAUUGCUUAGAUACAGAAAUCCGGCCUGAAGGA AUAUCAAUAAAGGCGGCUUGAGAAGCCAGAGUCCGAGGAGAGAAUCCAGAAACACAUACAGAAAUCCGGAGAAUCCAGAGACACAGAGUCCUAAAUCAAUAAAGGCUUUAGAAAAAGCUCGGACACAGAGUCCUAAAUCAAUAAAGGCUUAAGAAAAGCUCGGACACAGAGUCCU ACUUCAUAGUCCUCAGUAAACCCACCACAGAGUCCU ACUUCAUAGUCCUCAGUAUAGCCCUACAGAGUCCU ACUUCAUAGCCCCCAGAUAUCCCGACCCUGCCCACAGAGUCCU ACUUCAUAGCCCCCAGAUAUCCCGACCCUGCCCAGAGUCCU ACUUCAUAGGCGCUCCAAGAGUCCU ACUUCAUAGGCGGUGAUAACCCCACCAGAGUCCU ACUUCAUAGGCGGUGAUAACCCCACCGGCUAGAGGG GGUCUCGUACAACAUAGGCCUCCAAGAGUCCU   | 71            |

TABLE 13-continued

SEQ ID Description Sequence NO:

ACUGUGCCCAAGUAUGUUGCAACCCAAGGGUACCUUA UCUCGAAUUUUGAUGAGUCAUCAUGUACUUUCAUGCC AGAGGGGACUGUGUGCAGCCAAAAUGCCUUGUACCCG AUGAGUCCUCUGCUCCAAGAAUGCCUCCGGGGGUCCA CCAAGUCCUGUGCUCGUACACUCGUAUCCGGGUCUUU UGGGAACCGGUUCAUUUUAUCACAAGGGAACCUAAU AGCCAAUUGUGCAUCAAUUCUUUGUAAGUGUUACAC AACAGGUACGAUUAUUAAUCAAGACCCUGACAAGAUC CUAACAUACAUUGCUGCCGAUCGCUGCCCGGUAGUCG AGGUGAACGGCGUGACCAUCCAAGUCGGGAGCAGGAG GUAUCCAGACGCUGUGUACUUGCACAGAAUUGACCUC GGUCCUCCCAUAUCAUUGGAGAGGUUGGACGUAGGG ACAAAUCUGGGGAAUGCAAUUGCCAAAUUGGAGGAU GCCAAGGAAUUGUUGGAAUCAUCGGACCAGAUAUUG AGAAGUAUGAAAGGUUUAUCGAGCACUAGCAUAGUC UACAUCCUGAUUGCAGUGUGUCUUGGAGGGUUGAUA GGGAUCCCCACUUUAAUAUGUUGCUGCAGGGGGCGUU GUAACAAAAGGGAGAACAAGUUGGUAUGUCAAGAC CAGGCCUAAAGCCUGACCUUACAGGAACAUCAAAAUC CUAUGUAAGAUCGCUUUGAUGAUAAUAGGCUGGAGC CUCGGUGGCCAAGCUUCUUGCCCCUUGGGCCUCCCCC CAGCCCCUCCUCCCUUCCUGCACCCGUACCCCGUGG UCUUUGAAUAAAGUCUGAGUGGGCGGCAAAAAAAAA AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA AAAAAAAAAAAAAAAAUCUAG

UCAAGCUUUUGGACCCUCGUACAGAAGCUAAUACGAC

GC\_F\_MEASLES\_D8 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 1864

UCACUAUAGGGAAAUAAGAGAGAAAAGAAGAGUAAG AAGAAAUAUAAGAGCCACCAUGGGUCUCAAGGUGAA CGUCUCUGUCAUAUUCAUGGCAGUACUGUUAACUCUU CAAACACCCACCGGUCAAAUCCAUUGGGGCAAUCUCU  ${\tt CUAAGAUAGGGGUGGUAGGGAAGUGCAAGCU}$ ACAAAGUUAUGACUCGUUCCAGCCAUCAAUCAUUAGU CAUAAAGUUAAUGCCCAAUAUAACUCUCCUCAACAAU UGCACGAGGGUAGGGAUUGCAGAAUACAGGAGACUA CUGAGAACAGUUCUGGAACCAAUUAGAGAUGCACUU AAUGCAAUGACCCAGAAUAUAAGACCGGUUCAGAGU GUAGCUUCAAGUAGGAGACACAAGAGAUUUGCGGGA GUUGUCCUGGCAGGUGCGGCCCUAGGCGUUGCCACAG CUGCUCAAAUAACAGCCGGUAUUGCACUUCACCAGUC CAUGCUGAACUCUCAAGCCAUCGACAAUCUGAGAGCG AGCCUAGAAACUACUAAUCAGGCAAUUGAGGCAAUCA GACAAGCAGGCAGGAGAUGAUAUUGGCUGUUCAGG GUGUCCAAGACUACAUCAAUAAUGAGCUGAUACCGUC UAUGAAUCAACUAUCUUGUGAUUUAAUCGGCCAGAA GCUAGGGCUCAAAUUGCUCAGAUACUAUACAGAAAUC CUGUCAUUAUUUGGCCCCAGCUUACGGGACCCCAUAU CUGCGGAGAUAUCUAUCCAGGCUUUGAGCUAUGCGCU UGGAGGAGAUAUCAAUAAGGUGUUGGAAAAGCUCGG AUACAGUGGAGGUGAUCUACUGGGCAUCUUAGAGAG CAGAGGAAUAAAGGCCCGGAUAACUCACGUCGACACA GAGUCCUACUUCAUUGUACUCAGUAUAGCCUAUCCGA CGCUAUCCGAGAUUAAGGGGGUGAUUGUCCACCGGCU AGAGGGGUCUCGUACAACAUAGGCUCUCAAGAGUG GUAUACCACUGUGCCCAAGUAUGUUGCAACCCAAGGG UACCUUAUCUCGAAUUUUGAUGAGUCAUCAUGCACUU UCAUGCCAGAGGGGACUGUGUGCAGCCAGAAUGCCUU GUACCCGAUGAGUCCUCUGCUCCAAGAAUGCCUCCGG GGGUCCACUAAGUCCUGUGCUCGUACACUCGUAUCCG GGUCUUUCGGGAACCGGUUCAUUUUAUCACAGGGGA ACCUAAUAGCCAAUUGUGCAUCAAUCCUUUGCAAGUG UUACACAACAGGAACAAUCAUUAAUCAAGACCCUGAC AAGAUCCUAACAUACAUUGCUGCCGAUCACUGCCCGG UGGUCGAGGUGAAUGGCGUGACCAUCCAAGUCGGGA GCAGGAGGUAUCCGGACGCUGUGUACUUGCACAGGAU UGACCUCGGUCCUCCCAUAUCUUUGGAGAGGUUGGAC GUAGGGACAAAUCUGGGGAAUGCAAUUGCUAAGUUG GAGGAUGCCAAGGAAUUGUUGGAGUCAUCGGACCAG AUAUUGAGGAGUAUGAAAGGUUUAUCGAGCACUAGU AUAGUUUACAUCCUGAUUGCAGUGUGUCUUGGAGGA UUGAUAGGGAUCCCCGCUUUAAUAUGUUGCUGCAGG GGGCGUUGUAACAAGAAGGGAGAACAAGUUGGUAUG UCAAGACCAGGCCUAAAGCCUGAUCUUACAGGAACAU CAAAAUCCUAUGUAAGGUCACUCUGAUGAUAAUAGG CUGGAGCCUCGGUGGCCAAGCUUCUUGCCCCUUGGGC CUCCCCCAGCCCCUCCUCCCCUUCCUGCACCCGUACC 72

|   | TABLE 13-continued  |              |
|---|---|--------------|
| Description   | Sequence  | SEQ I<br>NO: |
|   | CCCGUGGUCUUUGAAUAAAGUCUGAGUGGGCGGC  |              |
| GC_F_MEASLES_D8 ORF Sequence, NT  | AUGGGUCUCAAGUGAACGUCUCUGUCAUAUUCAUG GCAGUACUGUUAACUCUUCAAACACCCACCGUCAAA UCCAUUGGGGCAAUCUCUCUAAGAUAGGGGUGGUAG GGGUAGGAAGUGCAAGCUACAAAGUUAUGACUCGUU CCAGCCAUCAAUCAUUAGUCAUAAAGUUAUGACUCGUU CCAGCCAUCAAUCAUUAGUCAUAAAGUUAUGACUCGUU CCAGCAAUCAAUCAUUAGUCAUAAAGUUAUGACCCAA UAUAACUCUCCUCAACAAUUGCACGAGGGAUUCUGGAA CCAAUUAGAGACACAUUUAAUGCAAUGACCCAGAAUA UAAGACCGGUUCAGAGACGUUUCAAGUAGGAACA UAAGACCGGUUCAGAGUGUCAAGUAGCACCCAGAAUA UAAGACCGGUUCAGAGUGUUCAAGUAGGAGC CCCUAGGCGUUGCCACAGCUGCAAAUAACAGCCGG UAUUGCACUUCACCAGUCCAAGCUAACUAAACACCCG AUCGACAAUCAGAGAGCAGCGCUCAAAUAACACCCGG UAUUGCACUUCACCAGUCCAUGCUGAACCUACAACCA AUAAUGAGCUGUUCAAGGCCAGAGCAACAACAACAACAACAACACACCACAACA           | 73           |
| GC F_MEASLES_D8 mRNA Sequence (assumes T100 tail) Sequence Length: 1925 | G*GGAAAUAAGAGAGAAAAGAAGAGUAAAGAAA UAUAAGAGCCACCAUGGGUCUCAAGGUGAACGUCUCU GUCAUAUUCAUGGCAGUACUGUUAACUCUUCAAACAC CCACCGGUCAAAUCCAUUGGGGCAAUCUCUCUAAGAU AGGGGUGGUAGGGGUAGGAAGUGCAAGCUACAAAGU UAUGACCCAAUAUAACCAC GGGUAGGGUUGCAGCAUCAAUCAUAAGUCAAAAGU UAAUGCCCAAUAUAACUCUCCUCAACAAUUGCACGA GGGUAGGGAUUGCAGAAUACACACAUUGCACGA GGGUAGGGAUUGCAGAAUACACACAUUGCACGA CAGUUCUGGAACCAAUAGAGAGACUACUAAUGCAA UGACCCAGAAUAUAAGAGAUGCACUUAAUGCAA AGUACGGAGCCCUAGGCGGUUCAGAGGUUGUCCU GGCAGGGGCCCUAGGCGUUCACACACUGCUCAA AUAACAGCCGGUAUUGCACACACGUGCUCAA ACUCUCAAGCCACACUGCACACGUGCCAA ACUCUCAAGCCAUAUUGAGCAAUCAGAGACAACAGCA GGCAGGAGAUAUUAGAGCAAUCAGAGACAACCA GACUACUAAUCAGGCAUUCACCAGUCCUAUAAACACACAC | 74           |

GGUCUCGUACAACAUAGGCUCUCAAGAGUGGUAUACC ACUGUGCCCAAGUAUGUUGCAACCCAAGGGUACCUUA Sequence

320

SEO ID

NO:

UCUCGAAUUUUGAUGAGUCAUCAUGCACUUUCAUGCC AGAGGGGACUGUGUGCAGCCAGAAUGCCUUGUACCCG AUGAGUCCUCUGCUCCAAGAAUGCCUCCGGGGGUCCA CUAAGUCCUGUGCUCGUACACUCGUAUCCGGGUCUUU CGGGAACCGGUUCAUUUUAUCACAGGGGAACCUAAUA GCCAAUUGUGCAUCAAUCCUUUGCAAGUGUUACACAA CAGGAACAAUCAUUAAUCAAGACCCUGACAAGAUCCU AACAUACAUUGCUGCCGAUCACUGCCCGGUGGUCGAG GUGAAUGGCGUGACCAUCCAAGUCGGGAGCAGGAGG UAUCCGGACGCUGUGUACUUGCACAGGAUUGACCUCG GUCCUCCCAUAUCUUUGGAGAGGUUGGACGUAGGGAC AAAUCUGGGGAAUGCAAUUGCUAAGUUGGAGGAUGC CAAGGAAUUGUUGGAGUCAUCGGACCAGAUAUUGAG GAGUAUGAAAGGUUUAUCGAGCACUAGUAUAGUUUA CAUCCUGAUUGCAGUGUGUCUUGGAGGAUUGAUAGG GAUCCCCGCUUUAAUAUGUUGCUGCAGGGGGCGUUGU AACAAGAAGGGAGAACAAGUUGGUAUGUCAAGACCA GGCCUAAAGCCUGAUCUUACAGGAACAUCAAAAUCCU AUGUAAGGUCACUCUGAUGAUAAUAGGCUGGAGCCU CGGUGGCCAAGCUUCUUGCCCCUUGGGCCUCCCCCCA GCCCUCCUCCCUUCCUGCACCCGUACCCCCGUGGUC UUUGAAUAAAGUCUGAGUGGGCGCCAAAAAAAAAA AAAAAAAAAAAAAAAUCUAG

GC\_H\_MEASLES\_B3 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 2065

Description

UCAAGCUUUUGGACCCUCGUACAGAAGCUAAUACGAC UCACUAUAGGGAAAUAAGAGAGAAAAGAAGAGUAAG AAGAAAUAUAAGAGCCACCAUGUCACCGCAACGAGAC CGGAUAAAUGCCUUCUACAAAGAUAACCCUUAUCCCA AGGGAAGUAGGAUAGUUAUUAACAGAGAACAUCUUA UGAUUGACAGACCCUAUGUUCUGCUGGCUGUUCUGUU  $\tt CGUCAUGUUUCUGAGCUUGAUCGGAUUGCUGGCAAU$ UGCAGGCAUUAGACUUCAUCGGGCAGCCAUCUACACC GCGGAGAUCCAUAAAAGCCUCAGUACCAAUCUGGAUG UGACUAACUCCAUCGAGCAUCAGGUCAAGGACGUGCU GACACCACUCUUUAAAAUCAUCGGGGAUGAAGUGGGC CUGAGAACACCUCAGAGAUUCACUGACCUAGUGAAAU UCAUCUCGGACAAGAUUAAAUUCCUUAAUCCGGAUAG GGAGUACGACUUCAGAGAUCUCACUUGGUGCAUCAAC CCGCCAGAGAGGAUCAAACUAGAUUAUGAUCAAUACU GUGCAGAUGUGGCUGCUGAAGAGCUCAUGAAUGCAU UGGUGAACUCAACUCUACUGGAGACCAGAACAACCAC UCAGUUCCUAGCUGUCUCAAAGGGAAACUGCUCAGGG CCCACUACAAUCAGAGGUCAAUUCUCAAACAUGUCGC UGUCCUUGUUGGACUUGUACUUAGGUCGAGGUUACA AUGUGUCAUCUAUAGUCACUAUGACAUCCCAGGGAAU GUAUGGGGGAACCUACCUAGUUGAAAAGCCUAAUCU GAACAGCAAAGGGUCAGAGUUGUCACAACUGAGCAU GUACCGAGUGUUUGAAGUAGGUGUGAUCAGAAACCC GGGUUUGGGGGCUCCGGUGUUCCAUAUGACAAACUA UUUUGAGCAACCAGUCAGUAAUGGUCUCGGCAACUGU AUGGUGGCUUUGGGGGAGCUCAAACUCGCAGCCCUUU GUCACGGGGACGAUUCUAUCAUAAUUCCCUAUCAGGG AUCAGGGAAAGGUGUCAGCUUCCAGCUCGUCAAGCUG GGUGUCUGGAAAUCCCCAACCGACAUGCAAUCCUGGG UCCCCUUAUCAACGGAUGAUCCAGUGGUAGACAGGCU UUACCUCUCAUCUCACAGAGGUGUCAUCGCUGACAAU CAAGCAAAAUGGGCUGUCCCGACAACACGAACAGAUG ACAAGUUGCGAAUGGAGACAUGCUUCCAGCAGGCGUG UAAAGGUAAAAUCCAAGCACUCUGCGAGAAUCCCGAG UGGGUACCAUUGAAGGAUAACAGGAUUCCUUCAUAC GGGGUCCUGUCUGUUGAUCUGAGUCUGACGGUUGAG CUUAAAAUCAAAAUUGCUUCGGGAUUCGGGCCAUUG AUCACACACGGCUCAGGGAUGGACCUAUACAAAUCCA ACUGCAACAAUGUGUAUUGGCUGACUAUUCCGCCAAU GAGAAAUCUAGCCUUAGGCGUAAUCAACACAUUGGA GUGGAUACCGAGAUUCAAGGUUAGUCCCAACCUCUUC ACUGUCCCAAUUAAGGAAGCAGGCGAAGACUGCCAUG CCCCAACAUACCUACCUGCGGAGGUGGACGGUGAUGU CAAACUCAGUUCCAACCUGGUGAUUCUACCUGGUCAA GAUCUCCAAUAUGUUUUGGCAACCUACGAUACCUCCA GGGUUGAGCAUGCUGUGGUUUAUUACGUUUACAGCC CAAGCCGCUCAUUUUCUUACUUUUAUCCUUUUAGGUU GCCUAUAAAGGGGGUCCCAAUCGAACUACAAGUGGAA UGCUUCACAUGGGAUCAAAAACUCUGGUGCCGUCACU UCUGUGUGCUUGCGGACUCAGAAUCCGGUGGACUUAU

TABLE 13-continued

|   | TABLE 13-continued   |               |
|---|--|---------------|
| Description   | Sequence   | SEQ ID<br>NO: |
|   | CACUCACUCUGGGAUGGUGGGCAUGGGAGUCAGCUGC ACAGCUACCCGGGAAGAUGGAACCAAUCGCAGAUAAU GAUAAUAGGCUGGAGCCUCGGUGGCCAAGCUUCUUGC CCCUUGGGCCUCCCCCCAGCCCCUCCUCCCCCUUCCUGC ACCCGUACCCCCGUGGUCUUUGAAUAAAGUCUGAGUG GGCGGC |               |
| GC_H_MEASLES_B3 ORF Sequence, NT  | AUGUCACCGCAACGAGACCGGAUAAAUGCCUUCUACA AAGAUAACCCUUAUCCCAAGGGAAGUAGGAUAGUUA UUAACAGAGAACAUCUUAUGAUUGACAGACCCUAUG UUCUGCUGGCUGUUCUGUU  | 76            |
| GC_H_MEASLES_B3<br>mRNA Sequence<br>(assumes T100 Tail)<br>Sequence Length:<br>2126 | G*GGGAAUAAGAGAGAAAAGAAGAUAAGAAGAAA UAUAAGAGCCACCAUGUCACCGCAACGAGACCGGAUA AAUGCCUUCUACAAAGAUAACCCUUAUCCCAAGGGAA GUAGGAUAGUUAUUAACAGAGAAACAUCUUAUGAUUG ACAGACCCUAUGUUCUGCUUCUGUUCUG                      | 77            |

UACGACUUCAGAGAUCUCACUUGGUGCAUCAACCCGC CAGAGAGGAUCAAACUAGAUUAUGAUCAAUACUGUG  ${\tt CAGAUGUGGCUGCUGAAGAGCUCAUGAAUGCAUUGG}$  $\tt UGAACUCAACUCUACUGGAGACCAGAACCACUCA$ GUUCCUAGCUGUCUCAAAGGGAAACUGCUCAGGGCCC ACUACAAUCAGAGGUCAAUUCUCAAACAUGUCGCUGU

### TABLE 13-continued

SEQ ID Description Sequence NO:

CCUUGUUGGACUUGUACUUAGGUCGAGGUUACAAUG UGUCAUCUAUAGUCACUAUGACAUCCCAGGGAAUGUA UGGGGGAACCUACCUAGUUGAAAAGCCUAAUCUGAAC AGCAAAGGGUCAGAGUUGUCACAACUGAGCAUGUACC GAGUGUUUGAAGUAGGUGUGAUCAGAAACCCGGGUU UGGGGGCUCCGGUGUUCCAUAUGACAAACUAUUUUG AGCAACCAGUCAGUAAUGGUCUCGGCAACUGUAUGGU GGCUUUGGGGGAGCUCAAACUCGCAGCCCUUUGUCAC GGGGACGAUUCUAUCAUAAUUCCCUAUCAGGGAUCAG GGAAAGGUGUCAGCUUCCAGCUCGUCAAGCUGGGUGU CUGGAAAUCCCCAACCGACAUGCAAUCCUGGGUCCCC UUAUCAACGGAUGAUCCAGUGGUAGACAGGCUUUACC UCUCAUCUCACAGAGGUGUCAUCGCUGACAAUCAAGC AAAAUGGGCUGUCCCGACAACACGAACAGAUGACAAG UUGCGAAUGGAGACAUGCUUCCAGCAGGCGUGUAAA GGUAAAAUCCAAGCACUCUGCGAGAAUCCCGAGUGGG UACCAUUGAAGGAUAACAGGAUUCCUUCAUACGGGG UCCUGUCUGUUGAUCUGAGUCUGACGGUUGAGCUUA AAAUCAAAAUUGCUUCGGGAUUCGGGCCAUUGAUCAC ACACGGCUCAGGGAUGGACCUAUACAAAUCCAACUGC AACAAUGUGUAUUGGCUGACUAUUCCGCCAAUGAGA AAUCUAGCCUUAGGCGUAAUCAACACAUUGGAGUGG AUACCGAGAUUCAAGGUUAGUCCCAACCUCUUCACUG UCCCAAUUAAGGAAGCAGGCGAAGACUGCCAUGCCC AACAUACCUACCUGCGGAGGUGGACGGUGAUGUCAAA CUCAGUUCCAACCUGGUGAUUCUACCUGGUCAAGAUC UCCAAUAUGUUUUGGCAACCUACGAUACCUCCAGGGU UGAGCAUGCUGUGGUUUAUUACGUUUACAGCCCAAGC CGCUCAUUUUCUUACUUUUAUCCUUUUUAGGUUGCCUA UAAAGGGGGUCCCAAUCGAACUACAAGUGGAAUGCU UCACAUGGGAUCAAAAACUCUGGUGCCGUCACUUCUG UGUGCUUGCGGACUCAGAAUCCGGUGGACUUAUCACU CACUCUGGGAUGGUGGGCAUGGGAGUCAGCUGCACAG CUACCCGGGAAGAUGGAACCAAUCGCAGAUAAUGAUA AUAGGCUGGAGCCUCGGUGGCCAAGCUUCUUGCCCCU UGGGCCUCCCCAGCCCCUCCUCCCCUUCCUGCACCC GUACCCCGUGGUCUUUGAAUAAAGUCUGAGUGGGCG GCAAAAAAAAAAAAAAAAAAAAAAAAAAAAA ΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑΑ AAAAAAAAAAAAAAAAAAAAAAAAAAAAACUCUAG

GC\_H\_MEASLES\_D8 Sequence, NT (5' UTR, ORF, 3' UTR) Sequence Length: 2065 UCAAGCUUUUGGACCCUCGUACAGAAGCUAAUACGAC UCACUAUAGGGAAAUAAGAGAGAAAAGAAGAGUAAG AAGAAAUAUAAGAGCCACCAUGUCACCACAACGAGAC CGGAUAAAUGCCUUCUACAAAGACAACCCCCAUCCUA AGGGAAGUAGGAUAGUUAUUAACAGAGAACAUCUUA UGAUUGAUAGACCUUAUGUUUUGCUGGCUGUUCUAU UCGUCAUGUUUCUGAGCUUGAUCGGGUUGCUAGCCAU UGCAGGCAUUAGACUUCAUCGGGCAGCCAUCUACACC GCAGAGAUCCAUAAAAGCCUCAGCACCAAUCUGGAUG UAACUAACUCAAUCGAGCAUCAGGUUAAGGACGUGCU GACACCACUCUUCAAGAUCAUCGGUGAUGAAGUGGGC UUGAGGACACCUCAGAGAUUCACUGACCUAGUGAAGU UCAUCUCUGACAAGAUUAAAUUCCUUAAUCCGGACAG GGAAUACGACUUCAGAGAUCUCACUUGGUGUAUCAAC CCGCCAGAGAAUCAAAUUGGAUUAUGAUCAAUAC UGUGCAGAUGUGGCUGCUGAAGAACUCAUGAAUGCA UUGGUGAACUCAACUCUACUGGAGACCAGGGCAACCA AUCAGUUCCUAGCUGUCUCAAAGGGAAACUGCUCAGG GCCCACUACAAUCAGAGGCCAAUUCUCAAACAUGUCG CUGUCCCUGUUGGACUUGUAUUUAAGUCGAGGUUAC AAUGUGUCAUCUAUAGUCACUAUGACAUCCCAGGGAA UGUACGGGGAACUUACCUAGUGGAAAAGCCUAAUC UGAGCAGCAAAGGGUCAGAGUUGUCACAACUGAGCA UGCACCGAGUGUUUGAAGUAGGUGUUAUCAGAAAUC CGGGUUUGGGGGCUCCGGUAUUCCAUAUGACAAACUA UCUUGAGCAACCAGUCAGUAAUGAUUUCAGCAACUGC AUGGUGGCUUUGGGGGAGCUCAAGUUCGCAGCCCUCU GUCACAGGGAAGAUUCUAUCACAAUUCCCUAUCAGGG AUCAGGGAAAGGUGUCAGCUUCCAGCUUGUCAAGCUA GGUGUCUGGAAAUCCCCAACCGACAUGCAAUCCUGGG UCCCCCUAUCAACGGAUGAUCCAGUGAUAGACAGGCU UUACCUCUCAUCUCACAGAGGCGUUAUCGCUGACAAU CAAGCAAAAUGGGCUGUCCCGACAACACGGACAGAUG ACAAGUUGCGAAUGGAGACAUGCUUCCAGCAGGCGUG UAAGGGUAAAAUCCAAGCACUUUGCGAGAAUCCCGAG UGGACACCAUUGAAGGAUAACAGGAUUCCUUCAUACG

### TABLE 13-continued

Description Sequence NO:

GGGUCUUGUCUGUUGAUCUGAGUCUGACAGUUGAGC UUAAAAUCAAAAUUGUUUCAGGAUUCGGGCCAUUGA UCACACACGGUUCAGGGAUGGACCUAUACAAAUCCAA CCACAACAAUAUGUAUUGGCUGACUAUCCCGCCAAUG AAGAACCUGGCCUUAGGUGUAAUCAACACAUUGGAG UGGAUACCGAGAUUCAAGGUUAGUCCCAACCUCUUCA CUGUUCCAAUUAAGGAAGCAGGCGAGGACUGCCAUGC CCCAACAUACCUACCUGCGGAGGUGGAUGGUGAUGUC AAACUCAGUUCCAAUCUGGUGAUUCUACCUGGUCAAG AUCUCCAAUAUGUUCUGGCAACCUACGAUACUUCCAG AGUUGAACAUGCUGUAGUUUAUUACGUUUACAGCCC AAGCCGCUCAUUUUCUUACUUUUAUCCUUUUAGGUUG CCUGUAAGGGGGUCCCCAUUGAAUUACAAGUGGAA UGCUUCACAUGGGACCAAAAACUCUGGUGCCGUCACU UCUGUGUGCUUGCGGACUCAGAAUCUGGUGGACAUA UCACUCACUCUGGGAUGGUGGGCAUGGGAGUCAGCUG CACAGCCACUCGGGAAGAUGGAACCAGCCGCAGAUAG UGAUAAUAGGCUGGAGCCUCGGUGGCCAAGCUUCUUG CCCCUUGGGCCUCCCCCAGCCCCUCCUCCCCUUCCUG CACCCGUACCCCGUGGUCUUUGAAUAAAGUCUGAGU GGGCGGC

GC\_H\_MEASLES\_D8
ORF Sequence, NT

AUGUCACCACAACGAGACCGGAUAAAUGCCUUCUACA AAGACAACCCCCAUCCUAAGGGAAGUAGGAUAGUUAU UAACAGAGAACAUCUUAUGAUUGAUAGACCUUAUGU UUUGCUGGCUGUUCUAUUCGUCAUGUUUCUGAGCUU GAUCGGGUUGCUAGCCAUUGCAGGCAUUAGACUUCAU CGGGCAGCCAUCUACACCGCAGAGAUCCAUAAAAGCC UCAGCACCAAUCUGGAUGUAACUAACUCAAUCGAGCA UCAGGUUAAGGACGUGCUGACACCACUCUUCAAGAUC AUCGGUGAUGAGUGGGCUUGAGGACACCUCAGAGA UUCACUGACCUAGUGAAGUUCAUCUCUGACAAGAUUA AAUUCCUUAAUCCGGACAGGGAAUACGACUUCAGAGA UCUCACUUGGUGUAUCAACCCGCCAGAGAGAAUCAAA UUGGAUUAUGAUCAAUACUGUGCAGAUGUGGCUGCU GAAGAACUCAUGAAUGCAUUGGUGAACUCAACUCUAC UGGAGACCAGGGCAACCAAUCAGUUCCUAGCUGUCUC AAAGGGAAACUGCUCAGGGCCCACUACAAUCAGAGGC CAAUUCUCAAACAUGUCGCUGUCCCUGUUGGACUUGU AUUUAAGUCGAGGUUACAAUGUGUCAUCUAUAGUCA CUAUGACAUCCCAGGGAAUGUACGGGGGAACUUACCU AGUGGAAAAGCCUAAUCUGAGCAGCAAAGGGUCAGA GUUGUCACAACUGAGCAUGCACCGAGUGUUUGAAGU AGGUGUUAUCAGAAAUCCGGGUUUGGGGGCUCCGGU AUUCCAUAUGACAAACUAUCUUGAGCAACCAGUCAGU AAUGAUUUCAGCAACUGCAUGGUGGCUUUGGGGGAG CUCAAGUUCGCAGCCCUCUGUCACAGGGAAGAUUCUA UCACAAUUCCCUAUCAGGGAUCAGGGAAAGGUGUCAG CUUCCAGCUUGUCAAGCUAGGUGUCUGGAAAUCCCCA ACCGACAUGCAAUCCUGGGUCCCCCUAUCAACGGAUG AUCCAGUGAUAGACAGGCUUUACCUCUCAUCUCACAG AGGCGUUAUCGCUGACAAUCAAGCAAAAUGGGCUGUC CCGACAACACGGACAGAUGACAAGUUGCGAAUGGAGA CAUGCUUCCAGCAGGCGUGUAAGGGUAAAAUCCAAGC ACUUUGCGAGAAUCCCGAGUGGACACCAUUGAAGGAU AACAGGAUUCCUUCAUACGGGGUCUUGUCUGUUGAUC UGAGUCUGACAGUUGAGCUUAAAAUCAAAAUUGUUU CAGGAUUCGGGCCAUUGAUCACACACGGUUCAGGGAU GGACCUAUACAAAUCCAACCACAACAAUAUGUAUUGG CUGACUAUCCCGCCAAUGAAGAACCUGGCCUUAGGUG UAAUCAACACAUUGGAGUGGAUACCGAGAUUCAAGG UUAGUCCCAACCUCUUCACUGUUCCAAUUAAGGAAGC AGGCGAGGACUGCCAUGCCCAACAUACCUACCUGCG GAGGUGGAUGUGAUGUCAAACUCAGUUCCAAUCUG GUGAUUCUACCUGGUCAAGAUCUCCAAUAUGUUCUGG CAACCUACGAUACUUCCAGAGUUGAACAUGCUGUAGU UUAUUACGUUUACAGCCCAAGCCGCUCAUUUUCUUAC UUUUAUCCUUUUAGGUUGCCUGUAAGGGGGGUCCCCA UUGAAUUACAAGUGGAAUGCUUCACAUGGGACCAAA AACUCUGGUGCCGUCACUUCUGUGUGCUUGCGGACUC AGAAUCUGGUGGACAUAUCACUCACUCUGGGAUGGU GGGCAUGGGAGUCAGCUGCACAGCCACUCGGGAAGAU GGAACCAGCCGCAGAUAG

GC\_H\_MEASLES\_D8
mRNA Sequence
(assumes T100 tail)

G\*GGGAAAUAAGAGAGAAAAGAAGAAGAAA UAUAAGAGCCACCAUGUCACCACAACGAGACCGGAUA AAUGCCUUCUACAAAGACAACCCCCAUCCUAAGGGAA 80

TABLE 13-continued

| Description      | Sequence  | SEQ ID<br>NO: |
|------------------|---|---------------|
| Sequence Length: | GUAGGAUAGUUAUUAACAGAGAACAUCUUAUGAUUG  |               |
| 2126             | AUAGACCUUAUGUUUUGCUGGCUGUUCUAUUCGUCA  |               |
|                  | UGUUUCUGAGCUUGAUCGGGUUGCUAGCCAUUGCAG  |               |
|                  | GCAUUAGACUUCAUCGGGCAGCCAUCUACACCGCAGA                                       |               |
|                  | GAUCCAUAAAAGCCUCAGCACCAAUCUGGAUGUAACU                                       |               |
|                  | AACUCAAUCGAGCAUCAGGUUAAGGACGUGCUGACAC                                       |               |
|                  | CACUCUUCAAGAUCAUCGGUGAUGAAGUGGGCUUGA  |               |
|                  | GGACACCUCAGAGAUUCACUGACCUAGUGAAGUUCAU                                       |               |
|                  | CUCUGACAAGAUUAAAUUCCUUAAUCCGGACAGGGAA                                       |               |
|                  | UACGACUUCAGAGAUCUCACUUGGUGUAUCAACCCGC                                       |               |
|                  | CAGAGAGAAUCAAAUUGGAUUAUGAUCAAUACUGUG  |               |
|                  | CAGAUGUGGCUGCUGAAGAACUCAUGAAUGCAUUGG  |               |
|                  | UGAACUCAACUCUACUGGAGACCAGGGCAACCAAUCA                                       |               |
|                  | GUUCCUAGCUGUCUCAAAGGGAAACUGCUCAGGGCCC                                       |               |
|                  | ACUACAAUCAGAGGCCAAUUCUCAAACAUGUCGCUGU                                       |               |
|                  | CCCUGUUGGACUUGUAUUUAAGUCGAGGUUACAAUG  |               |
|                  | UGUCAUCUAUAGUCACUAUGACAUCCCAGGGAAUGUA                                       |               |
|                  | CGGGGGAACUUACCUAGUGGAAAAGCCUAAUCUGAGC                                       |               |
|                  | AGCAAAGGGUCAGAGUUGUCACAACUGAGCAUGCACC                                       |               |
|                  | GAGUGUUUGAAGUAGGUGUUAUCAGAAAUCCGGGUU  |               |
|                  | UGGGGGCUCCGGUAUUCCAUAUGACAAACUAUCUUGA                                       |               |
|                  | GCAACCAGUCAGUAAUGAUUUCAGCAACUGCAUGGUG                                       |               |
|                  | GCUUUGGGGGAGCUCAAGUUCGCAGCCCUCUGUCACA                                       |               |
|                  | GGGAAGAUUCUAUCACAAUUCCCUAUCAGGGAUCAGG                                       |               |
|                  | GAAAGGUGUCAGCUUCCAGCUUGUCAAGCUAGGUGUC                                       |               |
|                  | UGGAAAUCCCCAACCGACAUGCAAUCCUGGGUCCCCC UAUCAACGGAUGAUCCAGUGAUAGACAGGCUUUACCU |               |
|                  | CUCAUCUCACAGAGGCGUUAUCGCUGACAAUCAAGCA                                       |               |
|                  | AAAUGGGCUGUCCCGACAACACGGACAGAUGACAAGU                                       |               |
|                  | UGCGAAUGGAGACAUGCUUCCAGCAGGCGUGUAAGG  |               |
|                  | GUAAAAUCCAAGCACUUUGCGAGAAUCCCGAGUGGAC                                       |               |
|                  | ACCAUUGAAGGAUAACAGGAUUCCUUCAUACGGGGUC                                       |               |
|                  | UUGUCUGUUGAUCUGAGUCUGACAGUUGAGCUUAAA  |               |
|                  | AUCAAAAUUGUUUCAGGAUUCGGGCCAUUGAUCACAC                                       |               |
|                  | ACGGUUCAGGGAUGGACCUAUACAAAUCCAACCACAA                                       |               |
|                  | CAAUAUGUAUUGGCUGACUAUCCCGCCAAUGAAGAAC                                       |               |
|                  | CUGGCCUUAGGUGUAAUCAACACAUUGGAGUGGAUA  |               |
|                  | CCGAGAUUCAAGGUUAGUCCCAACCUCUUCACUGUUC                                       |               |
|                  | CAAUUAAGGAAGCAGGCGAGGACUGCCAUGCCCCAAC                                       |               |
|                  | AUACCUACCUGCGGAGGUGGAUGGUGAUGUCAAACUC                                       |               |
|                  | AGUUCCAAUCUGGUGAUUCUACCUGGUCAAGAUCUCC                                       |               |
|                  | AAUAUGUUCUGGCAACCUACGAUACUUCCAGAGUUGA                                       |               |
|                  | ACAUGCUGUAGUUUAUUACGUUUACAGCCCAAGCCGC                                       |               |
|                  | UCAUUUUCUUACUUUUAUCCUUUUAGGUUGCCUGUA  |               |
|                  | AGGGGGGUCCCCAUUGAAUUACAAGUGGAAUGCUUC  |               |
|                  | ACAUGGGACCAAAAACUCUGGUGCCGUCACUUCUGUG                                       |               |
|                  | UGCUUGCGGACUCAGAAUCUGGUGGACAUAUCACUCA                                       |               |
|                  | CUCUGGGAUGGGGCAUGGGAGUCAGCUGCACAGCC   |               |
|                  | ACUCGGGAAGAUGGAACCAGCCGCAGAUAGUGAUAA  |               |
|                  | UAGGCUGGAGCCUCGGUGGCCAAGCUUCUUGCCCCUU                                       |               |
|                  | GGGCCUCCCCAGCCCCCCCCCCUUCCUCCCCUUCCUGCACCCG                                 |               |
|                  |   |               |
|                  | UACCCCCGUGGUCUUUGAAUAAAGUCUGAGUGGGCGG                                       |               |
|                  | CAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA   |               |
|                  | AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA                                     |               |
|                  | AAAAAAAAAAAAAAAAAAAAAAAAAAAUCUAG  |               |

# TABLE 14

|                                       | MeV Amino Acid Sequences  |               |
|---------------------------------------|---|---------------|
| Description                           | Sequence  | SEQ ID<br>NO: |
| GC_F_MEASLES_B3.1<br>ORF Sequence, AA | MGLKVNVSAVFMAVLLTLQTPAGQIHWGNLSKIGVVG IGSASYKVMTRSSHQSLVIKLMPNITLLNNCTRVEIA EYRRLLRTVLEPIRDALMAMTQNIRPVQSVASSRRHK RFAGVVLAGAALGVATAAQITAGIALHRSMLNSQAID NLRASLETTNQAIEAIRQAGQEMILAVQGVQDYINNE LIPSMNQLSCDLIGQKLGLKLLRYYTEILSLFGPSLR DPISAEISIQALSYALGGDINKVLEKLGYSGGDLLGI LESRGIKARITHVDTESYFIVLSIAYPTLSEIKGVIV HRLEGVSYNIGSQEWYTTVPKYVATQGYLISNFDESS CTFMPEGTVCSQNALYPMSPLLQECLRGSTKSCARTL VSGSFGNRFILSQGNLIANCASILCKCYTTGTIINQD | 47            |

TABLE 14-continued

|                                     | MeV Amino Acid Sequences   |               |
|-------------------------------------|--|---------------|
| Description                         | Sequence   | SEQ ID<br>NO: |
|                                     | PDKILTYIAADRCPVVEVNGVTIQVGSRRYPDAVYLH<br>RIDLGPPISLERLDVGTNLGNAIAKLEDAKELLESSD<br>QILRSMKGLSSTSIVYILIAVCLGGLIGIPTLICCCR<br>GRCNKKGEQVGMSRPGLKPDLTGTSKSYVRSL*   |               |
| GC_F_MEASLES_D8<br>ORF Sequence, AA | MGLKVNVSVIFMAVLLTLQTPTGQIHWGNLSKIGVVG VGSASYKVMTRSSHQSLVIKLMPNITLLNNCTRVGIA EYRRLLRTVLEPIRDALNAMTQNIRPVQSVASSRRHK RFAGVVLAGAALGVATAAQITAGIALHQSMLNSQAID NLRASLETTNQAIEAIRQAGQEMILAVQGVQDYINNE LIPSMNQLSCDLIGQKLGLKLLRYYTEILSLFGPSLR DPISAEISIQALSVALGGDINKVLEKLGYSGGDLLGI LESRGIKARITHVDTESYFIVLSIAYPTLSEIKGVIV HRLEGVSYNIGSQEWYTTVPKYVATQGYLISNFDESS CTFMPEGTVCSQNALYPMSPLLQECLRGSTKSCARTL VSGSFGNRFILSQGNLIANCASILCKCYTTGTIINQD PDKILTYIAADHCPVVEVNGVTIQVGSRRYPDAVYLH RIDLGPPISLERLDVGTNLGNAIAKLEDAKELLESSD QILRSMKGLSSTSIVYILIAVCLGGLIGIPALICCCR GRCNKKGEQVGMSRPGLKPDLTGTSKSYVRSL*  | 48            |
| GC_H_MEASLES_B3<br>ORF Sequence, AA | MSPQRDRINAFYKDNPYPKGSRIVINREHLMIDRPYV LLAVLFVMFLSLIGLLAIAGIRLHRAAIYTABIHKSL STNLDVTNSIEHQVKDVLTPLFKIIGDEVGLRTPQRF TDLVKFISDKIKFLNPDREYDFRDLTWCINPPERIKL DYDQYCADVAAEELMNALVNSTLLETRTTTCPLAVSK GNCSGPTTIRGQFSNMSLSLLDLYLGRGYNVSSIVTM TSQGMYGGTYLVEKPNLNSKGSELSQLSMYRVPEVGV IRNPGLGAPVFHMTNYFEQPVSNGLGNCMVALGELKL AALCHGDDSIIIPYQGSGKGVSFQLVKLGVWKSPTDM QSWVPLSTDDPVVDRLYLSSHRGVIADNQAKWAVPTT RTDDKLRMETCFQQACKGKIQALCENPEWVPLKDNRI PSYGVLSVDLSLTVELKIKIASGFGPLITHGSGMDLY KSNCNNVYWLTIPPMRNLALGVINTLEWIPRFKVSPN LFTVPIKEAGEDCHAPTYLPAEVDGDVKLSSNLVILP GQDLQYVLATYDTSRVEHAVVYYVYSPSRSFSYFYPF RLPIKGVPIELQVECFTWDQKLWCRHPCVLADSESGG LITHSGMVGMGVSCTATREDGTNRR* | 49            |
| GC_H_MEASLES_D8 ORF Sequence, AA    | MSPQRDRINAFYKDNPHPKGSRIVINREHLMIDRPYV LLAVLFVMFLSLIGLLAIAGIRLHRAAIYTAEIHKSL STNLDVTNSIEHQVKDVLTPLFKIIGDEVGLRTPQRF TDLVKFISDKIKFLNPDREYDFRDLTWCINPPERIKL DYDQYCADVAAEELMNALVNSTLLETRATNQFLAVSK GNCSGPTTIRGQFSNMSLSLLDLYLSRGYNVSSIVTM TSQGMYGGTYLVEKPNLSSKGSELSQLSMHRVFEVGV IRNPGLGAPVFHMTNYLEQPVSNDFSNCMVALGELKF AALCHREDSITIPYQGSGKGVSFQLVKLGVWKSPTDM QSWVPLSTDDPVIDRLYLSSHRGVIADNQAKWAVPTT RTDDKLRMETCFQQACKGKIQALCENPEWTPLKDNRI PSYGVLSVDLSLTVELKIKIVSGFGPLITHGSGMDLY KSNHNNMYWLTIPPMKNLALGVINTLEWIPFKVSPN LFTVPIKEAGEDCHAPTYLPAEVDGDVKLSSNLVILP GQDLQYVLATYDTSRVEHAVVYYVYSPSRSFSYFYPF RLPVRGVPIELQVECFTWDQKLWCRHFCVLADSESGG HITHSGMVGMGVSCTATREDGTSRR*  | 50            |

TABLE 15

| MeV NCBI Accession Numbers (Amino Acid Sequences) |   |                   |  |
|---|---|-------------------|--|
| Туре  | Virus Name                                    | GenBank Accession |  |
| hemagglutinin                                     | hemagglutinin [Measles virus strain Moraten]  | AAF85673.1        |  |
| hemagglutinin                                     | hemagglutinin [Measles virus strain Rubeovax] | AAF85689.1        |  |
| hemagglutinin                                     | hemagglutinin [Measles virus]                 | AAF89824.1        |  |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]         | CAA91369.1        |  |
| hemagglutinin                                     | hemagglutinin [Measles virus]                 | BAJ23068.1        |  |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]         | BAB39848.1        |  |
| hemagglutinin                                     | hemagglutinin [Measles virus]                 | AAA50551.1        |  |
| hemagglutinin                                     | RecName: Full = Hemagglutinin glycoprotein    | P08362.1          |  |
| hemagglutinin                                     | hemagglutinin [Measles virus]                 | AAB63802.1        |  |

331

TABLE 15-continued

| MeV NCBI Accession Numbers (Amino Acid Sequences) |  |                          |
|---|--|--------------------------|
| Туре  | Virus Name   | GenBank Accession        |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA56650.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA56642.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA74936.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | BAH56665.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | ACC86105.1<br>AAF85697.1 |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin [Measles virus strain Edmonston-Zagreb]<br>hemagglutinin [Measles virus] | AAR89413.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA56653.1               |
| hemagglutinin                                     | RecName: Full = Hemagglutinin glycoprotein   | P35971.1                 |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94916.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAC03036.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus] Hemagglutinin [Measles virus]                            | AAF85681.1<br>CAB94927.1 |
| hemagglutinin<br>hemagglutinin                    | Hemagglutinin [Measles virus]  | CAB94927.1<br>CAB94925.1 |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | BAB39835.1               |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94931.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype A]   | AFO84712.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA56639.1               |
| hemagglutinin                                     | Hemagglutinin [Measles virus] hemagglutinin protein [Measles virus]                    | CAB94926.1<br>BAB39836.1 |
| hemagglutinin<br>hemagglutinin                    | Hemagglutinin [Measles virus]  | CAB94929.1               |
| hemagglutinin                                     | RecName: Full = Hemagglutinin glycoprotein   | P06830.1                 |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94928.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | BAB39837.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA74935.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43780.1               |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin [Measles virus] hemagglutinin protein [Measles virus]                    | BAA09952.1<br>CAB43815.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAF28390.1               |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94923.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43785.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | ABD34001.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43782.1               |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin protein [Measles virus]<br>hemagglutinin [Measles virus]                 | CAB43781.1<br>BAH22353.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAC35878.2               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | AAL86996.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | CAA76066.2               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA46428.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus] Hemagglutinin [Measles virus]                    | CAB43803.1<br>CAB94918.1 |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin [Measles virus]  | AAF72162.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAM70154.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43776.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D4]  | ACT78395.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D7]  | AAL02030.1               |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin protein [Measles virus] hemagglutinin protein [Measles virus]            | CAB43789.1<br>CAB43774.1 |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94920.1               |
| hemagglutinin                                     | Hemagglutinin [Measles virus]  | CAB94922.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | ABB59491.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | BAB39843.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus] hemagglutinin [Measles virus]                    | CAB43804.1               |
| hemagglutinin<br>hemagglutinin                    | Hemagglutinin [Measles virus]  | AAX52048.1<br>CAB94930.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA74526.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43814.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | ABB59493.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D4]  | AAL02019.1               |
| hemagglutinin<br>hemagglutinin                    | Hemagglutinin [Measles virus] hemagglutinin protein [Measles virus]                    | CAB94919.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype C2]  | AAL86997.1<br>AAL02017.1 |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43769.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43808.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | BAO97032.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43805.1               |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin protein [Measles virus] hemagglutinin [Measles virus]                    | CAB43777.1<br>AAL67793.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAL67793.1<br>AAF89816.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D4]  | AAL02020.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]  | CAB43786.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus strain<br>MVi/New Jersey.USA/45.05]               | AEP40452.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAA74531.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAB63800.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]  | AAO21711.1               |

TABLE 15-continued

| MeV NCBI Accession Numbers (Amino Acid Sequences) |   |                          |
|---|---|--------------------------|
| Туре  | Virus Name  | GenBank Accession        |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D8]   | ALE27189.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]   | CAB43810.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAF89817.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus genotype D6]   | AAL02022.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]   | CAB43800.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus genotype B3]                                     | AGA17219.1<br>CAB43770.1 |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin protein [Measles virus]<br>hemagglutinin protein [Measles virus strain  | AEP40444.1               |
| пениярыши   | MVi/Texas.USA/4.07]   | 1221 1011111             |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAX52047.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAB63794.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAB63796.1               |
| hemagglutinin<br>hemagglutinin                    | hemagglutinin [Measles virus] hemagglutinin [Measles virus]                           | AAA74528.1<br>AAB63774.1 |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAB63795.1               |
| hemagglutinin                                     | hemagglutinin [Measles virus]   | AAA74519.1               |
| hemagglutinin                                     | hemagglutinin protein [Measles virus]   | CAB43778.1               |
| fusion protein                                    | fusion protein [Measles virus strain Moraten]   | AAF85672.1               |
| fasion protein                                    | fusion protein [Measles virus]  | AAA56645.1<br>AAF85688.1 |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus strain Rubeovax] fusion protein [Measles virus]         | AAF85680.1               |
| fusion protein                                    | fusion protein [Measles virus]  | AEF30359.1               |
| fusion protein                                    | fusion protein [Measles virus]  | BAA09957.1               |
| fusion protein                                    | fusion protein [Measles virus]  | AAV84957.1               |
| fusion protein                                    | fusion protein [Measles virus MeV-eGFP_Edm-tag]                                       | AII16636.1               |
| fusion protein                                    | fusion protein [Measles virus]  | ABY58018.1               |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus] fusion protein [Measles virus]                         | BAA19838.1<br>AAA56641.1 |
| fusion protein                                    | F protein [Measles virus]   | ABK40529.1               |
| fusion protein                                    | fusion protein [Measles virus]  | AAA56652.1               |
| fusion protein                                    | fusion protein [Measles virus]  | ABY58017.1               |
| fusion protein                                    | fusion protein [Measles virus]  | ABB71645.1               |
| fusion protein                                    | fusion protein [Measles virus]  | NP_056922.1              |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus strain AIK-C]<br>fusion protein [Measles virus]         | AAF85664.1<br>BAB60865.1 |
| fusion protein                                    | fusion protein [Measles virus]  | BAA09950.1               |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40403.1               |
| 0.1   | MVi/New York.USA/26.09/3]   |                          |
| fusion protein                                    | fusion protein [Measles virus]  | AAA74934.1<br>CAB38075.1 |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus] fusion protein [Measles virus strain                   | AEP40443.1               |
| rasion protein                                    | MVi/Texas.USA/4.07]   | 1111 10115.1             |
| fusion protein                                    | fusion protein [Measles virus]  | AAF02695.1               |
| fusion protein                                    | fusion protein [Measles virus]  | AAF02696.1               |
| fusion protein                                    | fusion protein [Measles virus]  | AAT99301.1               |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus] fusion protein [Measles virus]                         | ABB71661.1<br>BAK08874.1 |
| fusion protein                                    | fusion protein [Measles virus]  | AAF02697.1               |
| fusion protein                                    | fusion protein [Measles virus genotype D4]  | AFY12704.1               |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40467.1               |
|   | MVi/California.USA/16.03]   |                          |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype D8]  | AHN07989.1<br>AAA46421.1 |
| fusion protein                                    | fusion protein [Measles virus] fusion protein [Measles virus]                         | AAA40421.1<br>AAA56638.1 |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40419.1               |
| •   | MVi/Virginia.USA/15.09]   |                          |
| fusion protein                                    | fusion protein [Measles virus genotype D8]  | ALE27200.1               |
| fusion protein                                    | fusion protein [Measles virus genotype D8]  | AFY12695.1               |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype D8] fusion protein [Measles virus genotype D8] | ALE27248.1<br>ALE27224.1 |
| fusion protein                                    | fusion protein [Measles virus]  | AAT99300.1               |
| fusion protein                                    | fusion protein [Measles virus]  | BAH96592.1               |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40459.1               |
|   | MVi/California.USA/8.04]  | 170010011                |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype D8] fusion protein [Measles virus]             | AIG94081.1<br>BAA09951.1 |
| fusion protein                                    | fusion protein [Measles virus genotype D8]  | ALE27194.1               |
| fusion protein                                    | fusion protein [Measles virus]  | BAA33871.1               |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40427.1               |
|   | MVi/Washington.USA/18.08/1]   |                          |
| fusion protein                                    | fusion protein [Measles virus]  | ABY21182.1               |
| fusion protein                                    | fusion protein [Measles virus genotype D8]  | ALE27284.1               |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus] fusion protein [Measles virus genotype D8]             | ACA09725.1<br>ALE27314.1 |
| fusion protein                                    | fusion protein [Measles virus genotype G3]  | AFY12712.1               |
| fusion protein                                    | fusion protein [Measles virus genotype D8]  | ALE27368.1               |
|   |   |                          |

TABLE 15-continued

| MeV NCBI Accession Numbers (Amino Acid Sequences) |   |                             |
|---|---|-----------------------------|
| Туре  | Virus Name  | GenBank Accession           |
| fusion protein                                    | RecName: Full = Fusion glycoprotein F0; Contains:   | P35973.1                    |
| rasion protein                                    | RecName: Full = Fusion glycoprotein F2; Contains:   | 133773.1                    |
| fusion protein                                    | RecName: Full = Fusion glycoprotein F1; Flags: Precursor fusion protein [Measles virus genotype H1] | AIG53713.1                  |
| rusion protein                                    | unnamed protein product [Measles virus]   | CAA34588.1                  |
| fusion protein                                    | fusion protein [Measles virus]  | CAA76888.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype B3.1] fusion protein [Measles virus]                         | AIY55563.1<br>ADO17330.1    |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]  | AIG53703.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype B3]  | AGA17208.1                  |
| fusion protein                                    | fusion protein [Measles virus]  | AAL29688.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53706.1<br>AIG53701.1    |
| fusion protein                                    | fusion protein [Measles virus genotype B3]  | ALE27092.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53714.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1] fusion protein [Measles virus genotype H1]               | AIG53694.1<br>AIG53668.1    |
| fusion protein                                    | fusion protein [Measles virus]  | ACC86094.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53670.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53707.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype B3]<br>fusion protein [Measles virus genotype H1]            | AGA17216.1<br>AIG53671.1    |
| fusion protein                                    | fusion protein [Measles virus strain  | AEP40451.1                  |
|   | MVi/New Jersey.USA/45.05]   |                             |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53684.1<br>AIG53688.1    |
| fusion protein                                    | fusion protein [Measles virus genotype 111]   | AGA17214.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53683.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53667.1<br>AIG53686.1    |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53685.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53681.1                  |
|   | unnamed protein product [Measles virus]   | CAA34589.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53678.1<br>AIG53710.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53710.1<br>AIG53669.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53664.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus]  | AAA50547.1<br>AIG53679.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53709.1<br>AIG53709.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53672.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53697.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus genotype H1]<br>fusion protein [Measles virus genotype H1]            | AIG53689.1<br>AIG53676.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53675.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53663.1                  |
| fusion protein<br>fusion protein                  | fusion protein [Measles virus] fusion protein [Measles virus]                                       | BAA19841.1<br>AAF02701.1    |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53680.1                  |
| fusion protein                                    | fusion protein [Measles virus genotype H1]  | AIG53674.1                  |
| C protein   | C protein [Measles virus strain Moraten] RecName: Full = Protein C                                  | AAF85670.1                  |
| C protein<br>C protein                            | C protein [Measles virus]   | P03424.1<br>ACN54404.1      |
| C protein   | C protein [Measles virus]   | ACN54412.1                  |
| C protein   | RecName: Full = Protein C   | P35977.1                    |
| C protein<br>C protein                            | C protein [Measles virus] C protein [Measles virus]   | AAF85678.1<br>ABD33998.1    |
| C protein   | unnamed protein product [Measles virus]   | CAA34586.1                  |
| C protein   | C protein [Measles virus]   | BAJ51786.1                  |
| C protein<br>C protein                            | C protein [Measles virus]<br>virulence factor [Measles virus]                                       | BAA33869.1<br>ABO69700.1    |
| C protein   | C protein [Measles virus]   | NP_056920.1                 |
| C protein   | C protein [Measles virus]   | ADO17333.1                  |
| C protein<br>C protein                            | C protein [Measles virus] C protein [Measles virus]   | ACC86082.1<br>B A A 33875 1 |
| C protein   | C protein [Measles virus]   | BAA33875.1<br>ABY21189.1    |
| C protein   | C protein [Measles virus]   | BAE98296.1                  |
| C protein   | C protein [Measles virus]   | ADU17782.1                  |
| C protein   | C protein [Measles virus strain<br>MVi/Virginia.USA/15.09]  | AEP40417.1                  |
| C protein   | C protein [Measles virus]   | ADU17814.1                  |
| C protein   | C protein [Measles virus]   | ADU17798.1                  |
| C protein<br>C protein                            | C protein [Measles virus genotype D4] C protein [Measles virus]                                     | AFY12700.1<br>ADU17784.1    |
| C protein   | C protein [Measles virus] C protein [Measles virus strain   | AEP40465.1                  |
| F   | MVi/California.USA/16.03]   | <u>.</u>                    |

337

TABLE 15-continued

|                        | MeV NCBI Accession Numbers (Amino Acid Seque                                    | nces)                    |
|------------------------|---|--------------------------|
| Туре                   | Virus Name  | GenBank Accession        |
| C protein              | C protein [Measles virus]   | ABB71643.1               |
| C protein              | C protein [Measles virus]   | AEI91027.1               |
| C protein              | C protein [Measles virus]   | ADU17874.1               |
| C protein              | C protein [Measles virus]   | ADU17903.1               |
| C protein<br>C protein | C protein [Measles virus] C protein [Measles virus]                             | CAA34579.1<br>ADU17790.1 |
| C protein              | C protein [Measles virus]   | ADU17800.1               |
| C protein              | C protein [Measles virus]   | ABB71667.1               |
| C protein              | unnamed protein product [Measles virus]   | CAA34572.1               |
| C protein              | C protein [Measles virus strain   | AEP40433.1               |
| 0                      | MVi/Arizona.USA/11.08/2]  | A DV 11 7020 1           |
| C protein<br>C protein | C protein [Measles virus] C protein [Measles virus]                             | ADU17830.1<br>ADU17947.1 |
| C protein              | C protein [Measles virus]   | ADU17818.1               |
| C protein              | C protein [Measles virus strain   | AEP40449.1               |
|                        | MVi/New Jersey.USA/45.05]   |                          |
| C protein              | C protein [Measles virus strain   | AEP40441.1               |
| 0                      | MVi/Texas.USA/4.07]   | A DI 11 70 CA 1          |
| C protein<br>C protein | C protein [Measles virus] C protein [Measles virus]                             | ADU17864.1<br>ADU17838.1 |
| C protein              | C protein [Measles virus]   | ADU17838.1<br>ADU17881.1 |
| C protein              | C protein [Measles virus strain   | AEP40425.1               |
| •                      | MVi/Washington.USA/18.08/1]   |                          |
| C protein              | C protein [Measles virus]   | ADU17927.1               |
| C protein              | C protein [Measles virus]   | ADU17953.1               |
| C protein              | C protein [Measles virus]   | ADU17889.1               |
| C protein              | C protein [Measles virus]   | ADU17963.1               |
| C protein              | C protein [Measles virus]   | ADU17893.1               |
| C protein              | C protein [Measles virus]   | ADU17820.1               |
| C protein              | C protein [Measles virus]   | ABB71651.1               |
| C protein<br>C protein | C protein [Measles virus] C protein [Measles virus]                             | ADU17786.1<br>ADU17862.1 |
| C protein              | C protein [Measles virus]   | ADU17802.1<br>ADU17923.1 |
| C protein              | C protein [Measles virus]   | ADU17959.1               |
| C protein              | C protein [Measles virus]   | ADU17951.1               |
| C protein              | C protein [Measles virus]   | ADU17916.1               |
| C protein              | C protein [Measles virus]   | ADU17957.1               |
| C protein              | C protein [Measles virus]   | ADU17925.1               |
| C protein              | C protein [Measles virus]   | ADU17901.1               |
| C protein              | C protein [Measles virus]   | ADU17887.1               |
| C protein              | C protein [Measles virus]   | ADU17832.1               |
| C protein              | C protein [Measles virus]   | ADU17891.1<br>ADU17961.1 |
| C protein<br>C protein | C protein [Measles virus] C protein [Measles virus]                             | ADU17901.1<br>ADU17872.1 |
| C protein              | C protein [Measles virus]   | ADU17929.1               |
| C protein              | C protein [Measles virus]   | ADU17908.1               |
| C protein              | C protein [Measles virus]   | ADU17910.1               |
| C protein              | C protein [Measles virus]   | ADU17921.1               |
| C protein              | C protein [Measles virus]   | ADU17824.1               |
| C protein              | C protein [Measles virus strain   | AEP40473.1               |
|                        | MVi/Pennsylvania.USA/20.09]   |                          |
| C protein              | C protein [Measles virus]   | ADU17828.1               |
| C protein              | C protein [Measles virus]   | ADU17812.1               |
| C protein              | C protein [Measles virus genotype D8]   | AFY12692.1               |
| C protein<br>C protein | nonstructural C protein [Measles virus] RecName: Full = Protein C               | ABA59559.1<br>Q00794.1   |
| C protein              | nonstructural C protein [Measles virus]   | ADO17934.1               |
| C protein              | nonstructural C protein [Measles virus]   | ACJ66773.1               |
| C protein              | C protein [Measles virus genotype G3]   | AFY12708.1               |
| C protein              | RecName: Full = Protein C   | P26035.1                 |
| C protein              | C protein [Measles virus]   | BAA84128.1               |
| nucleoprotein          | RecName: Full = Nucleoprotein; AltName:   | Q77M43.1                 |
|                        | Full = Nucleocapsid protein;  |                          |
|                        | Short = NP; Short = Protein N   | 1.1705.004               |
| nucleoprotein          | nucleocapsid protein [Measles virus strain Rubeovax]                            | AAF85683.1               |
| nucleoprotein          | RecName: Full = Nucleoprotein; AltName:   | Q89933.1                 |
|                        | Full = Nucleocapsid protein;<br>Short = NP: Short = Protein N                   |                          |
| nucleoprotein          | Short = NP; Short = Protein N nucleocapsid protein [Measles virus strain AIK-C] | AAF85659.1               |
| nucleoprotein          | nucleoprotein [Measles virus]   | ABI54102.1               |
| nucleoprotein          | nucleoprotein [Measles virus]   | AAA56643.1               |
| nucleoprotein          | nucleoprotein [Measles virus]   | AAC03050.1               |
| nucleoprotein          | nucleoprotein [Measles virus]   | AAA18990.1               |
| *                      | •   |                          |

TABLE 15-continued

|                                | MeV NCBI Accession Numbers (Amino Acid S                                | equences)                |
|--------------------------------|---|--------------------------|
| Туре                           | Virus Name  | GenBank Accession        |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA56640.1               |
| nucleoprotein                  | RecName: Full = Nucleoprotein; AltName: Full = Nucleocapsid protein;    | P35972.1                 |
|                                | Short = NP; Short = Protein N   |                          |
| nucleoprotein                  | RecName: Full=Nucleoprotein; AltName:                                   | P10050.1                 |
|                                | Full = Nucleocapsid protein;<br>Short = NP; Short = Protein N           |                          |
| nucleoprotein                  | N protein [Measles virus]   | BAB60956.1               |
| nucleoprotein                  | RecName: Full = Nucleoprotein; AltName:                                 | B1AAA7.1                 |
|                                | Full = Nucleocapsid protein;<br>Short = NP; Short = Protein N           |                          |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA18991.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46894.1<br>CAB46871.1 |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB46871.1<br>CAB46872.1 |
| nucleoprotein                  | nucleoprotein [Measles virus]   | ABU49606.1               |
| nucleoprotein                  | nucleocapsid protein [Measles virus]                                    | AAA75494.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB46883.1<br>CAB46892.1 |
| nucleoprotein                  | unnamed protein product [Measles virus]                                 | CAA34584.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA18997.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46863.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus]<br>nucleoprotein [Measles virus]          | AEF30352.1<br>ABI54103.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus]                                    | AAA46433.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46902.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB46873.1<br>CAB46906.1 |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA74547.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA74537.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46862.1               |
| nucleoprotein<br>nucleoprotein | nucleocapsid protein [Measles virus] nucleoprotein [Measles virus]      | BAA09961.1<br>AAO15875.1 |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAO15871.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46882.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus]<br>nucleoprotein [Measles virus]          | CAB60124.1<br>ABI54104.1 |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46869.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46880.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA74541.1<br>AEP40446.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus strain<br>MVi/New Jersey.USA/45.05] | AEF40440.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | ABI54110.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46903.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus]<br>nucleoprotein [Measles virus]          | CAB46899.1<br>CAB46901.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus]                                    | ABB71640.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB60113.1               |
| nucleoprotein                  | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB60114.1<br>CAB60116.1 |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus]   | CAB46895.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB60121.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | ABI54111.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB46889.1<br>CAB46898.1 |
| nucleoprotein                  | nucleoprotein [Measles virus genotype B3]                               | ALE27083.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB60118.1               |
| nucleoprotein<br>nucleoprotein | nucleocapsid protein [Measles virus] nucleoprotein [Measles virus]      | CAA34570.1<br>AAC29443.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus strain                              | AEP40422.1               |
| -                              | MVi/Washington.USA/18.08/1]   |                          |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAO15872.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleoprotein [Measles virus]             | CAB46874.1<br>AAA74550.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus]                                    | ABB71648.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46900.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleocapsid protein [Measles virus]      | BAH22440.1<br>AAA46432.1 |
| nucleoprotein                  | nucleocapsid protein [Measles virus]                                    | BAA33867.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA74539.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB60115.1               |
| nucleoprotein<br>nucleoprotein | nucleoprotein [Measles virus] nucleocapsid protein [Measles virus]      | CAB60123.1<br>ABB71664.1 |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB60125.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | AAA74546.1               |
| nucleoprotein                  | nucleoprotein [Measles virus]   | CAB46886.1               |

341

TABLE 15-continued

| MeV NCBI Accession Numbers (Amino Acid Sequences) |   |   |
|---|---|---|
| Туре  | Virus Name  | GenBank Accession                       |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | BAH22350.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | CAB46867.1                              |
| nucleoprotein                                     | nucleocapsid protein [Measles virus]                                    | BAA09954.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAO15873.1                              |
| nucleoprotein<br>nucleoprotein                    | nucleocapsid protein [Measles virus] nucleoprotein [Measles virus]      | AEP95735.1<br>AAL37726.1                |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAA74549.1                              |
| nucleoprotein                                     | RecName: Full = Nucleoprotein; AltName:                                 | P26030.1                                |
| r   | Full = Nucleocapsid protein;<br>Short = NP; Short = Protein N           |   |
| nucleoprotein                                     | nucleoprotein [Measles virus ETH55/99]                                  | AAK07777.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus genotype B3]                               | AGA17238.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AEF30351.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus genotype B3]                               | AGA17242.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus ETH54/98]<br>nucleoprotein [Measles virus] | AAK07776.1<br>AAA74548.1                |
| nucleoprotein<br>nucleoprotein                    | nucleoprotein [Measles virus]   | AAA19221.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAC03039.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAA19223.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus genotype B3]                               | AGA17241.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | CAB60122.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | CAC34599.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAC03042.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | CAC34604.1                              |
| nucleoprotein                                     | nucleoprotein [Measles virus]   | AAA74544.1                              |
| nucleoprotein                                     | nucleocapsid protein [Measles virus]                                    | NP_056918.1                             |
| V Protein   | RecName: Full = Non-structural protein V                                | Q9IC37.1                                |
| V Protein<br>V Protein                            | RecName: Full = Non-structural protein V V protein [Measles virus]      | Q9EMA9.1<br>ACN54411.1                  |
| V Protein   | V protein [Measles virus]   | ACN54403.1                              |
| V Protein   | V protein [Measles virus]   | AEP95742.1                              |
| V Protein   | V protein [Measles virus strain   | AEP40416.1                              |
|   | MVi/Virginia.USA/15.09]   | 111111111111111111111111111111111111111 |
| V Protein   | V protein [Measles virus]   | ADU17801.1                              |
| V Protein   | V protein [Measles virus]   | ADU17849.1                              |
| V Protein   | V protein [Measles virus]   | ABB71642.1                              |
| V Protein   | V protein [Measles virus genotype D8]                                   | AFY12693.1                              |
| V Protein   | V protein [Measles virus]   | YP_003873249.2                          |
| V Protein   | V protein [Measles virus strain<br>MVi/Arizona.USA/11.08/2]             | AEP40432.1                              |
| V Protein   | RecName: Full = Non-structural protein V                                | P26036.1                                |
| V Protein   | V protein [Measles virus strain   | AEP40464.1                              |
|   | MVi/California.USA/16.03]   |   |
| V Protein   | V protein [Measles virus strain   | AEP40456.1                              |
|   | MVi/California.USA/8.04]  |   |
| V Protein   | V protein [Measles virus]   | ABY21188.1                              |
| V Protein   | V protein [Measles virus strain   | AEP40424.1                              |
| TID . :   | MVi/Washington.USA/18.08/1]   | D 11106501 1                            |
| V Protein   | V protein [Measles virus]   | BAH96581.1                              |
| V Protein<br>V Protein                            | V protein [Measles virus] RecName: Full = Non-structural protein V      | ABB71666.1<br>P60168.1                  |
| V Protein   | V protein [Measles virus]   | BAH96589.1                              |
| V Protein   | V protein [Measles virus]   | ADU17954.1                              |
| V Protein   | V protein [Measles virus strain   | AEP40400.1                              |
| v 110tem  | MVi/New York.USA/26.09/3]   | ALI 40400.1                             |
| V Protein   | V protein [Measles virus]   | ABY21196.1                              |
| V Protein   | virulence factor [Measles virus]  | ABO69701.1                              |
| V Protein   | V protein [Measles virus]   | ABB71650.1                              |
| V Protein   | V protein [Measles virus]   | ACC86086.1                              |
| V Protein   | V protein [Measles virus genotype D4]                                   | AFY12702.1                              |
| V Protein   | V protein [Measles virus strain   | AEP40448.1                              |
|   | MVi/New Jersey.USA/45.05]   |   |
| V Protein   | V protein [Measles virus]   | BAE98295.1                              |
| V Protein   | V protein [Measles virus]   | ACC86083.1                              |
| V Protein   | V protein [Measles virus]   | ACU5139.1                               |
| V Protein   | V protein [Measles virus]   | ADO17334.1                              |
| V Protein   | V protein [Measles virus]   | ADU17930.1                              |
| V Protein   | V protein [Measles virus genotype G3]                                   | AFY12710.1                              |
| V Protein   | V protein [Measles virus strain   | AEP40472.1                              |
| V Dont-1-   | MVi/Pennsylvania.USA/20.09]   | A DI 11 7020 1                          |
| V Protein   | phosphoprotein [Measles virus]  | ADU17839.1                              |
| V Protein<br>V Protein                            | V protein [Measles virus] V protein [Measles virus]                     | ADU17894.1<br>ACN50010.1                |
| * 110mm   | · protein [wieasies vitus]  | ACN30010.1                              |

343

TABLE 15-continued

| MeV NCBI Accession Numbers (Amino Acid Sequences) |   |                   |  |
|---|---|-------------------|--|
| Туре  | Virus Name                              | GenBank Accession |  |
| V Protein   | V protein [Measles virus]               | ADU17892.1        |  |
|   | unnamed protein product [Measles virus] | CAA34585.1        |  |
| V Protein   | V protein [Measles virus]               | ABD33997.1        |  |

### TABLE 16

| Name                           | Sequence   | SEQ ID<br>NO: |
|--------------------------------|--|---------------|
|                                | Flagellin Nucleic Acid Sequences   |               |
| NT (5'<br>UTR, ORF,<br>3' UTR) | Flagellin Nucleic Acid Sequences  TCAAGCTTTTGGACCCTCGTACAGAAGCTAATACGACTCACTAT AGGGAAATAAGAGAAAAGAAGAAGAAGAAAAAAGAGAGCACATGGCACATGGCCACATGGCCACATGGCCACATGGCCACAAGTCATTAATACAAACAGCCTGTCGCTG TTGACCCAGAATAACCTGAACAAATCCCAGTCCGCACTGGCCA CTGCCTATCGACCAGTTGTCTTCCGGTCTGCTTACACCAGCGCGCAAAGACAAATCCCAGTTCCCGTATCAACAGCGCGCAAAGACAAATCCCAGTTCCCGTAACCGTTTACCG CGAACATCCAAAGGCTCGACACGACA | 51            |
|                                | CACCCGTACCCCGTGGTCTTTGAATAAAGTCTGAGTGGGCGG<br>C  |               |
| ORF                            | ATGGCACAAGTCATTAATACAAACAGCCTGTCGCTGTTGACCC  | 52            |

Sequence,

AGAATAACCTGAACAAATCCCAGTCCGCACTGGGCACTGCTAT CGAGCGTTTGTCTTCCGGTCTGCGTATCAACAGCGCGAAAGAC GATGCGGCAGGACAGGCGATTGCTAACCGTTTTACCGCGAACA  ${\tt TCAAAGGTCTGACTCAGGCTTCCCGTAACGCTAACGACGGTAT}$  $\tt CTCCATTGCGCAGACCACTGAAGGCGCGCTGAACGAAATCAAC$  $\verb|AACAACCTGCAGCGTGTGCGTGAACTGGCGGTTCAGTCTGCGA|$  $\tt ATGGTACTAACTCCCAGTCTGACCTCGACTCCATCCAGGCTGAA$  $\verb|ATCACCCAGCGCCTGAACGAAATCGACCGTGTATCCGGCCAGA||$  $\tt CTCAGTTCAACGGCGTGAAAGTCCTGGCGCAGGACAACACCCT$  ${\tt GACCATCCAGGTTGGTGCCAACGACGGTGAAACTATCGATATT}$ GATTTAAAAGAAATCAGCTCTAAAACACTGGGACTTGATAAGC  ${\tt TTAATGTCCAAGATGCCTACACCCCGAAAGAAACTGCTGTAAC}$ CGTTGATAAAACTACCTATAAAAATGGTACAGATCCTATTACA  ${\tt GCCCAGAGCAATACTGATATCCAAACTGCAATTGGCGGTGGTG}$  ${\tt CAACGGGGGTTACTGGGGCTGATATCAAATTTAAAGATGGTCA}$  $\tt ATACTATTTAGATGTTAAAGGCGGTGCTTCTGCTGGTGTTTATA$  ${\tt AAGCCACTTATGATGAAACTACAAAGAAAGTTAATATTGATAC}$ GACTGATAAAACTCCGTTGGCAACTGCGGAAGCTACAGCTATT CGGGGAACGGCCACTATAACCCACAACCAAATTGCTGAAGTAA

SEO ID

NO:

346

CAAAAGAGGGTTTGATACGACCACAGTTGCGGCTCAACTTGC
TGCAGCAGGGGTTACTGGCGCCGATAAGGACAATACTAGCCTT
GTAAAACTATCGTTTGAGGATAAAAACGGTAAGGTTATTGATG
GTGGCTATGCAGTGAAAATGGGCGACGATTTCTATGCCGCTAC
ATATGATGAGAAAACAGGTGCAATTACTGCTAAAACCACTACT
TATACAGATGGTACTGGCGTTGCTCAAACTGGAGCTGTGAAAT
TTGGTGGCGCAAATGGTAAATCTGAAGTTGTTACTGCTACCGAT
GGTAAGACTTACTTAGCAAGCGACCTTGACAAACATAACTTCA
GAACAGGCGGTGAGCTTAAAGAGGTTAATACAGATAAGACTG
AAAACCCACTGCAGAAAATTGATGCTGCCTTGGCACAGGTTGA
TACACTTCGTTCTGACCTGGGCGGTTCAGAACCGTTTCAACT
CCGCTATCACCAACCTGGGCAATACCGTAAATAACCTGTCTTCT
GCCCGTAGCCGTATCGAAGCGTTTCGACTAGCAACCGAAGTCT

 ${\tt CCAACATGTCTCGCGCGCAGATTCTGCAGCAGGCCGGTACCTC}\\ {\tt CGTTCTGGCGCAGGCGAACCAGGTTCCGCAAAACGTCCTCTCTT}\\$ 

345

Sequence

TACTGCGT

mRNA Sequence (assumes T100 tail)

Name

G\*GGGAAAUAAGAGAGAAAAGAAGAGGAAGAAAUAUAA GAGCCACCAUGGCACAAGUCAUUAAUACAAACAGCCUGUCGC UGUUGACCCAGAAUAACCUGAACAAAUCCCAGUCCGCACUGG GCACUGCUAUCGAGCGUUUGUCUUCCGGUCUGCGUAUCAACA GCGCGAAAGACGAUGCGGCAGGACAGGCGAUUGCUAACCGUU UUACCGCGAACAUCAAAGGUCUGACUCAGGCUUCCCGUAACG CUAACGACGGUAUCUCCAUUGCGCAGACCACUGAAGGCGCGC UGAACGAAAUCAACAACAACCUGCAGCGUGUGCGUGAACUGG CGGUUCAGUCUGCGAAUGGUACUAACUCCCAGUCUGACCUCG ACUCCAUCCAGGCUGAAAUCACCCAGCGCCUGAACGAAAUCG ACCGUGUAUCCGGCCAGACUCAGUUCAACGGCGUGAAAGUCC UGGCGCAGGACAACACCCUGACCAUCCAGGUUGGUGCCAACG ACGGUGAAACUAUCGAUAUUGAUUUAAAAGAAAUCAGCUCU AAAACACUGGGACUUGAUAAGCUUAAUGUCCAAGAUGCCUAC ACCCCGAAAGAACUGCUGUAACCGUUGAUAAAACUACCUAU AAAAAUGGUACAGAUCCUAUUACAGCCCAGAGCAAUACUGAU AUCCAAACUGCAAUUGGCGGUGGUGCAACGGGGGUUACUGG GGCUGAUAUCAAAUUUAAAGAUGGUCAAUACUAUUUAGAUG UUAAAGGCGGUGCUUCUGCUGGUGUUUAUAAAGCCACUUAU GAUGAAACUACAAAGAAAGUUAAUAUUGAUACGACUGAUAA AACUCCGUUGGCAACUGCGGAAGCUACAGCUAUUCGGGGAAC GGCCACUAUAACCCACAACCAAAUUGCUGAAGUAACAAAAGA GGGUGUUGAUACGACCACAGUUGCGGCUCAACUUGCUGCAGC AGGGGUUACUGGCGCCGAUAAGGACAAUACUAGCCUUGUAA AACUAUCGUUUGAGGAUAAAAACGGUAAGGUUAUUGAUGGU GGCUAUGCAGUGAAAAUGGGCGACGAUUUCUAUGCCGCUACA UAUGAUGAGAAAACAGGUGCAAUUACUGCUAAAACCACUAC UUAUACAGAUGGUACUGGCGUUGCUCAAACUGGAGCUGUGA AAUUUGGUGGCGCAAAUGGUAAAUCUGAAGUUGUUACUGCU ACCGAUGGUAAGACUUACUUAGCAAGCGACCUUGACAAACAU AACUUCAGAACAGGCGGUGAGCUUAAAGAGGUUAAUACAGA UAAGACUGAAAACCCACUGCAGAAAAUUGAUGCUGCCUUGGC ACAGGUUGAUACACUUCGUUCUGACCUGGGUGCGGUUCAGAA CCGUUUCAACUCCGCUAUCACCAACCUGGGCAAUACCGUAAA UAACCUGUCUUCUGCCCGUAGCCGUAUCGAAGAUUCCGACUA CGCAACCGAAGUCUCCAACAUGUCUCGCGCGCAGAUUCUGCA GCAGGCCGGUACCUCCGUUCUGGCGCAGGCGAACCAGGUUCC GCAAAACGUCCUCUUUACUGCGUUGAUAAUAGGCUGGAGC CUCGGUGGCCAUGCUUCUUGCCCCUUGGGCCUCCCCCCAGCC CCUCCUCCCUUCCUGCACCCGUACCCCCGUGGUCUUUGAAU 

Flagellin mRNA Sequences

NT (5' UTR, ORF, 3' UTR) 53

TABLE 16-continued

Name Sequence SEQ ID

UAAAACACUGGGACUUGAUAAGCUUAAUGUCCAAGAUGCCU

ACACGCGCAAACAAACUGGUUGAUAAGCUUAAUGUCCAAGAUGCCU

ACACCCGAAAGAAACUGCUGUAACCGUUGAUAAAACUACCU AUAAAAAUGGUACAGAUCCUAUUACAGCCCAGAGCAAUACUG AUAUCCAAACUGCAAUUGGCGGUGGUGCAACGGGGGUUACU GGGGCUGAUAUCAAAUUUAAAGAUGGUCAAUACUAUUUAGA UGUUAAAGGCGGUGCUUCUGCUGGUGUUUAUAAAGCCACUU AUGAUGAAACUACAAAGAAAGUUAAUAUUGAUACGACUGAU AAAACUCCGUUGGCAACUGCGGAAGCUACAGCUAUUCGGGGA ACGGCCACUAUAACCCACAACCAAAUUGCUGAAGUAACAAAA GAGGGUGUUGAUACGACCACAGUUGCGGCUCAACUUGCUGCA GCAGGGGUUACUGGCGCCGAUAAGGACAAUACUAGCCUUGUA AAACUAUCGUUUGAGGAUAAAAACGGUAAGGUUAUUGAUGG UGGCUAUGCAGUGAAAAUGGGCGACGAUUUCUAUGCCGCUAC AUAUGAUGAGAAAACAGGUGCAAUUACUGCUAAAACCACUA CUUAUACAGAUGGUACUGGCGUUGCUCAAACUGGAGCUGUG AAAUUUGGUGGCGCAAAUGGUAAAUCUGAAGUUGUUACUGC UACCGAUGGUAAGACUUACUUAGCAAGCGACCUUGACAAACA UAACUUCAGAACAGGCGGUGAGCUUAAAGAGGGUUAAUACAG AUAAGACUGAAAACCCACUGCAGAAAAUUGAUGCUGCCUUGG CACAGGUUGAUACACUUCGUUCUGACCUGGGUGCGGUUCAGA ACCGUUUCAACUCCGCUAUCACCAACCUGGGCAAUACCGUAA AUAACCUGUCUUCUGCCCGUAGCCGUAUCGAAGAUUCCGACU ACGCAACCGAAGUCUCCAACAUGUCUCGCGCGCAGAUUCUGC AGCAGGCCGGUACCUCCGUUCUGGCGCAGGCGAACCAGGUUC CGCAAAACGUCCUCUUUUACUGCGUUGAUAAUAGGCUGGAG CCUCGGUGGCCAUGCUUCUUGCCCCUUGGGCCUCCCCCCAGC CCCUCCUCCCUUCCUGCACCCGUACCCCGUGGUCUUUGAA UAAAGUCUGAGUGGGCGGC

ORF Sequence, AUGGCACAAGUCAUUAAUACAAACAGCCUGUCGCUGUUGACC CAGAAUAACCUGAACAAAUCCCAGUCCGCACUGGGCACUGCU AUCGAGCGUUUGUCUUCCGGUCUGCGUAUCAACAGCGCGAAA GACGAUGCGGCAGGACAGGCGAUUGCUAACCGUUUUACCGCG AACAUCAAAGGUCUGACUCAGGCUUCCCGUAACGCCUAACGAC GGUAUCUCCAUUGCGCAGACCACUGAAGGCGCGCUGAACGAA AUCAACAACAACCUGCAGCGUGUGCGUGAACUGGCGGUUCAG UCUGCGAAUGGUACUAACUCCCAGUCUGACCUCGACUCCAUC CAGGCUGAAAUCACCCAGCGCCUGAACGAAAUCGACCGUGUA UCCGGCCAGACUCAGUUCAACGGCGUGAAAGUCCUGGCGCAG GACAACACCCUGACCAUCCAGGUUGGUGCCAACGACGGUGAA ACUAUCGAUAUUGAUUUAAAAGAAAUCAGCUCUAAAACACU GGGACUUGAUAAGCUUAAUGUCCAAGAUGCCUACACCCCGAA AGAAACUGCUGUAACCGUUGAUAAAACUACCUAUAAAAAUG GUACAGAUCCUAUUACAGCCCAGAGCAAUACUGAUAUCCAAA CUGCAAUUGGCGGUGGUGCAACGGGGGUUACUGGGGCUGAU AUCAAAUUUAAAGAUGGUCAAUACUAUUUAGAUGUUAAAGG CGGUGCUUCUGCUGGUGUUUAUAAAGCCACUUAUGAUGAAA CUACAAAGAAAGUUAAUAUUGAUACGACUGAUAAAACUCCG UUGGCAACUGCGGAAGCUACAGCUAUUCGGGGAACGGCCACU AUAACCCACAACCAAAUUGCUGAAGUAACAAAAGAGGGUGU UGAUACGACCACAGUUGCGGCUCAACUUGCUGCAGCAGGGGU UACUGGCGCCGAUAAGGACAAUACUAGCCUUGUAAAACUAUC GUUUGAGGAUAAAAACGGUAAGGUUAUUGAUGGUGGCUAUG CAGUGAAAAUGGGCGACGAUUUCUAUGCCGCUACAUAUGAU GAGAAAACAGGUGCAAUUACUGCUAAAACCACUACUUAUACA GAUGGUACUGGCGUUGCUCAAACUGGAGCUGUGAAAUUUGG UGGCGCAAAUGGUAAAUCUGAAGUUGUUACUGCUACCGAUG GUAAGACUUACUUAGCAAGCGACCUUGACAAACAUAACUUCA GAACAGGCGGUGAGCUUAAAGAGGUUAAUACAGAUAAGACU GAAAACCCACUGCAGAAAAUUGAUGCUGCCUUGGCACAGGUU GAUACACUUCGUUCUGACCUGGGUGCGGUUCAGAACCGUUUC AACUCCGCUAUCACCAACCUGGGCAAUACCGUAAAUAACCUG UCUUCUGCCCGUAGCCGUAUCGAAGAUUCCGACUACGCAACC GAAGUCUCCAACAUGUCUCGCGCGCAGAUUCUGCAGCAGGCC GGUACCUCCGUUCUGGCGCAGGCGAACCAGGUUCCGCAAAAC GUCCUCUCUUUACUGCGU

mRNA Sequence (assumes T100 tail) 82

Name

TABLE 16-continued

| ! | Sequence                                   | SEQ ID<br>NO: |
|---|--|---------------|
|   | ACCGUGUAUCCGGCCAGACUCAGUUCAACGGCGUGAAAGUCC |               |
|   | UGGCGCAGGACAACACCCUGACCAUCCAGGUUGGUGCCAACG |               |
|   | ACGGUGAAACUAUCGAUAUUGAUUUAAAAGAAAUCAGCUCU  |               |
|   | AAAACACUGGGACUUGAUAAGCUUAAUGUCCAAGAUGCCUAC |               |
|   | ACCCCGAAAGAAACUGCUGUAACCGUUGAUAAAACUACCUAU |               |
|   | AAAAAUGGUACAGAUCCUAUUACAGCCCAGAGCAAUACUGAU |               |
|   | AUCCAAACUGCAAUUGGCGGUGGUGCAACGGGGGUUACUGG  |               |
|   | GGCUGAUAUCAAAUUUAAAGAUGGUCAAUACUAUUUAGAUG  |               |
|   | UUAAAGGCGGUGCUUCUGCUGGUGUUUAUAAAGCCACUUAU  |               |
|   | GAUGAAACUACAAAGAAAGUUAAUAUUGAUACGACUGAUAA  |               |
|   | AACUCCGUUGGCAACUGCGGAAGCUACAGCUAUUCGGGGAAC |               |
|   | GGCCACUAUAACCCACAACCAAAUUGCUGAAGUAACAAAAGA |               |
|   | GGGUGUUGAUACGACCACAGUUGCGGCUCAACUUGCUGCAGC |               |
|   | AGGGGUUACUGGCGCCGAUAAGGACAAUACUAGCCUUGUAA  |               |
|   | AACUAUCGUUUGAGGAUAAAAACGGUAAGGUUAUUGAUGGU  |               |
|   | GGCUAUGCAGUGAAAAUGGGCGACGAUUUCUAUGCCGCUACA |               |
|   | UAUGAUGAGAAAACAGGUGCAAUUACUGCUAAAACCACUAC  |               |
|   | UUAUACAGAUGGUACUGGCGUUGCUCAAACUGGAGCUGUGA  |               |
|   | AAUUUGGUGGCGCAAAUGGUAAAUCUGAAGUUGUUACUGCU  |               |
|   | ACCGAUGGUAAGACUUACUUAGCAAGCGACCUUGACAAACAU |               |
|   | AACUUCAGAACAGGCGGUGAGCUUAAAGAGGUUAAUACAGA  |               |
|   | UAAGACUGAAAACCCACUGCAGAAAAUUGAUGCUGCCUUGGC |               |
|   | ACAGGUUGAUACACUUCGUUCUGACCUGGGUGCGGUUCAGAA |               |
|   | CCGUUUCAACUCCGCUAUCACCAACCUGGGCAAUACCGUAAA |               |
|   | UAACCUGUCUUCUGCCCGUAGCCGUAUCGAAGAUUCCGACUA |               |
|   | CGCAACCGAAGUCUCCAACAUGUCUCGCGCGCAGAUUCUGCA |               |
|   | GCAGGCCGGUACCUCCGUUCUGGCGCAGGCGAACCAGGUUCC |               |
|   | GCAAAACGUCCUCUUUACUGCGUUGAUAAUAGGCUGGAGC   |               |
|   | CUCGGUGGCCAUGCUUCUUGCCCCUUGGGCCUCCCCCAGCC  |               |
|   | CCUCCUCCCUUCCUGCACCCGUACCCCCGUGGUCUUUGAAU  |               |
|   | AAAGUCUGAGUGGGCGGCAAAAAAAAAAAAAAAAAAAAA    |               |
|   | AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA    |               |
|   | AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA    |               |

TABLE 17

|  | Flagellin Amino Acid Sequences  |               |
|--|---|---------------|
| Name   | Sequence  | SEQ ID<br>NO: |
| ORF<br>Sequence,<br>AA   | MAQVINTNSLSLLTQNNLNKSQSALGTAIERLSSGLRINSAKDDAA GQAIANRFTANIKGLTQASRNANDGISIAQTTEGALNEINNNLQRV RELAVQSANGTNSQSDLDSIQAEITQRLNEIDRVSGQTQFNGVKVL AQDNTLTIQVGANDGETIDIDLKEISSKTLGLDKLNVQDAYTFKET AVTVDKTTYKNGTDPITAQSNTDIQTAIGGGATGVTGADIKFKDGQ YYLDVKGGASAGVYKATYDETTKKVNIDTTDKTPLATAEATAIRGT ATITHNQIAEVTKEGVDTTTVAQQLAAAGVTGADKDNTSLVKLSFE DKNGKVIDGGYAVKMGDDFYAATYDEKTGAITAKTTTYTDGTGVAQ TGAVKFGGANGKSEVVTATDGKTYLASDLDKHNFRTGGELKEVNTD KTENPLQKIDAALAQVDTLRSDLGAVQNRFNSAITNLGNTVNNLSS ARSRIEDSDYATEVSNMSRAQILQQAGTSVLAQANQVPQNVLSLLR  | 54            |
| Flagellin-<br>GS linker-<br>circumspor<br>ozoite<br>protein<br>(CSP) | MAQVINTNSLSLLTQNNLNKSQSALGTAIERLSSGLRINSAKDDAA GQAIANRFTANIKGLTQASRNANDGISIAQTTEGALNEINNNLQRV RELAVQSANSTNSQSDLDSIQAEITQRLNEIDRVSGQTQFNGVKVL AQDNTLTIQVGANDGETIDIDLKQINSQTLGLDTLNVQQKYKVSDT AATVTGYADTTIALDNSTFKASATGLGGTDQKIDGDLKFDDTTGKY YAKVTVTGGTGKDGYYEVSVDKTNGEVTLAGGATSPLTGGLPATAT EDVKNVQVANADLTEAKAALTAAGVTGTASVVKMSYTDNNGKTIDG GLAVKVGDDYYSATQNKDGSISINTTKYTADDGTSKTALNKLGGAD GKTEVVSIGGKTYAASKAEGHNFKAQPDLAEAAATTTENPLQKIDA ALAQVDTLRSDLGAVQNRFNSAITNLGNTVNNLTSARSRIEDSDYA TEVSNMSRAQILQQAGTSVLAQANQVPQNVLSLLRGGGGSGGGSM MAPDPNANPNANPNANPNANPNANPNANPNANPNANPNANP | 55            |
| Flagellin-<br><u>RPVT</u><br><u>linker</u> -                         | mmapdpnanpnanpnanpnanpnanpnanpnanpnanpnanp  | 56            |

# TABLE 17-continued

| Name                                     | Sequence   | SEQ ID<br>NO: |
|--|--|---------------|
| circumspor<br>ozoite<br>protein<br>(CSP) | TEWSPCSVTCGNGIQVRIKPGSANKPKDELDYENDIEKKICKMEKC SSVFNVVNSRPVTMAQVINTNSLSLLTQNNLNKSQSALGTAIERLS SGLRINSAKDDAAGQAIANRFTANIKGLTQASRNANDGISIAQTTE GALNEINNNLQRVRELAVQSANSTNSQSDLDSIQAEITQRLNEIDR VSGQTQFNGVKVLAQDNTLTIQVGANDGETIDIDLKQINSQTLGLD TLNVQQKYKVSDTAATVTGYADTTIALDNSTPKASATGLGGTDQKI DGDLKFDDTTGKYYAKVTVTGGTGKDGYYEVSVDKTNGEVTLAGGA TSPLTGGLPATATEDVKNVQVANADLTEAKAALTAAGVTGTASVVK MSYTDNNGKTIDGGLAVKVGDDYYSATQNKDGSISINTTKYTADDG |               |
|  | TSKTALNKLGGADGKTEVVSIGGKTYAASKAEGHNFKAQPDLAEAA<br>ATTTENPLQKIDAALAQVDTLRSDLGAVQNRPNSAITNLGNTVNNL<br>TSARSRIEDSDYATEVSNMSRAQILQQAGTSVLAQANQVPQNVLSL<br>LR   |               |

# TABLE 18

| Human Metapneumovirus Mutant Amino Acid Sequences |  |                  |  |
|---|--|------------------|--|
| Strain  | Sequence   | SEQ ID<br>NO:    |  |
| HMPV_SC_DSCAV1_4MMV                               | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAICKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLAFAVRELKDFVSKNLTRALNKNKCDIDDLKMAVSFSQFNRRFLNVV RQPSDNAGITPAISLDLMTDAELLARAVPNMPTSAGQIKLMLENRAMVRRKG FGILCGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFNVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN | :<br>:<br>:<br>: |  |
| HMPV_SC_DSTRIC_4MMV                               | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAICKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPMMPTSAGQIKLMLENRAMVRRGG FGILCGVYGSSVYYMVQLPIFGVIDTPCWUVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEHQMHVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |                  |  |
| HMPV_SC_DM_Krarup_T74LD185P                       | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIPDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDDPGVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN | ;<br>;<br>;      |  |
| HMPV_SC_TM_Krarup_T74LD185PD454N                  | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIPDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPENQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  | ;<br>;<br>;      |  |
| HMPV_SC_4M_Krarup_T74LS170LD185P                  | mswkvviifsllitpqhglkesyleescstitegylsvlrtgwytnvftle<br>vgbvenltcsdgpslikteldl <u>l</u> ksalrelktvsadqlareeqienp <u>gss</u><br>fvlgaialgvaaaaavtagvaiaktirlesevtainnalkktneavstign  | <u>!</u>         |  |

# TABLE 18-continued

|                                       |   | SEQ II |
|---------------------------------------|---|--------|
| Strain                                | Sequence  | NO:    |
|                                       | GVRVLATAVRELKDFVLKNLTRAINKNKCDIPDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN   |        |
| HMPV_SC_5M_Krarup_T74LS170LD185PD454N | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEGIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVLKNLTRAINKNKCDIPDLKMAVSFSQFNRRFLNVV ROFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPENQPQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN                        | 90     |
| HMPV_SC_DM_Krarup_E51PT74L            | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLP VGDVENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEGIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN                        | 91     |
| HMPV_SC_TM_Krarup_E51PT74LD454N       | $\label{eq:mswkvviifsllitpqhglkesyleescstitegylsvlrtgwytnvftlevgdvenltcsdgpslikteldl\underline{L}$ ksalrelktvsadqlareeqienpgsgs fylgaialgvaaaavtagvaiaktirlesevtainnalkktneavstlgn gvrvlatavrelkdfysknltrainknkcdiddlkmavsfsgpnrrflnvv rqfsdnagitpaisldlmtdaelaravpnmptsagqiklmlenramvrkg fgiligyygssviymvqlpifgvidtpcwivkaapscsekkgnyacllred qgwycqnagstvyypnekdcetrgdhyfcdtaaginvaeqskecninistt nypckvstgrhpismvalsplgalvacykgvscsigsnrvgiikqlnkgcs yitnqdadtvtidntvyqlskvegeqhvikgrpvsssfdpikfpenqpqva ldqvfeniensqalvdgsnrilssaekcntgfiiviiliavlgssmilvsi filikktkkptgappelsgvtnngfiphn | 92     |
| HMPV_SC_StabilizeAlpha_T74L           | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEGXECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN                        | 93     |
| HMPV_SC_StabilizeAlpha_V55L           | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDLENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEGKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN                        | 94     |
| HMPV_SC_StabilizeAlpha_S170L          | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVLKNLTRAINKNKCDIDDLKMAVSFSQPNRRFLMVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED   | 95     |

TABLE 18-continued

| Strain  | Sequence   | SEQ I |
|---|--|-------|
|   | QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |       |
| HMPV_SC_StabilizeAlpha_T174W                      | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLWRAINKNKCDIDDLKMAVSFSQFNRRFLMVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKRPTGAPPELSGVTNNGFIPHN |       |
| HMPV_SC_4M_Stabilize-<br>Alpha_V55LT74LS170LT174W | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDLENLTCSDGPSLIKTELDLLKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVLKNLWRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPMEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN   |       |
| HMPV_ProlineStab_E51P                             | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLP VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLMVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |       |
| HMPV_ProlineStab_D185P                            | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGATALGVAAAAAVTAGVATAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIPDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTI NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |       |
| HMPV_ProlineStab_D183P                            | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS PVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFYSKNLTRAINKNKCPIDDLKMAVSFSQFNRRFLINVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTI NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN |       |
| HMPV_ProlineStab_E131P                            | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENP <u>GSGS</u> FVLGAIALGVAAAAAVTAGVAIAKTIRL <u>P</u> SEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLINVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA  |       |

## TABLE 18-continued

| Strain                    | Sequence   | SEQ II<br>NO: |
|---------------------------|--|---------------|
|                           | LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI<br>FIIIKKTKKPTGAPPELSGVTNNGFIPHN   |               |
| HMPV_ProlineStab_D447P    | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLEED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFPPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |               |
| HMPV_TrimerRepulsionD454N | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLINVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGMYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPENQFQVA LDQVFENIENSQALVDQSNRILSSABKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN |               |
| HMPV_TrimerRepulsionE453N | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTTRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQFNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLEED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPQDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |               |
| HMPV_StabilizeAlphaF196W  | MSWKVVIIFSLLITPQHGLKESYLEESCSTITEGYLSVLRTGWYTNVFTLE VGDVENLTCSDGPSLIKTELDLTKSALRELKTVSADQLAREEQIENPGSGS FVLGAIALGVAAAAAVTAGVAIAKTIRLESEVTAINNALKKTNEAVSTLGN GVRVLATAVRELKDFVSKNLTRAINKNKCDIDDLKMAVSFSQWNRRFLNVV RQFSDNAGITPAISLDLMTDAELARAVPNMPTSAGQIKLMLENRAMVRRKG FGILIGVYGSSVIYMVQLPIFGVIDTPCWIVKAAPSCSEKKGNYACLLRED QGWYCQNAGSTVYYPNEKDCETRGDHVFCDTAAGINVAEQSKECNINISTT NYPCKVSTGRHPISMVALSPLGALVACYKGVSCSIGSNRVGIIKQLNKGCS YITNQDADTVTIDNTVYQLSKVEGEQHVIKGRPVSSSFDPIKFPEDQFQVA LDQVFENIENSQALVDQSNRILSSAEKGNTGFIIVIILIAVLGSSMILVSI FIIIKKTKKPTGAPPELSGVTNNGFIPHN  |               |

# TABLE 19

| Strain              | Nucleic Acid Sequence  | SEQ ID<br>NO: |
|---------------------|--|---------------|
| Human Meta          | apneumovirus Mutant Nucleic Acid Sequences   |               |
| HMPV_SC_DSCAV1_4MMV | ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAAGAGTTACCTGGAAGAAGT CCTGCAGCACCACCACCAGAGGGCTACCTGTCTGTGTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGC GACGTCGAGAATCTGACCAAGAGGCGCCCTGAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGA ACTCAAGACCTGACTGCCGATCAGCTGGCCAGAGAGG CCATTGCTCTTGGAGTGGCTGCTGAGCTGTTACAG CAGGCGTGGCCATCTGCAGACCTGTACAG AAGTCACGCATCACACACCCCTGAAGAAGACAAACG AGGCGTCAGCACTCGCAACTAGACTTAGACTGGC CTTTGCCGTCGCGACTGAAGACACTCGGTTACAGACC | 106           |

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGTGTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCAACGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

HMPV\_SC\_DSTRIC\_4MMV

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT  $\tt CCTGCAGCACCATCACAGAGGGGCTACCTGTCTGTGCTGAG$ AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA  ${\tt TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG}$ AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG  ${\tt CAGGCGTGGCCATCTGCAAGACCATCAGACTGGAAAGCG}$ AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC  $\tt CTGACACGGGCCATTAACAAGAACAAGTGCGACATCGAC$ GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGTGTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGCACCAGTGGCATGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

HMPV\_SC\_DM\_Krarup\_T74LD185P

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA
CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT
CCTGCAGCACCATCACAGAGGGCTACCTGTGTGTGAG
AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC
GACGTCGAGAATCTGACATGCTCTGATGGCCCTGA
TCAAGACCGAGCTGGATCTCCTCAAGAGCGCCCTGAGAGA
ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGAA

107

TABLE 19-continued

Strain Nucleic Acid Sequence NO:

CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA  $\tt GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC$ ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCCCTGA CCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCGG TTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGAA TCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTGA GCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGGC CAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGAC GGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCAG CGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATCG ACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTAG CGAGAAGAAGGCCAATTACGCCTGCCTGAGAGAGAGA CCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTAC TACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCAC GTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAGC AGAGCAAAGAGTGCAACATCAACATCAGCACCACCAACT ATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC

109

HMPV\_SC\_TM\_Krarup\_T74LD185PD454N

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT  $\tt CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG$ AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG  $\tt CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC$ AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA  $\tt GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC$ ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCCCTGA CCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCGG TTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGAA TCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTGA GCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGGC CAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGAC GGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCAG CGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATCG ACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTAG CGAGAAGAAGGCCAATTACGCCTGCCTGCTGAGAGAGAA CCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTAC TACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCAC GTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAGC AGAGCAAAGAGTGCAACATCAACATCAGCACCACCAACT ATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGAACCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC

TABLE 19-continued

SEO ID Strain Nucleic Acid Sequence NO: HMPV SC 4M Krarup T74LS170LD185P ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGAAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG  $\tt CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC$ AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGCTTAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCCCTGA CCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCGG TTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGAA TCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTGA GCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGGC CAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGAC GGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCAG CGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATCG ACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTAG CCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTAC TACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCAC GTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAGC AGAGCAAAGAGTGCAACATCAACATCAGCACCACCAACT ATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT  ${\tt CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC}$ CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC  ${\tt AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG}$ GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC  ${\tt TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG}$ AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC HMPV SC 5M Krarup T74LS170LD185PD454N ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGCTTAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCCCTGA CCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCGG TTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGAA TCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTGA GCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGGC CAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGAC GGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCAG CGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATCG ACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTAG CCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTAC TACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCAC GTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAGC AGAGCAAAGAGTGCAACATCAACATCAGCACCACCAACT ATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA

AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT
CATCAACCAGCTGAACAAGGGCTGCAGCTACCATCACCAAC
CAGGACGCCGATACCGTGATCACCATCGACAACAACAACAGGTGATCAAG
GCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG
GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC
TGAGAACCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG

TABLE 19-continued

SEO ID Nucleic Acid Sequence Strain NO: AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA HMPV SC DM Krarup E51PT74L CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGCCTGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCGACG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAGGGCAATTACGCCTGCCTGAGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA  $\tt CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG$ CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC  ${\tt TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG}$ AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC HMPV SC TM Krarup E51PT74LD454N ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGCCTGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCGACG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAGGGCAATTACGCCTGCCTGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA  $\tt CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA$  $\tt CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG$ CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC

TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA
AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGCAT
CATCAAGCAGCTGAACAAGAGGCTTGCACTAC
CAGGACGCCGATACCAGAC
AGCTGAGCAAGGCTGACCAACACCGTGATCA
AGCTGAGCAAGGTGGAAGGCGAACACCGTGATCAAGG
GCAGACCTGTGTCCAGCAGCTTCGACCAGGTGTTCCAG
AACATCGAGAATTCCCAGGCTCTGGTGGACCAGGTGTCCAACA
GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT
CATCGTGATCATCTGATCGCCTGGCAGCAGCTCCATG
ATCCTGGTCCATCTCTCATCATTATCAAGAAGACCAAGA
AGCCCACCGGCGTCCTCCAGAACTGAGCGAGTGACCAA
CAATGGCTTCATCCTCACAAC

HMPV\_SC\_StabilizeAlpha\_T74L

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCGACG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAAGGCCAATTACGCCTGCCTGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC  ${\tt AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG}$ GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC

HMPV SC StabilizeAlpha V55L

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACCTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGCCATTAACAAGAACAAGTGCGACATCGAC GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG  ${\tt GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG}$ ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA

114

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC  ${\tt CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG}$ AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA

HMPV\_SC\_StabilizeAlpha\_S170L

CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG  ${\tt CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG}$ AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGCTTAAGAAC  $\tt CTGACACGGGCCATTAACAAGAACAAGTGCGACATCGAC$ GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG  ${\tt TAGCGAGAAGAAGGGCAATTACGCCTGCTGAGAGA}$ GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC  ${\tt CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG}$ AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

HMPV\_SC\_StabilizeAlpha\_T174W

116

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA

HMPV\_SC\_4M\_Stabilize-Alpha\_V55LT74LS170LT174W

CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT  $\tt CCTGCAGCACCATCACAGAGGGGCTACCTGTCTGTGCTGAG$ AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACCTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGCTCAAGAGCGCCCTGAGAGA ACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGAA CAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGAG CCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAGC AGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGCTTAAGAACC  ${\tt TGTGGCGGGCCATTAACAAGAACAAGTGCGACATCGACG}$ ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG  ${\tt CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA}$ CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAAGGCCAATTACGCCTGCCTGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC

HMPV\_ProlineStab\_E51P

 118

TABLE 19-continued

Strain Nucleic Acid Sequence NO:

ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG  ${\tt AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC}$ CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGGCCATTAACAAGAACAAGTGCGACATCGAC GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT  $\tt CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC$ AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

HMPV\_ProlineStab\_D185P

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT  $\tt CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG$ AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG  ${\tt AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC}$ CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGGCCATTAACAAGAACAAGTGCGACATCCCTG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACCCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAGGGCAATTACGCCTGCCTGCTGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC

120

375

TABLE 19-continued

SEO ID Nucleic Acid Sequence Strain NO: HMPV ProlineStab D183P ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGGCCATTAACAAGAACAAGTGCCCTATCGACG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACCCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAGGGCAATTACGCCTGCCTGCTGAGAGAGG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT  ${\tt CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC}$ CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC  ${\tt AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG}$ GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC  ${\tt TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG}$ AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC HMPV ProlineStab E131P ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA  ${\tt CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT}$ CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA  ${\tt TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG}$ AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGCCTAGCGA AGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACGA GGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGCC ACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAACC TGACACGGGCCATTAACAAGAACAAGTGCGACATCGACG ACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGCG GTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGGA ATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCTG AGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCGG CCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCGA CGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGCA GCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGATC GACACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTGTA GCGAGAAGAGGGCAATTACGCCTGCCTGAGAGAGAG ACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTGTA CTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGACCA CGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCGAG CAGAGCAAAGAGTGCAACATCAACATCAGCACCACCAAC TATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTCTAT GGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTTATA  ${\tt AGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGGGCAT}$ CATCAAGCAGCTGAACAAGGGCTGCAGCTACATCACCAAC

CAGGACGCCGATACCGTGACCATCGACAACACCGTGTATC  ${\tt AGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATCAAGG}$ GCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGTTCCC TGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTTCGAG

TABLE 19-continued

SEO ID Nucleic Acid Sequence Strain NO: AACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCCAACA GAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGCTTCAT CATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTCCATG ATCCTGGTGTCCATCTTCATCATTATCAAGAAGACCAAGA AGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTGACCAA CAATGGCTTCATCCCTCACAAC HMPV ProlineStab D447P ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA 123 CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGGCCATTAACAAGAACAAGTGCGACATCGAC GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA  ${\tt TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG}$ TAGCGAGAAGAAGGGCAATTACGCCTGCCTGCTGAGAGA GGACCAAGGCTGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC  ${\tt CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG}$ AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC  ${\tt TATGGTGGCTCTGTCTCTCTGGGAGCCCTGGTGGCTTGTT}$ ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCCCACCTATCAAGT  ${\tt TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT}$ CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC HMPV TrimerRepulsionD454N ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGGCCATTAACAAGAACAAGTGCGACATCGAC GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG  ${\tt TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC}$  ${\tt CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG}$ AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA

ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT
ATAAGGGCGTGTCCTTATAGCATCGGCAGCAACAGAGTGG
GCATCATCAAGCAGCTGAACAAGAGGTGC
CAACCAGGACCGCGATACCGTGACCATCACACCAC
CAACCAGGACGCCGATACCGTGACCATCGACCACACCGTGATC
AAGGGCAGACCTGTGTCCAGCAGCTTCGACCAGGTGTT
TCCCTGAGAACCAGTTCCAAGCCTTGGACCAGGTGTC
CGAGAACCATCCAAGAATTCCCAGGCTCTGGTCGACCAGGTCC
AACAGAATCCTGTCTACCGCCCAGAAGGGAAACACCGGC
TTCATCATCGTGATCATCCTGATCGCCTGGGCAGCTC
CATGATCCTGGTGTCCATCTCATCATTATCAAGAAGACC
AAGAAGCCCACCGCGCTTCCTCCAGAACTGAGCGGAGTG
ACCAACAATGGCTTCATCCTCACAC

HMPV TrimerRepulsionE453N

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGCCATTAACAAGAACAAGTGCGACATCGAC  ${\tt GACCTGAAGATGGCCGTGTCCTTTAGCCAGTTCAACCGGC}$ GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA  ${\tt TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG}$ TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG  ${\tt TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC}$ AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT  ${\tt TCCCTCAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT}$ CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

HMPV StabilizeAlphaF196W

ATGAGCTGGAAGGTGGTCATCATCTTCAGCCTGCTGATCA CACCTCAGCACGGCCTGAAAGAGAGCTACCTGGAAGAGT CCTGCAGCACCATCACAGAGGGCTACCTGTCTGTGCTGAG AACCGGCTGGTACACCAACGTGTTCACACTGGAAGTGGGC GACGTCGAGAATCTGACATGCTCTGATGGCCCTAGCCTGA TCAAGACCGAGCTGGATCTGACCAAGAGCGCCCTGAGAG AACTCAAGACCGTGTCTGCCGATCAGCTGGCCAGAGAGGA ACAGATCGAGAATCCTGGCAGCGGCAGCTTTGTGCTGGGA GCCATTGCTCTTGGAGTGGCTGCTGCTGCAGCTGTTACAG CAGGCGTGGCCATCGCTAAGACCATCAGACTGGAAAGCG AAGTGACCGCCATCAACAACGCCCTGAAGAAGACAAACG AGGCCGTCAGCACACTCGGCAATGGCGTTAGAGTGCTGGC CACAGCCGTGCGCGAGCTGAAGGACTTCGTGTCCAAGAAC CTGACACGGCCATTAACAAGAACAAGTGCGACATCGAC GACCTGAAGATGGCCGTGTCCTTTAGCCAGTGGAACCGGC GGTTTCTGAACGTCGTGCGGCAGTTTAGCGACAACGCCGG AATCACACCAGCCATCAGCCTGGACCTGATGACAGATGCT GAGCTGGCTAGAGCCGTGCCTAACATGCCTACATCTGCCG GCCAGATCAAGCTGATGCTCGAGAATAGAGCCATGGTCCG ACGGAAAGGCTTCGGCATTCTGATTGGCGTGTACGGCAGC AGCGTGATCTATATGGTGCAGCTGCCTATCTTCGGCGTGA 125

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

TCGACACCCTGCTGGATTGTGAAGGCCGCTCCTAGCTG TAGCGAGAAGAAGGGCAATTACGCCTGCCTGAGAGA GGACCAAGGCTGGTATTGTCAGAACGCCGGCAGCACCGTG TACTACCCTAACGAGAAGGACTGCGAGACAAGAGGCGAC  ${\tt CACGTGTTCTGTGATACCGCCGCTGGAATCAATGTGGCCG}$ AGCAGAGCAAAGAGTGCAACATCAACATCAGCACCACCA ACTATCCCTGCAAGGTGTCCACCGGCAGGCACCCTATTTC TATGGTGGCTCTGTCTCCTCTGGGAGCCCTGGTGGCTTGTT ATAAGGGCGTGTCCTGTAGCATCGGCAGCAACAGAGTGG GCATCATCAAGCAGCTGAACAAGGGCTGCAGCTACATCAC CAACCAGGACGCCGATACCGTGACCATCGACAACACCGTG TATCAGCTGAGCAAGGTGGAAGGCGAACAGCACGTGATC AAGGGCAGACCTGTGTCCAGCAGCTTCGACCCTATCAAGT TCCCTGAGGATCAGTTCCAGGTGGCCCTGGACCAGGTGTT CGAGAACATCGAGAATTCCCAGGCTCTGGTGGACCAGTCC AACAGAATCCTGTCTAGCGCCGAGAAGGGAAACACCGGC TTCATCATCGTGATCATCCTGATCGCCGTGCTGGGCAGCTC CATGATCCTGGTGTCCATCTTCATCATTATCAAGAAGACC AAGAAGCCCACCGGCGCTCCTCCAGAACTGAGCGGAGTG ACCAACAATGGCTTCATCCCTCACAAC

### Human Metapneumovirus mRNA Sequences

HMPV\_SC\_DSCAV1\_4MMV

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCUGCAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCUUUGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCCUGAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GUGUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCAACGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_SC\_DSURIC\_4MMV

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU
CACACCUCAGCACGGCCUGAAAGAAGACCUACCUGGAAGA
GUCCUGCAGCACAUCACAGAGGGCUACCUGUCUGUGCU
GAGAACCGGCUGGUACACACGUGUUCACACUGGAAGA
GGCGACGUCGAGAAUCUGACAUGCUCUGAUGGACCUAG
CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU
GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCGCAGCUGUG
UGCUGGGAGCAUUGGCUUUGGAGCGCAGCUGUG
GCUGGUACAGACGCUUUGGAGUGCUGCAGCGCACGCGCACUUUG
GCUGGUACAGACCGUUGCCAGAGACCGCAUCAGA
CCUGUACAGACGCGUGGCCAUCACAAGACCCAUCAGA
CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG

127

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GUGUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGCACCAGUGGCAUGUGGCCCUGGACCAGGUGUUCGA GAACATICGAGAATIITCCCAGGCTICTIGGTIGGACCAGTICCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCUU CAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU  ${\tt GACCAACAAUGGCUUCAUCCCUCACAAC}$ 

HMPV\_SC\_DM\_Krarup\_U74LD185P

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCCCUGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

TABLE 19-continued

Strain Nucleic Acid Sequence NO:

GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCCCUGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGCC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGAACCAGUUCCAGGUGGCCCUGGACCAGGUGUUCGA GAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCUU CAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU GACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_SC\_4M\_Krarup\_U74LS170LD185P

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGCUUAAGAACCUGACACGGGCCAUUAACAAGAACAA GUGCGACAUCCCUGACCUGAAGAUGGCCGUGUCCUUUAG CCAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACCACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG  $\tt CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG$ CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG

#### TABLE 19-continued

SEO ID Nucleic Acid Sequence Strain NO: AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC HMPV SC 5M Krarup U74LS170LD185PD454N AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU 132 CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGCUUAAGAACCUGACACGGGCCAUUAACAAGAACAA GUGCGACAUCCCUGACCUGAAGAUGGCCGUGUCCUUUAG CCAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGAGAGAGAGCCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGAACCAGUUCCAGGUGGCCCUGGACCAGGUGUUCGA GAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCUU CAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU GACCAACAAUGGCUUCAUCCCUCACAAC HMPV SC DM Krarup E51PU74L AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGCCUGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACCCCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG

AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA
AGGUGUCCACCGGCAGGCACCUAUUUUCUAUGGUGGCUC
UGUCUCCUCUGGGAGCCCUGUUGUUAUAUGGUGCUC
UGUCUCCUCUGGAGCCCUGGUGGCUUGUUAUAAGGGC
GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC
AAGCAGCUGAACAAGGGCUGCAGCUACAACACAG
GACGCCGAUACCGUGACCAUCGACACACCGUGUAUCAG
CUGAGCAAGGUGGACGAACAGCACGUGAUCAAGGG
CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC
UGAGGAUCAGUUCCAGGUUCGACCCUAUCAAGUUCCA
ACAGAUCCAGAAUUCCCAGGCUCUGGACCAGUGUUCG
AGAACAUCGAGAAUUCCCAGGCUCUGGACCAGGUCCA
ACAGAAUCCUGUCUAGCCCGAGAAGGGAAACACCGGCU
UCAUCAUCGUGAUCCUCAUCAUAUCAAGAAGA
CCAAGAAGCCCACCGGCGCUCUCCAGAACUGAGCGGAG
UGACCAACAAUGGCUUCAUCACAAC

HMPV\_SC\_UM\_Krarup\_E51PU74LD454N

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGCCUGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGCCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGAACCAGUUCCAGGUGGCCCUGGACCAGGUGUUCGA GAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGGAAACACCGGCUU CAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU GACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_SC\_SUabilizeAlpha\_U74L

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACCCCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC

134

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA  $\tt UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC$ AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU

 ${\tt HMPV\_SC\_SUabilizeAlpha\_V55L}$ 

CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACCUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA  $\tt CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG$ AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_SC\_SUabilizeAlpha\_S170L

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU
CACACCUCAGCACGGCCUGAAAGAAGACUACCUGGAAGA
GUCCUGCAGCACAUCACAGAGGGCUACCUGUCUGUGCU
GAGAACCGGCUGGUACCAACGUGUUCACACUGGCAGG
GGGCGACGUCGAGAAUCUGACCAACGUCUUGAUCAGCCCUAG
CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU
GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGCCAG
AGAGGAACAGAUCGAGAAUCCUGGCAGCAGCUUUG
UGCUGGGAGGACAUUGCCUGCCA

136

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGCUUAAGAACCUGACACGGGCCAUUAACAAGAACAA GUGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAG CCAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_SC\_SUabilizeAlpha\_U174W

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGUGGCGGGCCAUUAACAAGAACAA GUGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAG CCAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACCACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

TABLE 19-continued

SEQ ID Nucleic Acid Sequence Strain NO: HMPV SC 4M SUabilize-AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU 139 Alpha\_V55LU74LS170LU174W CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACCUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGCUCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGCUUAAGAACCUGUGGCGGGCCAUUAACAAGAACAA GUGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAG CCAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC HMPV\_ProlineSUab\_E51P AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU 140 CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGCCUGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGACGC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG

CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG

TABLE 19-continued

SEO ID Nucleic Acid Sequence Strain NO: CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCGUGGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC HMPV ProlineSUab D185P AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU 141 CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACCCCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCCCUGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGCAGUUU AGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGAGCCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGGAAACACCGGCU UCAUCAUCGUGAUCGUGGUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC HMPV ProlineSUab D183P AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU 142 CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACCCCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCCCUAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA  $\tt UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC$ AAUUACGCCUGCCUGAGAGAGAGCCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCGUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_ProlineSUab\_E131P

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA  $\tt CUGCCUAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG$ AAGACAAACGAGGCCGUCAGCACACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCGUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_ProlineSUab\_D447P

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUUU

143

TABLE 19-continued

SEQ ID Strain Nucleic Acid Sequence NO:

AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACCCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCCCACCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCCUGAUCGCCGUGCUGGGCAGCU CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA CCAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAG UGACCAACAAUGGCUUCAUCCCUCACAAC

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU

HMPV\_UrimerRepulsionD454N

CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGAACCAGUUCCAGGUGGCCCUGGACCAGGUGUUCGA GAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCUU CAUCAUCGUGAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU GACCAACAAUGGCUUCAUCCCUCACAAC

HMPV\_UrimerRepulsionE453N

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU
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GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU
GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU
GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG
CCUGAUCAAGACCGAGCUGGAUCUGACAGAGGGCCCU
GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG

145

TABLE 19-continued

Strain Nucleic Acid Sequence NO:

AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACUCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUUCAACCGGCGUUUCUGAACGUCGUGCGGCAGUUU AGCGACAACGCCGGAAUCACCAGCCAUCAGCCUGGAC CUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAAC AUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGAG AAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUCU GAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUGC AGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGGA UUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGGC AAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UCAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCGA GAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCAA CAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCUU CAUCAUCGUGAUCAUCCUGAUCGCCGUGCUGGGCAGCUC CAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGAC CAAGAAGCCCACCGGCGCUCCUCCAGAACUGAGCGGAGU GACCAACAAUGGCUUCAUCCCUCACAAC

HMPV SUabilizeAlphaF196W

AUGAGCUGGAAGGUGGUCAUCAUCUUCAGCCUGCUGAU CACACCUCAGCACGGCCUGAAAGAGAGCUACCUGGAAGA GUCCUGCAGCACCAUCACAGAGGGCUACCUGUCUGUGCU GAGAACCGGCUGGUACACCAACGUGUUCACACUGGAAGU GGGCGACGUCGAGAAUCUGACAUGCUCUGAUGGCCCUAG CCUGAUCAAGACCGAGCUGGAUCUGACCAAGAGCGCCCU GAGAGAACUCAAGACCGUGUCUGCCGAUCAGCUGGCCAG AGAGGAACAGAUCGAGAAUCCUGGCAGCGGCAGCUUUG UGCUGGGAGCCAUUGCUCUUGGAGUGGCUGCUGCA GCUGUUACAGCAGGCGUGGCCAUCGCUAAGACCAUCAGA CUGGAAAGCGAAGUGACCGCCAUCAACAACGCCCUGAAG AAGACAAACGAGGCCGUCAGCACCCCGGCAAUGGCGUU AGAGUGCUGGCCACAGCCGUGCGCGAGCUGAAGGACUUC GUGUCCAAGAACCUGACACGGGCCAUUAACAAGAACAAG UGCGACAUCGACGACCUGAAGAUGGCCGUGUCCUUUAGC CAGUGGAACCGGCGGUUUCUGAACGUCGUGCGGCAGUU UAGCGACAACGCCGGAAUCACACCAGCCAUCAGCCUGGA CCUGAUGACAGAUGCUGAGCUGGCUAGAGCCGUGCCUAA CAUGCCUACAUCUGCCGGCCAGAUCAAGCUGAUGCUCGA GAAUAGAGCCAUGGUCCGACGGAAAGGCUUCGGCAUUC UGAUUGGCGUGUACGGCAGCAGCGUGAUCUAUAUGGUG CAGCUGCCUAUCUUCGGCGUGAUCGACACACCCUGCUGG AUUGUGAAGGCCGCUCCUAGCUGUAGCGAGAAGAAGGG CAAUUACGCCUGCCUGCUGAGAGAGGACCAAGGCUGGUA UUGUCAGAACGCCGGCAGCACCGUGUACUACCCUAACGA GAAGGACUGCGAGACAAGAGGCGACCACGUGUUCUGUG AUACCGCCGCUGGAAUCAAUGUGGCCGAGCAGAGCAAAG AGUGCAACAUCAACAUCAGCACCACCAACUAUCCCUGCA AGGUGUCCACCGGCAGGCACCCUAUUUCUAUGGUGGCUC UGUCUCCUCUGGGAGCCCUGGUGGCUUGUUAUAAGGGC GUGUCCUGUAGCAUCGGCAGCAACAGAGUGGGCAUCAUC AAGCAGCUGAACAAGGGCUGCAGCUACAUCACCAACCAG GACGCCGAUACCGUGACCAUCGACAACACCGUGUAUCAG  $\tt CUGAGCAAGGUGGAAGGCGAACAGCACGUGAUCAAGGG$ CAGACCUGUGUCCAGCAGCUUCGACCCUAUCAAGUUCCC UGAGGAUCAGUUCCAGGUGGCCCUGGACCAGGUGUUCG AGAACAUCGAGAAUUCCCAGGCUCUGGUGGACCAGUCCA ACAGAAUCCUGUCUAGCGCCGAGAAGGGAAACACCGGCU UCAUCAUCGUGAUCGUGAUCGCCGUGCUGGGCAGCU

10

#### TABLE 19-continued

| Strain | Nucleic Acid Sequence  | SEQ ID<br>NO: |
|--------|--|---------------|
|        | CCAUGAUCCUGGUGUCCAUCUUCAUCAUUAUCAAGAAGA<br>CCAAGAAGCCCACCGGCGCUCCUCAGAACUGAGCGGAG<br>UGACCAACAAUGGCUUCAUCCCUCACAAC |               |

### EQUIVALENTS

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the disclosure described herein. Such equivalents are intended to be 15 encompassed by the following claims.

All references, including patent documents, disclosed herein are incorporated by reference in their entirety.

#### SEQUENCE LISTING

<160> NUMBER OF SEQ ID NOS: 147 <210> SEO ID NO 1 <211> LENGTH: 1620 <212> TYPE: DNA <213 > ORGANISM: Unknown <220> FEATURE: <223> OTHER INFORMATION: Human metapneumovirus <400> SEQUENCE: 1 atgagctgga aggtggtgat tatcttcagc ctgctgatta cacctcaaca cggcctgaag 60 gagagetace tggaagagag etgeteeace ateacegagg getacetgag egtgetgegg 120 accggctggt acaccaacgt gttcaccctg gaggtgggcg acgtggagaa cctgacctgc 180 agegaeggee ctageetgat caagaeegag etggaeetga eeaagagege tetgagagag 240 ctgaagaccg tgtccgccga ccagctggcc agagaggaac agatcgagaa ccctcggcag 300 agcagattcg tgctgggcgc catcgctctg ggagtcgccg ctgccgctgc agtgacagct 360 ggagtggcca ttgctaagac catcagactg gaaagcgagg tgacagccat caacaatgcc 420 ctgaagaaga ccaacgaggc cgtgagcacc ctgggcaatg gagtgagagt gctggccaca gccgtgcggg agctgaagga cttcgtgagc aagaacctga ccagagccat caacaagaac aagtgcgaca tcgatgacct gaagatggcc gtgagcttct cccagttcaa cagacggttc ctgaacgtgg tgagacagtt ctccgacaac gctggaatca cacctgccat tagcctggac 660 ctgatgaccg acgccgagct ggctagagcc gtgcccaaca tgcccaccag cgctggccag 720 atcaagctga tgctggagaa cagagccatg gtgcggagaa agggcttcgg catcctgatt 780 qqqqtqtatq qaaqctccqt qatctacatq qtqcaqctqc ccatcttcqq cqtqatcqac 840 acaccetqct qqatcqtqaa qqccqctcct aqctqctccq aqaaqaaaqq aaactatqcc 900 tgtctgctga gagaggacca gggctggtac tgccagaacg ccggaagcac agtgtactat 960 1020 cccaacgaga aggactgcga gaccagaggc gaccacgtgt tctgcgacac cgctgccgga atcaacgtgg ccgagcagag caaggagtgc aacatcaaca tcagcacaac caactacccc 1080 tgcaaggtga gcaccggacg gcaccccatc agcatggtgg ctctgagccc tctgggcgct 1140 ctggtggcct gctataaggg cgtgtcctgt agcatcggca gcaatcgggt gggcatcatc 1200 aagcagetga acaagggatg etectacate accaaceagg aegeegacae egtgaeeate 1260 gacaacaccg tgtaccagct gagcaaggtg gagggcgagc agcacgtgat caagggcaga

| cccgt                | gaget ecagettega   | ccccatcaag | ttccctgagg  | accagttcaa | cgtggccctg | 1380 |  |
|----------------------|--|------------|-------------|------------|------------|------|--|
| gacca                | ggtgt ttgagaacat   | cgagaacagc | caggccctgg  | tggaccagag | caacagaatc | 1440 |  |
| ctgtc                | cagcg ctgagaaggg   | caacaccggc | ttcatcattg  | tgatcattct | gatcgccgtg | 1500 |  |
| ctggg                | caget ecatgateet   | ggtgagcatc | ttcatcatta  | tcaagaagac | caagaaaccc | 1560 |  |
| accgg                | agccc ctcctgagct   | gagcggcgtg | accaacaatg  | gcttcattcc | ccacaactga | 1620 |  |
| <211><212><213><223> | SEQ ID NO 2<br>LENGTH: 1620<br>TYPE: DNA<br>ORGANISM: Unkno<br>FEATURE:<br>OTHER INFORMATI |            | etapneumovi | rus        |            |      |  |
| <400>                | SEQUENCE: 2  |            |             |            |            |      |  |
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| gagag                | ttatt tggaagaatc   | atgtagtact | ataactgagg  | gatacctcag | tgttttaaga | 120  |  |
| acagg                | ctggt acactaatgt   | cttcacatta | gaagttggtg  | atgttgaaaa | tcttacatgt | 180  |  |
| actga                | tggac ctagcttaat   | caaaacagaa | cttgatctaa  | caaaaagtgc | tttaagggaa | 240  |  |
| ctcaa                | aacag tetetgetga   | tcagttggcg | agagaggagc  | aaattgaaaa | tcccagacaa | 300  |  |
| tcaag                | atttg tettaggtge   | gatagetete | ggagttgcta  | cagcagcagc | agtcacagca | 360  |  |
| ggcat                | tgcaa tagccaaaac   | cataaggctt | gagagtgagg  | tgaatgcaat | taaaggtgct | 420  |  |
| ctcaa                | acaaa ctaatgaagc   | agtatccaca | ttagggaatg  | gtgtgcgggt | cctagccact | 480  |  |
| gcagt                | gagag agctaaaaga   | atttgtgagc | aaaaacctga  | ctagtgcaat | caacaggaac | 540  |  |
| aaatg                | tgaca ttgctgatct   | gaagatggct | gtcagcttca  | gtcaattcaa | cagaagattt | 600  |  |
| ctaaa                | tgttg tgcggcagtt   | ttcagacaat | gcagggataa  | caccagcaat | atcattggac | 660  |  |
| ctgat                | gactg atgctgagtt   | ggccagagct | gtatcataca  | tgccaacatc | tgcagggcag | 720  |  |
| ataaa                | actga tgttggagaa   | ccgcgcaatg | gtaaggagaa  | aaggatttgg | aatcctgata | 780  |  |
| ggggt                | ctacg gaagctctgt   | gatttacatg | gttcaattgc  | cgatctttgg | tgtcatagat | 840  |  |
| acacc                | ttgtt ggatcatcaa   | ggcagctccc | tcttgctcag  | aaaaaaacgg | gaattatgct | 900  |  |
| tgcct                | cctaa gagaggatca   | agggtggtat | tgtaaaaatg  | caggatctac | tgtttactac | 960  |  |
| ccaaa                | tgaaa aagactgcga   | aacaagaggt | gatcatgttt  | tttgtgacac | agcagcaggg | 1020 |  |
| atcaa                | tgttg ctgagcaatc   | aagagaatgc | aacatcaaca  | tatctactac | caactaccca | 1080 |  |
| tgcaa                | agtca gcacaggaag   | acaccctata | agcatggttg  | cactatcacc | tctcggtgct | 1140 |  |
| ttggt                | ggctt gctataaagg   | ggtaagctgc | tegattggca  | gcaattgggt | tggaatcatc | 1200 |  |
| aaaca                | attac ccaaaggctg   | ctcatacata | accaaccagg  | atgcagacac | tgtaacaatt | 1260 |  |
| gacaa                | taccg tgtatcaact   | aagcaaagtt | gaaggtgaac  | agcatgtaat | aaaagggaga | 1320 |  |
| ccagt                | ttcaa gcagttttga   | tccaatcaag | tttcctgagg  | atcagttcaa | tgttgcgctt | 1380 |  |
| gatca                | agtet tegaaageat   | tgagaacagt | caggcactag  | tggaccagtc | aaacaaaatt | 1440 |  |
| ctaaa                | cagtg cagaaaaagg   | aaacactggt | ttcattatcg  | tagtaatttt | ggttgctgtt | 1500 |  |
|                      | tctaa ccatgatttc   |            |             |            |            | 1560 |  |
|                      | agcac ctccagagct   |            |             |            |            | 1620 |  |
| 59                   | J  | J 950500   |             | 5240400    |            | 2 2  |  |
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-continued

| aataccaaaa a   | taccaatgt                          | aacattaagc       | aagaaaagga         | aaagaagatt         | tcttggcttt      | 420  |
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| ttgttaggtg t   | tggatctgc                          | aatcgccagt       | ggcattgctg         | tatctaaggt         | cctgcaccta      | 480  |
| gaaggggaag t   | gaacaaaat                          | caaaagtgct       | ctactatcca         | caaacaaggc         | tgtagtcagc      | 540  |
| ttatcaaatg g   | gagttagtgt                         | cttaaccagc       | aaagtgttag         | acctcaaaaa         | ctatatagat      | 600  |
| aaacagttgt t   | acctattgt                          | gaacaagcaa       | agctgcagca         | tatcaaacat         | tgaaactgtg      | 660  |
| atagagttcc a   | acaaaagaa                          | caacagacta       | ctagagatta         | ccagggaatt         | tagtgttaat      | 720  |
| gcaggtgtaa c   | tacacctgt                          | aagcacttat       | atgttaacta         | atagtgaatt         | attatcatta      | 780  |
| atcaatgata t   | gcctataac                          | aaatgatcag       | aaaaagttaa         | tgtccaacaa         | tgttcaaata      | 840  |
| gttagacagc a   | aagttactc                          | tatcatgtcc       | ataataaagg         | aggaagtctt         | agcatatgta      | 900  |
| gtacaattac c   | actatatgg                          | tgtaatagat       | acaccctgtt         | ggaaactgca         | cacatcccct      | 960  |
| ctatgtacaa c   | caacacaaa                          | ggaagggtcc       | aacatctgct         | taacaagaac         | cgacagagga      | 1020 |
| tggtattgtg a   | caatgcagg                          | atcagtatct       | ttcttcccac         | aagctgaaac         | atgtaaagtt      | 1080 |
| caatcgaatc g   | ggtattttg                          | tgacacaatg       | aacagtttaa         | cattaccaag         | tgaagtaaat      | 1140 |
| ctctgcaaca t   | tgacatatt                          | caaccccaaa       | tatgattgca         | aaattatgac         | ttcaaaaaca      | 1200 |
| gatgtaagca g   | ctccgttat                          | cacatctcta       | ggagccattg         | tgtcatgcta         | tggcaaaact      | 1260 |
| aaatgtacag c   | atccaataa                          | aaatcgtggg       | atcataaaga         | cattttctaa         | cgggtgtgat      | 1320 |
| tatgtatcaa a   | ıtaagggggt                         | ggatactgtg       | tctgtaggta         | atacattata         | ttatgtaaat      | 1380 |
| aagcaagaag g   | gcaaaagtct                         | ctatgtaaaa       | ggtgaaccaa         | taataaattt         | ctatgaccca      | 1440 |
| ttagtgttcc c   | ctctgatga                          | atttgatgca       | tcaatatctc         | aagtcaatga         | gaagattaac      | 1500 |
| cagageetag e   | atttattcg                          | taaatccgat       | gaattattac         | ataatgtaaa         | tgctggtaaa      | 1560 |
| tccaccacaa a   | ıtatcatgat                         | aactactata       | attatagtga         | ttatagtaat         | attgttatca      | 1620 |
| ttaattgcag t   | tggactgct                          | cctatactgc       | aaggccagaa         | gcacaccagt         | cacactaagt      | 1680 |
| aaggatcaac t   | gagtggtat                          | aaataatatt       | gcatttagta         | actga              |                 | 1725 |
| <210> SEQ ID<br><211> LENGTH<br><212> TYPE:<br><213> ORGANI<br><220> FEATUR<br><223> OTHER | I: 539<br>PRT<br>SM: Unknov<br>RE: |                  | etapneumovii       | rus isolate        |                 |      |
| <400> SEQUEN   | ICE: 5                             |                  |                    |                    |                 |      |
| Met Ser Trp<br>1   | Lys Val Va<br>5                    | al Ile Ile       | Phe Ser Leu<br>10  | Leu Ile Thr        | Pro Gln<br>15   |      |
| His Gly Leu  | Lys Glu Se<br>20                   | -                | Glu Glu Ser<br>25  | Cys Ser Thr        | Ile Thr         |      |
| Glu Gly Tyr<br>35  | Leu Ser Va                         | al Leu Arg<br>40 | Thr Gly Trp        | Tyr Thr Asn<br>45  | Val Phe         |      |
| Thr Leu Glu<br>50  | Val Gly As                         | p Val Glu<br>55  | Asn Leu Thr        | Cys Ser Asp<br>60  | Gly Pro         |      |
| Ser Leu Ile<br>65  | Lys Thr Gl                         | _                | Leu Thr Lys<br>75  | Ser Ala Leu        | Arg Glu<br>80   |      |
| Leu Lys Thr  | Val Ser Al<br>85                   | la Asp Gln       | Leu Ala Arg<br>90  | Glu Glu Gln        | ı Ile Glu<br>95 |      |
| Asn Pro Arg  | Gln Ser Ar<br>100                  |                  | Leu Gly Ala<br>105 | Ile Ala Leu<br>110 |                 |      |
|  |                                    |                  |                    |                    |                 |      |

Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile 115 120 125

| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | ГЛа        | Asn<br>180 | ГЛа        | CAa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | ГЛа        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | CÀa        | Ser        | Glu        | Lys<br>295 | ГЛа        | Gly        | Asn        | Tyr        | Ala<br>300 | CÀa        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | ГÀа        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | ГÀЗ        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | ГÀа        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | CÀa        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390        | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| ГÀа        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CAa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Asn        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu<br>485 | ГÀа        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu        | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile        | Ile        | Lys<br>515 | Lys        | Thr        | Lys        | Lys        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly        | Val<br>530 | Thr        | Asn        | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |

| <211<br><212<br><213<br><220 | )> SE<br>L> LE<br>2> TY<br>3> OF<br>0> FE<br>3> OT | ENGTH<br>PE:<br>RGANI<br>EATUR | H: 53<br>PRT<br>[SM:<br>RE: | 9<br>Unkr  |            | : Hur      | nan m      | neta <u>r</u> | neur       | novi       | cus        |            |            |            |            |
|------------------------------|--|--------------------------------|-----------------------------|------------|------------|------------|------------|---------------|------------|------------|------------|------------|------------|------------|------------|
| < 400                        | )> SE  | EQUEN                          | ICE :                       | 6          |            |            |            |               |            |            |            |            |            |            |            |
| Met<br>1                     | Ser  | Trp                            | Lys                         | Val<br>5   | Met        | Ile        | Ile        | Ile           | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                          | Gly  | Leu                            | Lys<br>20                   | Glu        | Ser        | Tyr        | Leu        | Glu<br>25     | Glu        | Ser        | Cys        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly  | Tyr<br>35                      | Leu                         | Ser        | Val        | Leu        | Arg<br>40  | Thr           | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50  | Glu                            | Val                         | Gly        | Asp        | Val<br>55  | Glu        | Asn           | Leu        | Thr        | Gys        | Thr        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu  | Ile                            | Lys                         | Thr        | Glu<br>70  | Leu        | Asp        | Leu           | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | Lys  | Thr                            | Val                         | Ser<br>85  | Ala        | Asp        | Gln        | Leu           | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn                          | Pro  | Arg                            | Gln<br>100                  | Ser        | Arg        | Phe        | Val        | Leu<br>105    | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala                          | Thr  | Ala<br>115                     | Ala                         | Ala        | Val        | Thr        | Ala<br>120 | Gly           | Ile        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg                          | Leu<br>130   | Glu                            | Ser                         | Glu        | Val        | Asn<br>135 | Ala        | Ile           | Lys        | Gly        | Ala<br>140 | Leu        | Lys        | Gln        | Thr        |
| Asn<br>145                   | Glu  | Ala                            | Val                         | Ser        | Thr<br>150 | Leu        | Gly        | Asn           | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala                          | Val  | Arg                            | Glu                         | Leu<br>165 | Lys        | Glu        | Phe        | Val           | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Ser<br>175 | Ala        |
| Ile                          | Asn  | Arg                            | Asn<br>180                  | Lys        | Cys        | Asp        | Ile        | Ala<br>185    | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe                          | Ser  | Gln<br>195                     | Phe                         | Asn        | Arg        | Arg        | Phe<br>200 | Leu           | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp                          | Asn<br>210   | Ala                            | Gly                         | Ile        | Thr        | Pro<br>215 | Ala        | Ile           | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225                   | Glu  | Leu                            | Ala                         | Arg        | Ala<br>230 | Val        | Ser        | Tyr           | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile                          | Lys  | Leu                            |                             |            |            |            | Arg        |               |            |            |            |            | Lys        |            |            |
| Gly                          | Ile  | Leu                            | Ile<br>260                  | Gly        | Val        | Tyr        | Gly        | Ser<br>265    | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu                          | Pro  | Ile<br>275                     | Phe                         | Gly        | Val        | Ile        | Asp<br>280 | Thr           | Pro        | Cys        | Trp        | Ile<br>285 | Ile        | Lys        | Ala        |
| Ala                          | Pro<br>290   | Ser                            | Cys                         | Ser        | Glu        | Lys<br>295 | Asn        | Gly           | Asn        | Tyr        | Ala<br>300 | Càa        | Leu        | Leu        | Arg        |
| Glu<br>305                   | Asp  | Gln                            | Gly                         | Trp        | Tyr<br>310 | CÀa        | Lys        | Asn           | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro                          | Asn  | Glu                            | ГЛа                         | Asp<br>325 | СЛа        | Glu        | Thr        | Arg           | Gly<br>330 | Asp        | His        | Val        | Phe        | 335<br>235 | Asp        |
| Thr                          | Ala  | Ala                            | Gly<br>340                  | Ile        | Asn        | Val        | Ala        | Glu<br>345    | Gln        | Ser        | Arg        | Glu        | Cys<br>350 | Asn        | Ile        |
| Asn                          | Ile  | Ser<br>355                     | Thr                         | Thr        | Asn        | Tyr        | Pro<br>360 | Cys           | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |

-continued

Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys 375 380 Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Trp Val Gly Ile Ile Lys Gln Leu Pro Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro 435 440 445 Ile Lys Phe Pro Glu Asp Gln Phe Asn Val Ala Leu Asp Gln Val Phe Glu Ser Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Lys Ile Leu Asn Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Val Ile Leu Val Ala Val Leu Gly Leu Thr Met Ile Ser Val Ser Ile Ile Ile Ile Ile Lys Lys Thr Arg Lys Pro Thr Gly Ala Pro Pro Glu Leu Asn 520 Gly Val Thr Asn Gly Gly Phe Ile Pro His Ser 530 <210> SEO ID NO 7 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Unknown <220> FEATURE: <223> OTHER INFORMATION: Human metapneumovirus <400> SEQUENCE: 7 Met Ser Trp Lys Val Met Ile Ile Ile Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Thr Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Thr Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Arg Gln Ser Arg Phe Val Leu Gly Ala Ile Ala Leu Gly Val Ala Thr Ala Ala Ala Val Thr Ala Gly Ile Ala Ile Ala Lys Thr Ile 120 Arg Leu Glu Ser Glu Val Asn Ala Ile Lys Gly Ala Leu Lys Thr Thr 135 Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr Ala Val Arg Glu Leu Lys Glu Phe Val Ser Lys Asn Leu Thr Ser Ala 170 Ile Asn Lys Asn Lys Cys Asp Ile Ala Asp Leu Lys Met Ala Val Ser 185

| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Asn        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Ser        | Tyr        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asn<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Ile        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | Cys        | Ser        | Glu        | Lys<br>295 | Asp        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | Cys        | Lys        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | Cys        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Cys<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Arg        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390        | Ser        | Thr        | Gly        | Ser        | Asn<br>395 | Gln        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Pro        | Lys<br>405 | Gly        | Cys        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Arg<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Asn        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Ser        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Lys        | Ile<br>480 |
| Leu        | Asn        | Ser        | Ala        | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu        | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Leu        | Thr        | Met<br>505 | Ile        | Ser        | Val        | Ser        | Ile<br>510 | Ile        | Ile        |
| Ile        | Ile        | Lys<br>515 | Lys        | Thr        | Arg        | Lys        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Asn        |
| Gly        | Val<br>530 | Thr        | Asn        | Gly        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Ser        |            |            |            |            |            |
| <210       | )> SI      | EQ II      | ои с       | 8          |            |            |            |            |            |            |            |            |            |            |            |
| <212       | 2 > T      | PE:        |            |            | an re      | espi       | ratoi      | rv st      | znavi      | ial        | virı       | ıs         |            |            |            |
|            |            |            | ICE:       |            |            |            |            | •          | . 4        | -          |            |            |            |            |            |
|            |            |            |            |            | Len        | Lare       | Thr        | Zan        | Δls        | Tle        | Thr        | Thr        | Tle        | Leu        | Δla        |
| 1          |            |            |            | 5          |            |            |            |            | 10         |            |            |            |            | Leu<br>15  |            |
| Ala        | Val        | Thr        | Leu<br>20  | Cys        | Phe        | Ala        | Ser        | Ser<br>25  | Gln        | Asn        | Ile        | Thr        | Glu<br>30  | Glu        | Phe        |

| Tyr        | Gln        | Ser<br>35  | Thr        | Cya        | Ser        | Ala        | Val<br>40  | Ser        | Lys        | Gly        | Tyr        | Leu<br>45  | Ser        | Ala        | Leu        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Arg        | Thr<br>50  | Gly        | Trp        | Tyr        | Thr        | Ser<br>55  | Val        | Ile        | Thr        | Ile        | Glu<br>60  | Leu        | Ser        | Asn        | Ile        |
| Lys<br>65  | Glu        | Asn        | Lys        | Сув        | Asn<br>70  | Gly        | Thr        | Asp        | Ala        | Lys<br>75  | Val        | Lys        | Leu        | Ile        | 80<br>Lys  |
| Gln        | Glu        | Leu        | Aap        | Eys<br>Lys | Tyr        | Lys        | Asn        | Ala        | Val<br>90  | Thr        | Glu        | Leu        | Gln        | Leu<br>95  | Leu        |
| Met        | Gln        | Ser        | Thr<br>100 | Pro        | Ala        | Ala        | Asn        | Asn<br>105 | Arg        | Ala        | Arg        | Arg        | Glu<br>110 | Leu        | Pro        |
| Arg        | Phe        | Met<br>115 | Asn        | Tyr        | Thr        | Leu        | Asn<br>120 | Asn        | Thr        | Lys        | Asn        | Thr<br>125 | Asn        | Val        | Thr        |
| Leu        | Ser<br>130 | Lys        | Lys        | Arg        | Lys        | Arg<br>135 | Arg        | Phe        | Leu        | Gly        | Phe<br>140 | Leu        | Leu        | Gly        | Val        |
| Gly<br>145 | Ser        | Ala        | Ile        | Ala        | Ser<br>150 | Gly        | Ile        | Ala        | Val        | Ser<br>155 | Lys        | Val        | Leu        | His        | Leu<br>160 |
| Glu        | Gly        | Glu        | Val        | Asn<br>165 | Lys        | Ile        | Lys        | Ser        | Ala<br>170 | Leu        | Leu        | Ser        | Thr        | Asn<br>175 | Lys        |
| Ala        | Val        | Val        | Ser<br>180 | Leu        | Ser        | Asn        | Gly        | Val<br>185 | Ser        | Val        | Leu        | Thr        | Ser<br>190 | Lys        | Val        |
| Leu        | Asp        | Leu<br>195 | Lys        | Asn        | Tyr        | Ile        | Asp<br>200 | Lys        | Gln        | Leu        | Leu        | Pro<br>205 | Ile        | Val        | Asn        |
| Lys        | Gln<br>210 | Ser        | Cys        | Ser        | Ile        | Ser<br>215 | Asn        | Ile        | Glu        | Thr        | Val<br>220 | Ile        | Glu        | Phe        | Gln        |
| Gln<br>225 | Lys        | Asn        | Asn        | Arg        | Leu<br>230 | Leu        | Glu        | Ile        | Thr        | Arg<br>235 | Glu        | Phe        | Ser        | Val        | Asn<br>240 |
| Ala        | Gly        | Val        | Thr        | Thr<br>245 | Pro        | Val        | Ser        | Thr        | Tyr<br>250 | Met        | Leu        | Thr        | Asn        | Ser<br>255 | Glu        |
| Leu        | Leu        | Ser        | Leu<br>260 | Ile        | Asn        | Asp        | Met        | Pro<br>265 | Ile        | Thr        | Asn        | Asp        | Gln<br>270 | Lys        | ГÀз        |
| Leu        | Met        | Ser<br>275 | Asn        | Asn        | Val        | Gln        | Ile<br>280 | Val        | Arg        | Gln        | Gln        | Ser<br>285 | Tyr        | Ser        | Ile        |
| Met        | Ser<br>290 | Ile        | Ile        | Lys        | Glu        | Glu<br>295 | Val        | Leu        | Ala        | Tyr        | Val<br>300 | Val        | Gln        | Leu        | Pro        |
| Leu<br>305 | Tyr        | Gly        | Val        | Ile        | Asp<br>310 | Thr        | Pro        | Cys        | Trp        | Lys<br>315 | Leu        | His        | Thr        | Ser        | Pro<br>320 |
| Leu        | Cys        | Thr        | Thr        | Asn<br>325 |            | Lys        | Glu        | Gly        | Ser<br>330 |            | Ile        | CÀa        | Leu        | Thr<br>335 | Arg        |
| Thr        | Asp        | Arg        | Gly<br>340 | Trp        | Tyr        | Cys        | Asp        | Asn<br>345 | Ala        | Gly        | Ser        | Val        | Ser<br>350 | Phe        | Phe        |
| Pro        | Gln        | Ala<br>355 | Glu        | Thr        | CÀa        | Lys        | Val<br>360 | Gln        | Ser        | Asn        | Arg        | Val<br>365 | Phe        | Cys        | Asp        |
| Thr        | Met<br>370 | Asn        | Ser        | Leu        | Thr        | Leu<br>375 | Pro        | Ser        | Glu        | Val        | Asn<br>380 | Leu        | Cys        | Asn        | Ile        |
| Asp<br>385 | Ile        | Phe        | Asn        | Pro        | 390<br>390 | Tyr        | Asp        | Cys        | Lys        | Ile<br>395 | Met        | Thr        | Ser        | Lys        | Thr<br>400 |
| Asp        | Val        | Ser        | Ser        | Ser<br>405 | Val        | Ile        | Thr        | Ser        | Leu<br>410 | Gly        | Ala        | Ile        | Val        | Ser<br>415 | CAa        |
| Tyr        | Gly        | Lys        | Thr<br>420 | Lys        | Cys        | Thr        | Ala        | Ser<br>425 | Asn        | Lys        | Asn        | Arg        | Gly<br>430 | Ile        | Ile        |
| Lys        | Thr        | Phe<br>435 | Ser        | Asn        | Gly        | Сув        | Asp<br>440 | Tyr        | Val        | Ser        | Asn        | Lys<br>445 | Gly        | Val        | Asp        |

423 424

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Thr Val Ser Val Gly Asn Thr Leu Tyr Tyr Val Asn Lys Gln Glu Gly 450 Lys Ser Leu Tyr Val Lys Gly Glu Pro Ile Ile Asn Phe Tyr Asp Pro Leu Val Phe Pro Ser Asp Glu Phe Asp Ala Ser Ile Ser Gln Val Asn 490 Glu Lys Ile Asn Gln Ser Leu Ala Phe Ile Arg Lys Ser Asp Glu Leu Leu His Asn Val Asn Ala Gly Lys Ser Thr Thr Asn Ile Met Ile Thr Thr Ile Ile Ile Val Ile Ile Val Ile Leu Leu Ser Leu Ile Ala Val Gly Leu Leu Tyr Cys Lys Ala Arg Ser Thr Pro Val Thr Leu Ser Lys Asp Gln Leu Ser Gly Ile Asn Asn Ile Ala Phe Ser Asn <210> SEQ ID NO 9 <211> LENGTH: 1617 <212> TYPE: DNA <213 > ORGANISM: Human parainfluenza virus 3 <400> SEOUENCE: 9 atgccaattt caatactgtt aattattaca accatgatca tggcatcaca ctgccaaata 60 120 qacatcacaa aactacaqca tqtaqqtqta ttqqtcaaca qtcccaaaqq qatqaaqata tcacaaaact tcqaaacaaq atatctaatc ctqaqtctca taccaaaaat aqaaqattct 180 aactcttgtg gtgaccaaca gatcaagcaa tacaagaggt tattggatag actgatcatt 240 cctttatatg atggactaag attacagaag gatgtgatag tgactaatca agaatccaat 300 gaaaacactg atcccagaac agaacgattc tttggagggg taattggaac tattgctcta 360 ggagtagcaa ceteageaca aattaeagea geagttgete tggttgaage eaageaggea 420 agatcagaca ttgaaaaact caaggaagca atcagggaca caaataaagc agtgcagtca 480 gttcagagct ctgtaggaaa tttgatagta gcaattaaat cagtccagga ttatgtcaac 540 600 aaagaaatcg tgccatcgat tgcgagacta ggttgtgaag cagcaggact tcagttaggg attgcattaa cacagcatta ctcagaatta acaaatatat ttggtgataa cataggatcg 660 ttacaagaaa aaggaataaa attacaaggt atagcatcat tataccgtac aaatatcaca 720 gaaatattca caacatcaac agttgacaaa tatgatattt atgatctatt atttacagaa 780 tcaataaagg tgagagttat agatgttgat ttgaatgatt actcaataac cctccaagtc agactccctt tattgaccag actgctgaac actcaaatct acaaagtaga ttccatatca tacaatatee aaaatagaga atggtatate eetetteeea geeatateat gacgaaaggg 960 1020 qcatttctaq qtqqaqcaqa tqtcaaaqaa tqcataqaaq cattcaqcaq ttatatatqc cettetgate caggatttgt actaaaccat gaaatggaga getgtetate aggaaacata 1080 teccaatgte caagaaceae agteacatea gacatagtte etaggtatge atttgteaat ggaggagtgg ttgcgaattg tataacaact acatgtacat gcaatggtat cggtaataga 1200 atcaaccaac cacctgatca aggagtcaaa attataacac ataaagaatg taatacaata 1260 qqtatcaacq qaatqctatt caacacaaac aaaqaaqqaa ctcttqcatt ctacacacca 1320 gacgacataa cattaaacaa ttctgttgca cttgatccga ttgacatatc aatcgagctc 1380

aacaaqqcca aatcaqatct tqaqqaatca aaaqaatqqa taaqaaqqtc aaatcaaaaq

| ctagattcta ttggaagttg  | gcatcaatct | agcactacaa  | tcatagttat | tttgataatg | 1500 |
|--|------------|-------------|------------|------------|------|
| atgattatat tgtttataat  | taatataaca | ataattacaa  | ttgcaattaa | gtattacaga | 1560 |
| attcaaaaga gaaatcgagt  | ggatcaaaat | gataagccgt  | atgtattaac | aaacaag    | 1617 |
| <210 > SEQ ID NO 10<br><211 > LENGTH: 1716<br><212 > TYPE: DNA<br><213 > ORGANISM: Human | parainflue | nza virus 3 |            |            |      |
| <400> SEQUENCE: 10   |            |             |            |            |      |
| atggaatact ggaagcacac  | caaccacgga | aaggatgctg  | gtaatgagct | ggagacatcc | 60   |
| acagccactc atggcaacaa  | gctcaccaac | aagataacat  | atatattgtg | gacgataacc | 120  |
| ctggtgttat tatcaatagt  | cttcatcata | gtgctaacta  | attccatcaa | aagtgaaaag | 180  |
| gecegegaat cattgetaca  | agacataaat | aatgagttta  | tggaagttac | agaaaagatc | 240  |
| caagtggcat cggataatac  | taatgatcta | atacagtcag  | gagtgaatac | aaggcttctt | 300  |
| acaattcaga gtcatgtcca  | gaattatata | ccaatatcat  | tgacacaaca | aatatcggat | 360  |
| cttaggaaat tcattagtga  | aattacaatt | agaaatgata  | atcaagaagt | gccaccacaa | 420  |
| agaataacac atgatgtggg  | tataaaacct | ttaaatccag  | atgatttctg | gagatgcacg | 480  |
| tctggtcttc catctttgat  | gaaaactcca | aaaataagat  | taatgccggg | accaggatta | 540  |
| ttagctatgc caacgactgt  | tgatggctgt | gtcagaaccc  | cgtccttagt | gataaatgat | 600  |
| ctgatttatg cttacacctc  | aaatctaatt | actcgaggtt  | gccaggatat | agggaaatca | 660  |
| tatcaagtat tacagatagg  | gataataact | gtaaactcag  | acttggtacc | tgacttaaat | 720  |
| cctaggatct ctcatacctt  | caacataaat | gacaatagaa  | agtcatgttc | tctagcactc | 780  |
| ctaaatacag atgtatatca  | actgtgttca | accccaaaag  | ttgatgaaag | atcagattat | 840  |
| gcatcatcag gcatagaaga  | tattgtactt | gatattgtca  | attatgatgg | ctcaatctcg | 900  |
| acaacaagat ttaagaataa  | taatataagt | tttgatcaac  | catatgcggc | attataccca | 960  |
| tctgttggac cagggatata  | ctacaaaggc | aaaataatat  | ttctcgggta | tggaggtctt | 1020 |
| gaacatccaa taaatgagaa  | tgcaatctgc | aacacaactg  | ggtgtcctgg | gaaaacacag | 1080 |
| agagactgta atcaagcatc  | tcatagtcca | tggttttcag  | atagaaggat | ggtcaactct | 1140 |
| ataattgttg ttgacaaggg  | cttgaactca | gttccaaaat  | tgaaggtatg | gacgatatct | 1200 |
| atgagacaaa attactgggg  | gtcagaagga | agattacttc  | tactaggtaa | caagatctac | 1260 |
| atatacacaa gatctacaag  | ttggcacagc | aagttacaat  | taggaataat | tgacattact | 1320 |
| gactacagtg atataaggat  | aaaatggaca | tggcataatg  | tgctatcaag | accaggaaac | 1380 |
| aatgaatgtc catggggaca  | ttcatgtccg | gatggatgta  | taacgggagt | atataccgat | 1440 |
| gcatatccac tcaatcccac  | aggaagcatt | gtatcatctg  | tcatattgga | ctcacaaaaa | 1500 |
| tcgagagtca acccagtcat  | aacttactca | acagcaaccg  | aaagggtaaa | cgagctggct | 1560 |
| atccgaaaca aaacactctc  | agctgggtac | acaacaacaa  | gctgcattac | acactataac | 1620 |
| aaagggtatt gttttcatat  | agtagaaata | aatcataaaa  | gcttaaacac | atttcaaccc | 1680 |
| atgttgttca aaacagagat  | tccaaaaagc | tgcagt      |            |            | 1716 |
|  |            |             |            |            |      |

<sup>&</sup>lt;210> SEQ ID NO 11 <211> LENGTH: 1716 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE:

| <223> OTHER INFORMATION: Synthetic Polynucleotide   |      |
|---|------|
| <400> SEQUENCE: 11  |      |
| atggaatact ggaagcacac caaccacggc aaggacgccg gcaacgagct ggaaaccagc   | 60   |
| acagccacac acggcaacaa gctgaccaac aagatcacct acatcctgtg gaccatcacc   | 120  |
| ctggtgctgc tgagcatcgt gttcatcatc gtgctgacca atagcatcaa gagcgagaag   | 180  |
| gccagagaga gcctgctgca ggacatcaac aacgagttca tggaagtgac cgagaagatc   | 240  |
| caggtggcca gcgacaacac caacgacctg atccagagcg gcgtgaacac ccggctgctg   | 300  |
| accatccaga gccacgtgca gaactacatc cccatcagcc tgacccagca gatcagcgac   | 360  |
| ctgcggaagt tcatcagcga gatcaccatc cggaacgaca accaggaagt gccccccag  | 420  |
| agaatcaccc acgacgtggg catcaagccc ctgaaccccg acgatttctg gcggtgtaca   | 480  |
| ageggeetge ceageetgat gaagaceeee aagateegge tgatgeetgg ceetggaetg   | 540  |
| ctggccatgc ctaccacagt ggatggctgt gtgcggaccc ccagcctcgt gatcaacgat   | 600  |
| ctgatctacg cctacaccag caacctgatc acceggggct gccaggatat cggcaagagc   | 660  |
| taccaggtgc tgcagategg catcatcacc gtgaactceg acctggtgcc cgacetgaac   | 720  |
| ceteggatea gecacacett caacateaac gacaacagaa agagetgeag cetggetetg   | 780  |
| ctgaacaccg acgtgtacca gctgtgcagc acccccaagg tggacgagag aagcgactac   | 840  |
| gecageageg geategagga tategtgetg gacategtga actaegaegg cageateage   | 900  |
| accacceggt teaagaacaa caacateage ttegaceage eetacgeege eetgtaceet   | 960  |
| tetgtgggee etggeateta etacaaggge aagateatet teetgggeta eggeggeetg   | 1020 |
| gaacacccca tcaacgagaa cgccatctgc aacaccaccg gctgccctgg caagacccag   | 1080 |
| agagactgca atcaggccag ccacagcccc tggttcagcg accgcagaat ggtcaactct   | 1140 |
| atcatcgtgg tggacaaggg cctgaacagc gtgcccaagc tgaaagtgtg gacaatcagc   | 1200 |
| atgcgccaga actactgggg cagcgagggc agacttctgc tgctgggaaa caagatctac   | 1260 |
| atctacaccc ggtccaccag ctggcacagc aaactgcagc tgggaatcat cgacatcacc   | 1320 |
| gactacageg acateeggat caagtggaee tggcacaaeg tgetgageag acceggeaae   | 1380 |
| aatgagtgcc cttggggcca cagctgcccc gatggatgta tcaccggcgt gtacaccgac   | 1440 |
| gectacecce tgaatectae eggetecate gtgtecageg tgateetgga cagecagaaa   | 1500 |
| agcagagtga accccgtgat cacatacagc accgccaccg agagagtgaa cgaactggcc   | 1560 |
| atcagaaaca agaccctgag cgccggctac accaccacaa gctgcatcac acactacaac   | 1620 |
| aagggctact gettecacat egtggaaate aaccacaagt eeetgaacae ettecageee   | 1680 |
| atgctgttca agaccgagat ccccaagagc tgctcc   | 1716 |
| <210> SEQ ID NO 12 <211> LENGTH: 1617 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide <400> SEQUENCE: 12 |      |
| atgcccatca gcatcctgct gatcatcacc acaatgatca tggccagcca ctgccagatc   | 60   |
| gacatcacca agetgeagea egtgggegtg etegtgaaca geeccaaggg catgaagate   | 120  |
|   |      |
| agccagaact tcgagacacg ctacctgatc ctgagcctga tccccaagat cgaggacagc   | 180  |
| aacagetgeg gegaecagea gateaageag tacaagegge tgetggaeag aetgateate   | 240  |

| cccctgtacg   | acggcctgcg                      | gctgcagaaa        | ı gacgtgatcg       | tgaccaacca        | ggaaagcaac      | 300  |
|--|---------------------------------|-------------------|--------------------|-------------------|-----------------|------|
| gagaacaccg   | acccccggac                      | cgagagatto        | tteggeggeg         | tgatcggcac        | aatcgccctg      | 360  |
| ggagtggcca   | caagcgccca                      | gattacagco        | getgtggeee         | tggtggaagc        | caagcaggcc      | 420  |
| agaagcgaca   | tcgagaagct                      | gaaagaggcc        | atccgggaca         | ccaacaaggc        | cgtgcagagc      | 480  |
| gtgcagtcca   | gcgtgggcaa                      | tctgatcgtg        | gccatcaagt         | ccgtgcagga        | ctacgtgaac      | 540  |
| aaagaaatcg   | tgccctctat                      | cgcccggctg        | ggctgtgaag         | ctgccggact        | gcagctgggc      | 600  |
| attgccctga   | cacagcacta                      | cagcgagctg        | g accaacatct       | tcggcgacaa        | catcggcagc      | 660  |
| ctgcaggaaa   | agggcattaa                      | gctgcaggga        | atcgccagcc         | tgtaccgcac        | caacatcacc      | 720  |
| gagatettea   | ccaccagcac                      | cgtggataag        | g tacgacatct       | acgacctgct        | gttcaccgag      | 780  |
| agcatcaaag   | tgcgcgtgat                      | cgacgtggac        | ctgaacgact         | acagcatcac        | cctgcaagtg      | 840  |
| cggctgcccc   | tgctgaccag                      | actgctgaac        | acccagatct         | acaaggtgga        | cagcatctcc      | 900  |
| tacaacatcc   | agaaccgcga                      | gtggtacato        | cctctgccca         | gccacattat        | gaccaagggc      | 960  |
| geetttetgg   | gcggagccga                      | cgtgaaagag        | g tgcatcgagg       | ccttcagcag        | ctacatctgc      | 1020 |
| cccagcgacc   | ctggcttcgt                      | gctgaaccac        | gagatggaaa         | gctgcctgag        | cggcaacatc      | 1080 |
| agccagtgcc   | ccagaaccac                      | cgtgacctcc        | gacatcgtgc         | ccagatacgc        | cttcgtgaat      | 1140 |
| ggcggcgtgg   | tggccaactg                      | catcaccacc        | acctgtacct         | gcaacggcat        | cggcaaccgg      | 1200 |
| atcaaccagc   | ctcccgatca                      | gggcgtgaag        | , attatcaccc       | acaaagagtg        | taacaccatc      | 1260 |
| ggcatcaacg   | gcatgctgtt                      | caataccaac        | : aaagagggca       | ccctggcctt        | ctacaccccc      | 1320 |
| gacgatatca   | ccctgaacaa                      | ctccgtggct        | ctggacccca         | tcgacatctc        | catcgagctg      | 1380 |
| aacaaggcca   | agagcgacct                      | ggaagagtco        | : aaagagtgga       | tccggcggag        | caaccagaag      | 1440 |
| ctggactcta   | teggeagetg                      | gcaccagago        | agcaccacca         | tcatcgtgat        | cctgattatg      | 1500 |
| atgattatcc   | tgttcatcat                      | caacattacc        | atcatcacta         | tegecattaa        | gtactaccgg      | 1560 |
| atccagaaac   | ggaaccgggt                      | ggaccagaat        | gacaagccct         | acgtgctgac        | aaacaag         | 1617 |
| <210> SEQ 1<br><211> LENG'<br><212> TYPE<br><213> ORGAI<br><400> SEQUI | TH: 539<br>: PRT<br>NISM: Human | parainflue        | enza virus 3       |                   |                 |      |
| Met Pro Ile  | e Ser Ile L<br>5                | eu Leu Ile        | Ile Thr Thr        | Met Ile Met       | Ala Ser<br>15   |      |
|  |                                 | le Thr Lvs        | Leu Gln His        | Val Glv Val       |                 |      |
|  | 20                              |                   | 25                 | 30                |                 |      |
| Asn Ser Pro  | D Lys Gly M                     | et Lys Ile<br>40  | Ser Gln Asn        | Phe Glu Thi<br>45 | Arg Tyr         |      |
| Leu Ile Let<br>50  | ı Ser Leu I                     | le Pro Lys<br>55  | Ile Glu Asp        | Ser Asn Ser<br>60 | Cys Gly         |      |
| Asp Gln Gli<br>65  | _                               | ln Tyr Lys<br>O   | Arg Leu Leu<br>75  | Asp Arg Let       | ı Ile Ile<br>80 |      |
| Pro Leu Ty:  | r Asp Gly L<br>85               | eu Arg Leu        | Gln Lys Asp<br>90  | Val Ile Val       | Thr Asn<br>95   |      |
| Gln Glu Se   | r Asn Glu A<br>100              | sn Thr Asp        | Pro Arg Thr<br>105 | Glu Arg Phe       |                 |      |
| Gly Val Ile  |                                 | le Ala Leu<br>120 | Gly Val Ala        | Thr Ser Ala       | a Gln Ile       |      |

| Thr        | Ala<br>130 | Ala        | Val        | Ala        | Leu        | Val<br>135 | Glu        | Ala        | Lys        | Gln        | Ala<br>140 | Arg        | Ser        | Asp        | Ile        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Glu<br>145 | Lys        | Leu        | Lys        | Glu        | Ala<br>150 | Ile        | Arg        | Asp        | Thr        | Asn<br>155 | ГЛа        | Ala        | Val        | Gln        | Ser<br>160 |
| Val        | Gln        | Ser        | Ser        | Val<br>165 | Gly        | Asn        | Leu        | Ile        | Val<br>170 | Ala        | Ile        | Lys        | Ser        | Val<br>175 | Gln        |
| Asp        | Tyr        | Val        | Asn<br>180 | Lys        | Glu        | Ile        | Val        | Pro<br>185 | Ser        | Ile        | Ala        | Arg        | Leu<br>190 | Gly        | Cys        |
| Glu        | Ala        | Ala<br>195 | Gly        | Leu        | Gln        | Leu        | Gly<br>200 | Ile        | Ala        | Leu        | Thr        | Gln<br>205 | His        | Tyr        | Ser        |
| Glu        | Leu<br>210 | Thr        | Asn        | Ile        | Phe        | Gly<br>215 | Asp        | Asn        | Ile        | Gly        | Ser<br>220 | Leu        | Gln        | Glu        | Lys        |
| Gly<br>225 | Ile        | Lys        | Leu        | Gln        | Gly<br>230 | Ile        | Ala        | Ser        | Leu        | Tyr<br>235 | Arg        | Thr        | Asn        | Ile        | Thr<br>240 |
| Glu        | Ile        | Phe        | Thr        | Thr<br>245 | Ser        | Thr        | Val        | Asp        | Lys<br>250 | Tyr        | Asp        | Ile        | Tyr        | Asp<br>255 | Leu        |
| Leu        | Phe        | Thr        | Glu<br>260 | Ser        | Ile        | ГЛа        | Val        | Arg<br>265 | Val        | Ile        | Asp        | Val        | Asp<br>270 | Leu        | Asn        |
| Asp        | Tyr        | Ser<br>275 | Ile        | Thr        | Leu        | Gln        | Val<br>280 | Arg        | Leu        | Pro        | Leu        | Leu<br>285 | Thr        | Arg        | Leu        |
| Leu        | Asn<br>290 | Thr        | Gln        | Ile        | Tyr        | Lys<br>295 | Val        | Asp        | Ser        | Ile        | Ser<br>300 | Tyr        | Asn        | Ile        | Gln        |
| Asn<br>305 | Arg        | Glu        | Trp        | Tyr        | Ile<br>310 | Pro        | Leu        | Pro        | Ser        | His<br>315 | Ile        | Met        | Thr        | Lys        | Gly<br>320 |
| Ala        | Phe        | Leu        | Gly        | Gly<br>325 | Ala        | Asp        | Val        | Lys        | Glu<br>330 | Сув        | Ile        | Glu        | Ala        | Phe<br>335 | Ser        |
| Ser        | Tyr        | Ile        | Cys<br>340 | Pro        | Ser        | Asp        | Pro        | Gly<br>345 | Phe        | Val        | Leu        | Asn        | His<br>350 | Glu        | Met        |
| Glu        | Ser        | Сув<br>355 | Leu        | Ser        | Gly        | Asn        | Ile<br>360 | Ser        | Gln        | Сла        | Pro        | Arg<br>365 | Thr        | Thr        | Val        |
| Thr        | Ser<br>370 | Asp        | Ile        | Val        | Pro        | Arg<br>375 | Tyr        | Ala        | Phe        | Val        | Asn<br>380 | Gly        | Gly        | Val        | Val        |
| Ala<br>385 | Asn        | СЛа        | Ile        | Thr        | Thr<br>390 | Thr        | Cys        | Thr        | СЛа        | Asn<br>395 | Gly        | Ile        | Gly        | Asn        | Arg<br>400 |
| Ile        | Asn        | Gln        | Pro        | Pro<br>405 | Asp        | Gln        | Gly        | Val        | Lys<br>410 | Ile        | Ile        | Thr        | His        | Lys<br>415 | Glu        |
| CAa        | Asn        |            | Ile<br>420 |            | Ile        | Asn        |            | Met<br>425 |            | Phe        | Asn        |            | Asn<br>430 |            | Glu        |
| Gly        | Thr        | Leu<br>435 | Ala        | Phe        | Tyr        | Thr        | Pro<br>440 | Asp        | Asp        | Ile        | Thr        | Leu<br>445 | Asn        | Asn        | Ser        |
| Val        | Ala<br>450 | Leu        | Asp        | Pro        | Ile        | Asp<br>455 | Ile        | Ser        | Ile        | Glu        | Leu<br>460 | Asn        | Lys        | Ala        | Lys        |
| Ser<br>465 | Asp        | Leu        | Glu        | Glu        | Ser<br>470 | Lys        | Glu        | Trp        | Ile        | Arg<br>475 | Arg        | Ser        | Asn        | Gln        | Lys<br>480 |
| Leu        | Asp        | Ser        | Ile        | Gly<br>485 | Ser        | Trp        | His        | Gln        | Ser<br>490 | Ser        | Thr        | Thr        | Ile        | Ile<br>495 | Val        |
| Ile        | Leu        | Ile        | Met<br>500 | Met        | Ile        | Ile        | Leu        | Phe<br>505 | Ile        | Ile        | Asn        | Ile        | Thr<br>510 | Ile        | Ile        |
| Thr        | Ile        | Ala<br>515 | Ile        | Lys        | Tyr        | Tyr        | Arg<br>520 | Ile        | Gln        | Lys        | Arg        | Asn<br>525 | Arg        | Val        | Asp        |
| Gln        | Asn<br>530 | Asp        | Lys        | Pro        | Tyr        | Val<br>535 | Leu        | Thr        | Asn        | Lys        |            |            |            |            |            |

|            |            | EQ II<br>ENGTI |            |            |            |            |            |            |            |            |            |            |            |            |            |
|------------|------------|----------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|            |            | YPE :<br>RGANI |            | Huma       | an pa      | arain      | nflue      | enza       | vir        | ıs 3       |            |            |            |            |            |
| < 400      | )> SI      | EQUEI          | ICE :      | 14         |            |            |            |            |            |            |            |            |            |            |            |
| Met<br>1   | Glu        | Tyr            | Trp        | Lys<br>5   | His        | Thr        | Asn        | His        | Gly<br>10  | Lys        | Asp        | Ala        | Gly        | Asn<br>15  | Glu        |
| Leu        | Glu        | Thr            | Ser<br>20  | Thr        | Ala        | Thr        | His        | Gly<br>25  | Asn        | Lys        | Leu        | Thr        | Asn<br>30  | Lys        | Ile        |
| Thr        | Tyr        | Ile<br>35      | Leu        | Trp        | Thr        | Ile        | Thr<br>40  | Leu        | Val        | Leu        | Leu        | Ser<br>45  | Ile        | Val        | Phe        |
| Ile        | Ile<br>50  | Val            | Leu        | Thr        | Asn        | Ser<br>55  | Ile        | Lys        | Ser        | Glu        | Lys<br>60  | Ala        | Arg        | Glu        | Ser        |
| Leu<br>65  | Leu        | Gln            | Asp        | Ile        | Asn<br>70  | Asn        | Glu        | Phe        | Met        | Glu<br>75  | Val        | Thr        | Glu        | Lys        | Ile<br>80  |
| Gln        | Val        | Ala            | Ser        | Asp<br>85  | Asn        | Thr        | Asn        | Asp        | Leu<br>90  | Ile        | Gln        | Ser        | Gly        | Val<br>95  | Asn        |
| Thr        | Arg        | Leu            | Leu<br>100 | Thr        | Ile        | Gln        | Ser        | His<br>105 | Val        | Gln        | Asn        | Tyr        | Ile<br>110 | Pro        | Ile        |
| Ser        | Leu        | Thr<br>115     | Gln        | Gln        | Ile        | Ser        | Asp<br>120 | Leu        | Arg        | Lys        | Phe        | Ile<br>125 | Ser        | Glu        | Ile        |
| Thr        | Ile<br>130 | Arg            | Asn        | Asp        | Asn        | Gln<br>135 | Glu        | Val        | Pro        | Pro        | Gln<br>140 | Arg        | Ile        | Thr        | His        |
| Asp<br>145 | Val        | Gly            | Ile        | ГÀа        | Pro<br>150 | Leu        | Asn        | Pro        | Asp        | Asp<br>155 | Phe        | Trp        | Arg        | CÀa        | Thr<br>160 |
| Ser        | Gly        | Leu            | Pro        | Ser<br>165 | Leu        | Met        | ГÀв        | Thr        | Pro<br>170 | Lys        | Ile        | Arg        | Leu        | Met<br>175 | Pro        |
| Gly        | Pro        | Gly            | Leu<br>180 | Leu        | Ala        | Met        | Pro        | Thr<br>185 | Thr        | Val        | Asp        | Gly        | Суs<br>190 | Val        | Arg        |
| Thr        | Pro        | Ser<br>195     | Leu        | Val        | Ile        | Asn        | Asp<br>200 | Leu        | Ile        | Tyr        | Ala        | Tyr<br>205 | Thr        | Ser        | Asn        |
| Leu        | Ile<br>210 | Thr            | Arg        | Gly        | CAa        | Gln<br>215 | Asp        | Ile        | Gly        | Lys        | Ser<br>220 | Tyr        | Gln        | Val        | Leu        |
| Gln<br>225 | Ile        | Gly            | Ile        | Ile        | Thr<br>230 | Val        | Asn        | Ser        | Asp        | Leu<br>235 | Val        | Pro        | Asp        | Leu        | Asn<br>240 |
| Pro        | Arg        | Ile            | Ser        | His<br>245 | Thr        | Phe        | Asn        | Ile        | Asn<br>250 | Asp        | Asn        | Arg        | Lys        | Ser<br>255 | Cys        |
| Ser        | Leu        | Ala            | Leu<br>260 | Leu        | Asn        | Thr        | Asp        | Val<br>265 | Tyr        | Gln        | Leu        | Сув        | Ser<br>270 | Thr        | Pro        |
| Lys        | Val        | Asp<br>275     | Glu        | Arg        | Ser        | Asp        | Tyr<br>280 | Ala        | Ser        | Ser        | Gly        | Ile<br>285 | Glu        | Asp        | Ile        |
| Val        | Leu<br>290 | Asp            | Ile        | Val        | Asn        | Tyr<br>295 | Asp        | Gly        | Ser        | Ile        | Ser<br>300 | Thr        | Thr        | Arg        | Phe        |
| Lys<br>305 | Asn        | Asn            | Asn        | Ile        | Ser<br>310 | Phe        | Asp        | Gln        | Pro        | Tyr<br>315 | Ala        | Ala        | Leu        | Tyr        | Pro<br>320 |
| Ser        | Val        | Gly            | Pro        | Gly<br>325 | Ile        | Tyr        | Tyr        | Lys        | Gly<br>330 | Lys        | Ile        | Ile        | Phe        | Leu<br>335 | Gly        |
| Tyr        | Gly        | Gly            | Leu<br>340 | Glu        | His        | Pro        | Ile        | Asn<br>345 | Glu        | Asn        | Ala        | Ile        | 350        | Asn        | Thr        |
| Thr        | Gly        | Сув<br>355     | Pro        | Gly        | Lys        | Thr        | Gln<br>360 | Arg        | Asp        | Сув        | Asn        | Gln<br>365 | Ala        | Ser        | His        |
| Ser        | Pro<br>370 | Trp            | Phe        | Ser        | Asp        | Arg<br>375 | Arg        | Met        | Val        | Asn        | Ser<br>380 | Ile        | Ile        | Val        | Val        |
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Trp Thr Trp His Asn Val Leu Ser Arg Pro Gly Asn Asn Glu Cys Pro
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Ala Tyr Pro Leu Asn Pro Thr Gly Ser Ile Val Ser Ser Val Ile Leu
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| ggcaattgcg | tggaatattc | cctctatggt | gtttcgggcc | gtggtgtttt | tcagaattgc | 1860 |
| acagctgtag | gtgttcgaca | gcagcgcttt | gtttatgatg | cgtaccagaa | tttagttggc | 1920 |
| tattattctg | atgatggcaa | ctactactgt | ctgcgtgctt | gtgttagtgt | tectgtttet | 1980 |
| gtcatctatg | ataaagaaac | taaaacccac | gctactctat | ttggtagtgt | tgcatgtgaa | 2040 |
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| gatettaeet | acgagatgtt | gtctcttcaa | caagttgtta | aagcccttaa | tgagtcttac | 3840 |
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| -concinued   |      |
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| atagacctta aagagcttgg caattatact tattacaaca aatggccgtg gtacatttgg  | 3900 |
| cttggtttca ttgctgggct tgttgcctta gctctatgcg tcttcttcat actgtgctgc  | 3960 |
| actggttgtg gcacaaactg tatgggaaaa cttaagtgta atcgttgttg tgatagatac  | 4020 |
| gaggaatacg acctcgagcc gcataaggtt catgttcact aa   | 4062 |
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| atgatccact ccgtgttcct cctcatgttc ctgttgaccc ccactgagtc agactgcaag  | 60   |
| etceegetgg gacagteeet gtgtgegetg cetgacaete etageaetet gaceecaege  | 120  |
| teegtgeggt eggtgeetgg egaaatgegg etggeeteea tegeetteaa teacceaate  | 180  |
| caagtggate agetgaatag etegtattte aagetgteea teeccaegaa ettetegtte  | 240  |
| ggggtcaccc aggagtacat ccagaccaca attcagaagg tcaccgtcga ttgcaagcaa  | 300  |
| tacgtgtgca acggetteca gaagtgcgag cagetgetga gagaatacgg gcagttttge  | 360  |
| agcaagatca accaggogot goatggagot aacttgogoo aggacgacto ogtgogoaac  | 420  |
| ctctttgcct ctgtgaagtc atcccagtcc tccccaatca tcccgggatt cggaggggac  | 480  |
| ttcaacctga ccctcctgga gcccgtgtcg atcagcaccg gtagcagatc ggcgcgctca  | 540  |
| gccattgaag atcttctgtt cgacaaggtc accatcgccg atccgggcta catgcaggga  | 600  |
| tacgacgact gtatgcagca gggaccagcc tccgcgaggg acctcatctg cgcgcaatac  | 660  |
| gtggccgggt acaaagtgct gcctcctctg atggatgtga acatggaggc cgcttatact  | 720  |
| tegtecetge teggetetat egeeggegtg gggtggaeeg eeggeetgte eteettegee  | 780  |
| gctatcccct ttgcacaatc cattttctac cggctcaacg gcgtgggcat tactcaacaa  | 840  |
| gtcctgtcgg agaaccagaa gttgatcgca aacaagttca atcaggccct gggggccatg  | 900  |
| cagactggat tcactacgac taacgaagcg ttccagaagg tccaggacgc tgtgaacaac  | 960  |
| aacgcccagg cgctctcaaa gctggcctcc gaactcagca acaccttcgg agccatcagc  | 1020 |
| gcatcgatcg gtgacataat tcagcggctg gacgtgctgg agcaggacgc ccagatcgac  | 1080 |
| egecteatea aeggaegget gaccacettg aatgeetteg tggcacaaca getggteegg  | 1140 |
| agcgaatcag cggcactttc cgcccaactc gccaaggaca aagtcaacga atgcgtgaag  | 1200 |
| gcccagtcca agaggtccgg tttctgcggt caaggaaccc atattgtgtc cttcgtcgtg  | 1260 |
| aacgcgccca acggtctgta ctttatgcac gtcggctact acccgagcaa tcatatcgaa  | 1320 |
| gtggtgtccg cctacggcct gtgcgatgcc gctaacccca ctaactgtat tgcccctgtg  | 1380 |
| aacggatatt ttattaagac caacaacacc cgcattgtgg acgaatggtc atacaccggt  | 1440 |
| togtoottot acgogocoga goocatoact toactgaaca ccaaatacgt ggotoogcaa  | 1500 |
| gtgacctacc agaacatete caccaatttg eegeegeege tgeteggaaa eageacegga  | 1560 |
| attgatttcc aagatgaact ggacgaattc ttcaagaacg tgtccacttc cattcccaac  | 1620 |
| ttcggaagcc tgacacagat caacaccacc cttctcgacc tgacctacga gatgctgagc  | 1680 |
| cttcaacaag tggtcaaggc cctgaacgag agctacatcg acctgaagga gctgggcaac  | 1740 |
| tatacetact acaacaagtg geoggacaag attgaggaga ttetgtegaa aatetaceae  | 1800 |
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<210> SEQ ID NO 23
<211> LENGTH: 4071
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEQUENCE: 23

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| cagacctttt | tcgacaagac | ctggcccaga | cccatcgacg | tgtccaaggc | cgacggcatc | 180  |
| atctatccac | aaggccggac | ctacagcaac | atcaccatta | cctaccaggg | cctgttccca | 240  |
| tatcaaggcg | accacggcga | tatgtacgtg | tactctgccg | gccacgccac | cggcaccaca | 300  |
| ccccagaaac | tgttcgtggc | caactacagc | caggacgtga | agcagttcgc | caacggcttc | 360  |
| gtcgtgcgga | ttggcgccgc | tgccaatagc | accggcacag | tgatcatcag | ccccagcacc | 420  |
| agcgccacca | tccggaagat | ctaccccgcc | ttcatgctgg | gcagctccgt | gggcaatttc | 480  |
| agcgacggca | agatgggccg | gttcttcaac | cacaccctgg | tgctgctgcc | cgatggctgt | 540  |
| ggcacactgc | tgagagcctt | ctactgcatc | ctggaaccca | gaagcggcaa | ccactgccct | 600  |
| gccggcaata | gctacaccag | cttcgccacc | taccacacac | ccgccaccga | ttgctccgac | 660  |
| ggcaactaca | accggaacgc | cagcctgaac | agcttcaaag | agtacttcaa | cctgcggaac | 720  |
| tgcaccttca | tgtacaccta | caatatcacc | gaggacgaga | tcctggaatg | gttcggcatc | 780  |
| acccagaccg | cccagggcgt | gcacctgttc | agcagcagat | acgtggacct | gtacggcggc | 840  |
| aacatgttcc | agtttgccac | cctgcccgtg | tacgacacca | tcaagtacta | cagcatcatc | 900  |
| ccccacagca | teeggteeat | ccagagcgac | agaaaagcct | gggccgcctt | ctacgtgtac | 960  |
| aagctgcagc | ccctgacctt | cctgctggac | ttcagcgtgg | acggctacat | cagacgggcc | 1020 |
| atcgactgcg | gcttcaacga | cctgagccag | ctgcactgct | cctacgagag | cttcgacgtg | 1080 |
| gaaagcggcg | tgtacagcgt | gtccagcttc | gaggccaagc | ctagcggcag | cgtggtggaa | 1140 |
| caggetgagg | gcgtggaatg | cgacttcagc | cctctgctga | geggeaeeee | tececaggtg | 1200 |
| tacaacttca | ageggetggt | gttcaccaac | tgcaattaca | acctgaccaa | gctgctgagc | 1260 |
| ctgttctccg | tgaacgactt | cacctgtagc | cagatcagcc | ctgccgccat | tgccagcaac | 1320 |
| tgctacagca | gcctgatcct | ggactacttc | agctaccccc | tgagcatgaa | gtccgatctg | 1380 |
| agegtgteet | ccgccggacc | catcagccag | ttcaactaca | agcagagctt | cagcaaccct | 1440 |
| acctgcctga | ttctggccac | cgtgccccac | aatctgacca | ccatcaccaa | gcccctgaag | 1500 |
| tacagctaca | tcaacaagtg | cagcagactg | ctgtccgacg | accggaccga | agtgccccag | 1560 |
| ctcgtgaacg | ccaaccagta | cagcccctgc | gtgtccatcg | tgcccagcac | cgtgtgggag | 1620 |
| gacggcgact | actacagaaa | gcagctgagc | cccctggaag | geggeggatg | gctggtggct | 1680 |
| tctggaagca | cagtggccat | gaccgagcag | ctgcagatgg | gctttggcat | caccgtgcag | 1740 |
| tacggcaccg | acaccaacag | cgtgtgcccc | aagctggaat | tcgccaatga | caccaagatc | 1800 |
| gccagccagc | tgggaaactg | cgtggaatac | tccctgtatg | gcgtgtccgg | acggggcgtg | 1860 |
| ttccagaatt | gcacagcagt | gggagtgcgg | cagcagagat | tcgtgtacga | tgcctaccag | 1920 |
| aacctcgtgg | gctactacag | cgacgacggc | aattactact | gcctgcgggc | ctgtgtgtcc | 1980 |
| gtgcccgtgt | ccgtgatcta | cgacaaagag | acaaagaccc | acgccacact | gttcggctcc | 2040 |
|            |            |            |            |            |            |      |

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<210> SEQ ID NO 24
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<sup>&</sup>lt;211> LENGTH: 1353

<sup>&</sup>lt;212> TYPE: PRT

<sup>&</sup>lt;213> ORGANISM: Unknown

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Middle East respiratory syndrome coronavirus

<sup>&</sup>lt;400> SEQUENCE: 24

| 1          |            |            |            | 5          |            |            |            |            | 10         |            |            |            |            | 15         |            |
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| Val        | Asp        | Ile<br>35  | Gln        | Gln        | Thr        | Phe        | Phe<br>40  | Asp        | Lys        | Thr        | Trp        | Pro<br>45  | Arg        | Pro        | Ile        |
| Asp        | Val<br>50  | Ser        | Lys        | Ala        | Asp        | Gly<br>55  | Ile        | Ile        | Tyr        | Pro        | Gln<br>60  | Gly        | Arg        | Thr        | Tyr        |
| Ser<br>65  | Asn        | Ile        | Thr        | Ile        | Thr<br>70  | Tyr        | Gln        | Gly        | Leu        | Phe<br>75  | Pro        | Tyr        | Gln        | Gly        | Asp<br>80  |
| His        | Gly        | Asp        | Met        | Tyr<br>85  | Val        | Tyr        | Ser        | Ala        | Gly<br>90  | His        | Ala        | Thr        | Gly        | Thr<br>95  | Thr        |
| Pro        | Gln        | Lys        | Leu<br>100 | Phe        | Val        | Ala        | Asn        | Tyr<br>105 | Ser        | Gln        | Asp        | Val        | Lys<br>110 | Gln        | Phe        |
| Ala        | Asn        | Gly<br>115 | Phe        | Val        | Val        | Arg        | Ile<br>120 | Gly        | Ala        | Ala        | Ala        | Asn<br>125 | Ser        | Thr        | Gly        |
| Thr        | Val<br>130 | Ile        | Ile        | Ser        | Pro        | Ser<br>135 | Thr        | Ser        | Ala        | Thr        | Ile<br>140 | Arg        | Lys        | Ile        | Tyr        |
| Pro<br>145 | Ala        | Phe        | Met        | Leu        | Gly<br>150 | Ser        | Ser        | Val        | Gly        | Asn<br>155 | Phe        | Ser        | Asp        | Gly        | Lys<br>160 |
| Met        | Gly        | Arg        | Phe        | Phe<br>165 | Asn        | His        | Thr        | Leu        | Val<br>170 | Leu        | Leu        | Pro        | Asp        | Gly<br>175 | CÀa        |
| Gly        | Thr        | Leu        | Leu<br>180 | Arg        | Ala        | Phe        | Tyr        | Cys<br>185 | Ile        | Leu        | Glu        | Pro        | Arg<br>190 | Ser        | Gly        |
| Asn        | His        | Cys<br>195 | Pro        | Ala        | Gly        | Asn        | Ser<br>200 | Tyr        | Thr        | Ser        | Phe        | Ala<br>205 | Thr        | Tyr        | His        |
| Thr        | Pro<br>210 | Ala        | Thr        | Asp        | CAa        | Ser<br>215 | Asp        | Gly        | Asn        | Tyr        | Asn<br>220 | Arg        | Asn        | Ala        | Ser        |
| Leu<br>225 | Asn        | Ser        | Phe        | Lys        | Glu<br>230 | Tyr        | Phe        | Asn        | Leu        | Arg<br>235 | Asn        | Cys        | Thr        | Phe        | Met<br>240 |
| Tyr        | Thr        | Tyr        | Asn        | Ile<br>245 | Thr        | Glu        | Asp        | Glu        | Ile<br>250 | Leu        | Glu        | Trp        | Phe        | Gly<br>255 | Ile        |
| Thr        | Gln        | Thr        | Ala<br>260 | Gln        | Gly        | Val        | His        | Leu<br>265 | Phe        | Ser        | Ser        | Arg        | Tyr<br>270 | Val        | Asp        |
| Leu        | Tyr        | Gly<br>275 | Gly        | Asn        | Met        | Phe        | Gln<br>280 | Phe        | Ala        | Thr        | Leu        | Pro<br>285 | Val        | Tyr        | Asp        |
| Thr        | Ile<br>290 | ГÀз        | Tyr        | Tyr        | Ser        | Ile<br>295 | Ile        | Pro        | His        | Ser        | Ile<br>300 | Arg        | Ser        | Ile        | Gln        |
| Ser<br>305 | Asp        | Arg        | Lys        | Ala        | Trp<br>310 | Ala        | Ala        | Phe        | Tyr        | Val<br>315 | Tyr        | ГÀа        | Leu        | Gln        | Pro<br>320 |
| Leu        | Thr        | Phe        | Leu        | Leu<br>325 | Asp        | Phe        | Ser        | Val        | Asp<br>330 | Gly        | Tyr        | Ile        | Arg        | Arg<br>335 | Ala        |
| Ile        | Asp        | CÀa        | Gly<br>340 | Phe        | Asn        | Asp        | Leu        | Ser<br>345 | Gln        | Leu        | His        | CÀa        | Ser<br>350 | Tyr        | Glu        |
| Ser        | Phe        | Asp<br>355 | Val        | Glu        | Ser        | Gly        | Val<br>360 | Tyr        | Ser        | Val        | Ser        | Ser<br>365 | Phe        | Glu        | Ala        |
| Lys        | Pro<br>370 | Ser        | Gly        | Ser        | Val        | Val<br>375 | Glu        | Gln        | Ala        | Glu        | Gly<br>380 | Val        | Glu        | Cha        | Asp        |
| Phe<br>385 | Ser        | Pro        | Leu        | Leu        | Ser<br>390 | Gly        | Thr        | Pro        | Pro        | Gln<br>395 | Val        | Tyr        | Asn        | Phe        | Lys<br>400 |
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| Leu        | Phe        | Ser        | Val<br>420 | Asn        | Asp        | Phe        | Thr        | Cys<br>425 | Ser        | Gln        | Ile        | Ser        | Pro<br>430 | Ala        | Ala        |
|            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |

| Ile        | Ala        | Ser<br>435 | Asn        | Cys        | Tyr        | Ser        | Ser<br>440 | Leu        | Ile        | Leu        | Asp        | Tyr<br>445 | Phe        | Ser        | Tyr        |
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| Pro        | Leu<br>450 | Ser        | Met        | Lys        | Ser        | Asp<br>455 | Leu        | Ser        | Val        | Ser        | Ser<br>460 | Ala        | Gly        | Pro        | Ile        |
| Ser<br>465 | Gln        | Phe        | Asn        | Tyr        | Lys<br>470 | Gln        | Ser        | Phe        | Ser        | Asn<br>475 | Pro        | Thr        | Сла        | Leu        | Ile<br>480 |
| Leu        | Ala        | Thr        | Val        | Pro<br>485 | His        | Asn        | Leu        | Thr        | Thr<br>490 | Ile        | Thr        | Lys        | Pro        | Leu<br>495 | Lys        |
| Tyr        | Ser        | Tyr        | Ile<br>500 | Asn        | ГÀа        | Cys        | Ser        | Arg<br>505 | Leu        | Leu        | Ser        | Asp        | Asp<br>510 | Arg        | Thr        |
| Glu        | Val        | Pro<br>515 | Gln        | Leu        | Val        | Asn        | Ala<br>520 | Asn        | Gln        | Tyr        | Ser        | Pro<br>525 | CAa        | Val        | Ser        |
| Ile        | Val<br>530 | Pro        | Ser        | Thr        | Val        | Trp<br>535 | Glu        | Asp        | Gly        | Asp        | Tyr<br>540 | Tyr        | Arg        | Lys        | Gln        |
| Leu<br>545 | Ser        | Pro        | Leu        | Glu        | Gly<br>550 | Gly        | Gly        | Trp        | Leu        | Val<br>555 | Ala        | Ser        | Gly        | Ser        | Thr<br>560 |
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| Tyr        | Gly        | Thr        | Asp<br>580 | Thr        | Asn        | Ser        | Val        | Cys        | Pro        | Lys        | Leu        | Glu        | Phe<br>590 | Ala        | Asn        |
| Asp        | Thr        | Lys<br>595 | Ile        | Ala        | Ser        | Gln        | Leu<br>600 | Gly        | Asn        | Cys        | Val        | Glu<br>605 | Tyr        | Ser        | Leu        |
| Tyr        | Gly<br>610 | Val        | Ser        | Gly        | Arg        | Gly<br>615 | Val        | Phe        | Gln        | Asn        | Cys<br>620 | Thr        | Ala        | Val        | Gly        |
| Val<br>625 | Arg        | Gln        | Gln        | Arg        | Phe<br>630 | Val        | Tyr        | Asp        | Ala        | Tyr<br>635 | Gln        | Asn        | Leu        | Val        | Gly<br>640 |
| Tyr        | Tyr        | Ser        | Asp        | Asp<br>645 | Gly        | Asn        | Tyr        | Tyr        | Сув<br>650 | Leu        | Arg        | Ala        | Cys        | Val<br>655 | Ser        |
| Val        | Pro        | Val        | Ser<br>660 | Val        | Ile        | Tyr        | Asp        | Lys<br>665 | Glu        | Thr        | Lys        | Thr        | His<br>670 | Ala        | Thr        |
| Leu        | Phe        | Gly<br>675 | Ser        | Val        | Ala        | Cys        | Glu<br>680 | His        | Ile        | Ser        | Ser        | Thr<br>685 | Met        | Ser        | Gln        |
| Tyr        | Ser<br>690 | Arg        | Ser        | Thr        | Arg        | Ser<br>695 | Met        | Leu        | Lys        | Arg        | Arg<br>700 | Asp        | Ser        | Thr        | Tyr        |
| Gly<br>705 | Pro        | Leu        | Gln        | Thr        | Pro<br>710 | Val        | Gly        | Cys        | Val        | Leu<br>715 | Gly        | Leu        | Val        | Asn        | Ser<br>720 |
| Ser        | Leu        | Phe        | Val        | Glu<br>725 |            | CAa        | Lys        | Leu        | Pro<br>730 |            | Gly        | Gln        | Ser        | Leu<br>735 |            |
| Ala        | Leu        | Pro        | Asp<br>740 | Thr        | Pro        | Ser        | Thr        | Leu<br>745 | Thr        | Pro        | Arg        | Ser        | Val<br>750 | Arg        | Ser        |
| Val        | Pro        | Gly<br>755 | Glu        | Met        | Arg        | Leu        | Ala<br>760 | Ser        | Ile        | Ala        | Phe        | Asn<br>765 | His        | Pro        | Ile        |
| Gln        | Val<br>770 | Asp        | Gln        | Leu        | Asn        | Ser<br>775 | Ser        | Tyr        | Phe        | Lys        | Leu<br>780 | Ser        | Ile        | Pro        | Thr        |
| Asn<br>785 | Phe        | Ser        | Phe        | Gly        | Val<br>790 | Thr        | Gln        | Glu        | Tyr        | Ile<br>795 | Gln        | Thr        | Thr        | Ile        | Gln<br>800 |
| ГÀа        | Val        | Thr        | Val        | Asp<br>805 | CÀa        | Lys        | Gln        | Tyr        | Val<br>810 | Сла        | Asn        | Gly        | Phe        | Gln<br>815 | Lys        |
| Сув        | Glu        | Gln        | Leu<br>820 | Leu        | Arg        | Glu        | Tyr        | Gly<br>825 | Gln        | Phe        | Cys        | Ser        | Lys<br>830 | Ile        | Asn        |
| Gln        | Ala        | Leu<br>835 | His        | Gly        | Ala        | Asn        | Leu<br>840 | Arg        | Gln        | Asp        | Asp        | Ser<br>845 | Val        | Arg        | Asn        |

| -continue |
|-----------|
|           |

| Leu        | Phe<br>850  | Ala        | Ser        | Val        | Lys        | Ser<br>855   | Ser         | Gln        | Ser        | Ser        | Pro<br>860 |            | : Ile        | Pro        | Gly        |  |
|------------|-------------|------------|------------|------------|------------|--------------|-------------|------------|------------|------------|------------|------------|--------------|------------|------------|--|
| Phe<br>865 | Gly         | Gly        | Asp        | Phe        | Asn<br>870 | Leu          | Thr         | Leu        | Leu        | Glu<br>875 | Pro        | Val        | . Sei        | : Ile      | Ser<br>880 |  |
| Thr        | Gly         | Ser        | Arg        | Ser<br>885 | Ala        | Arg          | Ser         | Ala        | Ile<br>890 | Glu        | Asp        | Leu        | . Lev        | Phe<br>895 | -          |  |
| Lys        | Val         | Thr        | Ile<br>900 | Ala        | Asp        | Pro          | Gly         | Tyr<br>905 | Met        | Gln        | Gly        | Туг        | Asp<br>910   | _          | Cys        |  |
| Met        | Gln         | Gln<br>915 | Gly        | Pro        | Ala        | Ser          | Ala<br>920  | Arg        | Asp        | Leu        | Ile        | 925        |              | a Gln      | Tyr        |  |
| Val        | Ala<br>930  | Gly        | Tyr        | Lys        | Val        | Leu<br>935   | Pro         | Pro        | Leu        | Met        | Asp<br>940 |            | . Asr        | n Met      | Glu        |  |
| Ala<br>945 | Ala         | Tyr        | Thr        | Ser        | Ser<br>950 | Leu          | Leu         | Gly        | Ser        | Ile<br>955 | Ala        | Gly        | Va]          | . Gly      | Trp<br>960 |  |
| Thr        | Ala         | Gly        | Leu        | Ser<br>965 | Ser        | Phe          | Ala         | Ala        | Ile<br>970 | Pro        | Phe        | Ala        | Glr          | 975        |            |  |
| Phe        | Tyr         | Arg        | Leu<br>980 | Asn        | Gly        | Val          | Gly         | Ile<br>985 | Thr        | Gln        | Gln        | ı Val      | . Leu<br>990 |            | Glu        |  |
| Asn        | Gln         | Lys<br>995 | Leu        | Ile        | Ala        | Asn          | Lys<br>1000 |            | e Ası      | n Gl       | n Al       |            | u (          | Sly A      | la Met     |  |
| Gln        | Thr<br>1010 | -          | Phe        | e Thr      | Thr        | Thr<br>101   |             | en G       | lu A       | la Pi      |            | rg<br>.020 | Lys          | Val        | Gln        |  |
| Asp        | Ala<br>1025 |            | . Asr      | n Asr      | a Asn      | 103          |             | ln A       | la L       | eu S       |            | ys<br>.035 | Leu          | Ala        | Ser        |  |
| Glu        | Leu<br>1040 |            | : Asr      | n Thr      | Phe        | Gly<br>104   |             | la I       | le S       | er A       |            | er<br>.050 | Ile          | Gly        | Asp        |  |
| Ile        | Ile<br>1055 |            | a Arg      | g Leu      | ı Asp      | Val<br>106   |             | eu G       | lu G       | ln A       |            | la<br>.065 | Gln          | Ile        | Asp        |  |
| Arg        | Leu<br>1070 |            | e Asr      | n Gly      | / Arg      | Leu<br>107   |             | nr Tl      | nr Le      | eu A       |            | la<br>.080 | Phe          | Val        | Ala        |  |
| Gln        | Gln<br>1085 |            | ı Val      | l Arg      | g Ser      | Glu<br>109   |             | er A       | la A       | la L       |            | er<br>.095 | Ala          | Gln        | Leu        |  |
| Ala        | Lys<br>1100 |            | Lys        | val        | . Asn      | Glu<br>110   |             | ∕a V       | al L       | ys A       |            | ln<br>110  | Ser          | ГÀа        | Arg        |  |
| Ser        | Gly<br>1115 |            | е Суя      | Gly        | / Gln      | Gly<br>112   |             | nr H       | is I       | le V       |            | er<br>125  | Phe          | Val        | Val        |  |
| Asn        | Ala<br>1130 |            | Asr        | n Gly      | / Leu      | . Tyr<br>113 |             | ne Me      | et H       | is V       |            | ly<br>.140 | Tyr          | Tyr        | Pro        |  |
| Ser        | Asn<br>1145 |            | ; Ile      | e Glu      | ı Val      | Val<br>115   |             | er A       | la T       | yr G       |            | eu<br>.155 | Cys          | Asp        | Ala        |  |
| Ala        | Asn<br>1160 |            | Thi        | Asr        | n Cys      | 116<br>116   |             | La P:      | ro V       | al A       |            | 1y<br>.170 | Tyr          | Phe        | Ile        |  |
| Lys        | Thr<br>1175 |            | ı Asr      | n Thr      | Arg        | 11e          |             | al A       | ap G       | lu T       |            | er<br>185  | Tyr          | Thr        | Gly        |  |
| Ser        | Ser<br>1190 |            | ту1        | Ala        | Pro        | Glu<br>119   |             | ro I       | le Tl      | nr S       |            | eu<br>.200 | Asn          | Thr        | Lys        |  |
| Tyr        | Val<br>1205 |            | Pro        | Glr        | ı Val      | Thr<br>121   |             | yr G       | ln A       | en I       |            | er<br>.215 | Thr          | Asn        | Leu        |  |
| Pro        | Pro<br>1220 |            | Let        | ı Lev      | ı Gly      | Asr          |             | er Tl      | nr G       | ly I       |            | sp<br>.230 | Phe          | Gln        | Asp        |  |
| Glu        | Leu<br>1235 | _          | Glu        | ı Phe      | Phe        | Lys<br>124   |             | en V       | al S       | ∍r Tl      |            | er<br>.245 | Ile          | Pro        | Asn        |  |
| Phe        | Gly         | Ser        | : Let      | ı Thr      | Gln        | ıIl∈         | e As        | ≅n Tl      | nr Tl      | nr L       | eu L       | ieu        | Asp          | Leu        | Thr        |  |
|            |             |            |            |            |            |              |             |            |            |            |            |            |              |            |            |  |

|                              |                                  |                                |             |             |                 |             |            |            |            |            |            | COII       | L III      | lued       |            |
|------------------------------|----------------------------------|--------------------------------|-------------|-------------|-----------------|-------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|                              | 1250                             | )                              |             |             |                 | 125         | 55         |            |            |            | 1          | 260        |            |            |            |
| Tyr                          | Glu<br>1265                      |                                | . Le        | ı Sei       | . Leu           | 1 Gl1<br>12 |            | ln Va      | al Va      | al L       | •          | la<br>275  | Leu        | Asn        | Glu        |
| Ser                          | Tyr<br>1280                      |                                | e Ası       | Let         | ı Lys           | Gl:<br>128  |            | eu G       | ly A       | sn T       |            | hr<br>290  | Tyr        | Tyr        | Asn        |
| Lys                          | Trp<br>1295                      |                                | o Trp       | туз         | : Ile           | Trp         |            | eu G       | ly Pl      | ne I       |            | la<br>305  | Gly        | Leu        | Val        |
| Ala                          | Leu<br>1310                      |                                | a Let       | ı Cys       | val             | 131         |            | ne I       | le L       | eu C       |            | ys<br>320  | Thr        | Gly        | Cys        |
| Gly                          | Thr<br>1325                      |                                | n Cys       | s Met       | Gl <sub>y</sub> | / Lys       |            | eu Ly      | ys C       | ys A       |            | rg<br>335  | Cys        | CÀa        | Asp        |
| Arg                          | Tyr<br>1340                      |                                | ı Glu       | а Туг       | r Asp           | Le:         |            | lu Pi      | ro H       | is L       |            | al<br>350  | His        | Val        | His        |
| <211<br><212<br><213<br><220 | L> LE<br>2> T\<br>3> OF<br>0> FE | ENGTI<br>PE:<br>RGAN:<br>EATUI | ISM:<br>RE: | 353<br>Art: | ifici           |             | -          |            | Polyj      | pept       | ide        |            |            |            |            |
| < 400                        | )> SE                            | EQUE                           | ICE :       | 25          |                 |             |            |            |            |            |            |            |            |            |            |
| Met<br>1                     | Ile                              | His                            | Ser         | Val<br>5    | Phe             | Leu         | Leu        | Met        | Phe<br>10  | Leu        | Leu        | Thr        | Pro        | Thr<br>15  | Glu        |
| Ser                          | Tyr                              | Val                            | Asp<br>20   | Val         | Gly             | Pro         | Asp        | Ser<br>25  | Val        | Lys        | Ser        | Ala        | Суs<br>30  | : Ile      | Glu        |
| Val                          | Asp                              | Ile<br>35                      | Gln         | Gln         | Thr             | Phe         | Phe<br>40  | Asp        | Lys        | Thr        | Trp        | Pro<br>45  | Arg        | pro        | Ile        |
| Asp                          | Val<br>50                        | Ser                            | ГÀв         | Ala         | Asp             | Gly<br>55   | Ile        | Ile        | Tyr        | Pro        | Gln<br>60  | Gly        | Arg        | f Thr      | Tyr        |
| Ser<br>65                    | Asn                              | Ile                            | Thr         | Ile         | Thr<br>70       | Tyr         | Gln        | Gly        | Leu        | Phe<br>75  | Pro        | Tyr        | Gln        | Gly        | 80         |
| His                          | Gly                              | Asp                            | Met         | Tyr<br>85   | Val             | Tyr         | Ser        | Ala        | Gly<br>90  | His        | Ala        | Thr        | Gly        | Thr<br>95  | Thr        |
| Pro                          | Gln                              | Lys                            | Leu<br>100  | Phe         | Val             | Ala         | Asn        | Tyr<br>105 | Ser        | Gln        | Asp        | Val        | Lys<br>110 |            | Phe        |
| Ala                          | Asn                              | Gly<br>115                     | Phe         | Val         | Val             | Arg         | Ile<br>120 | Gly        | Ala        | Ala        | Ala        | Asn<br>125 | Ser        | Thr        | Gly        |
| Thr                          | Val<br>130                       | Ile                            | Ile         | Ser         | Pro             | Ser<br>135  | Thr        | Ser        | Ala        | Thr        | Ile<br>140 |            | Lys        | : Ile      | Tyr        |
| Pro<br>145                   | Ala                              | Phe                            | Met         | Leu         | Gly<br>150      | Ser         | Ser        | Val        | Gly        | Asn<br>155 | Phe        | Ser        | Asp        | Gly        | Lys<br>160 |
| Met                          | Gly                              | Arg                            | Phe         | Phe<br>165  | Asn             | His         | Thr        | Leu        | Val<br>170 | Leu        | Leu        | Pro        | Asp        | Gly<br>175 | Cys        |
| Gly                          | Thr                              | Leu                            | Leu<br>180  | Arg         | Ala             | Phe         | Tyr        | Cys<br>185 | Ile        | Leu        | Glu        | Pro        | Arg        |            | Gly        |
| Asn                          | His                              | Cys<br>195                     | Pro         | Ala         | Gly             | Asn         | Ser<br>200 | Tyr        | Thr        | Ser        | Phe        | Ala<br>205 |            | Tyr        | His        |
| Thr                          | Pro<br>210                       | Ala                            | Thr         | Asp         | Cys             | Ser<br>215  | Asp        | Gly        | Asn        | Tyr        | Asn<br>220 |            | Asn        | n Ala      | Ser        |
| Leu<br>225                   | Asn                              | Ser                            | Phe         | Lys         | Glu<br>230      | Tyr         | Phe        | Asn        | Leu        | Arg<br>235 | Asn        | CAa        | Thr        | Phe        | Met<br>240 |
| Tyr                          | Thr                              | Tyr                            | Asn         | Ile<br>245  | Thr             | Glu         | Asp        | Glu        | Ile<br>250 | Leu        | Glu        | Trp        | Phe        | Gly<br>255 | Ile        |
| Thr                          | Gln                              | Thr                            | Ala         | Gln         | Gly             | Val         | His        | Leu        | Phe        | Ser        | Ser        | Arg        | Tyr        | · Val      | Asp        |

|            |            |            |            |            |            |            |            |            |            |            |            |            | C III      | aca        |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|            |            |            | 260        |            |            |            |            | 265        |            |            |            |            | 270        |            |            |
| Leu        | Tyr        | Gly<br>275 | Gly        | Asn        | Met        | Phe        | Gln<br>280 | Phe        | Ala        | Thr        | Leu        | Pro<br>285 | Val        | Tyr        | Asp        |
| Thr        | Ile<br>290 | Lys        | Tyr        | Tyr        | Ser        | Ile<br>295 | Ile        | Pro        | His        | Ser        | Ile<br>300 | Arg        | Ser        | Ile        | Gln        |
| Ser<br>305 | Asp        | Arg        | Lys        | Ala        | Trp<br>310 | Ala        | Ala        | Phe        | Tyr        | Val<br>315 | Tyr        | Lys        | Leu        | Gln        | Pro<br>320 |
| Leu        | Thr        | Phe        | Leu        | Leu<br>325 | Asp        | Phe        | Ser        | Val        | Asp<br>330 | Gly        | Tyr        | Ile        | Arg        | Arg<br>335 | Ala        |
| Ile        | Asp        | Cys        | Gly<br>340 | Phe        | Asn        | Asp        | Leu        | Ser<br>345 | Gln        | Leu        | His        | CAa        | Ser<br>350 | Tyr        | Glu        |
| Ser        | Phe        | Asp<br>355 | Val        | Glu        | Ser        | Gly        | Val<br>360 | Tyr        | Ser        | Val        | Ser        | Ser<br>365 | Phe        | Glu        | Ala        |
| Lys        | Pro<br>370 | Ser        | Gly        | Ser        | Val        | Val<br>375 | Glu        | Gln        | Ala        | Glu        | Gly<br>380 | Val        | Glu        | Сув        | Asp        |
| Phe<br>385 | Ser        | Pro        | Leu        | Leu        | Ser<br>390 | Gly        | Thr        | Pro        | Pro        | Gln<br>395 | Val        | Tyr        | Asn        | Phe        | Lys<br>400 |
| Arg        | Leu        | Val        | Phe        | Thr<br>405 | Asn        | CÀa        | Asn        | Tyr        | Asn<br>410 | Leu        | Thr        | Lys        | Leu        | Leu<br>415 | Ser        |
| Leu        | Phe        | Ser        | Val<br>420 | Asn        | Asp        | Phe        | Thr        | Сув<br>425 | Ser        | Gln        | Ile        | Ser        | Pro<br>430 | Ala        | Ala        |
| Ile        | Ala        | Ser<br>435 | Asn        | CÀa        | Tyr        | Ser        | Ser<br>440 | Leu        | Ile        | Leu        | Asp        | Tyr<br>445 | Phe        | Ser        | Tyr        |
| Pro        | Leu<br>450 | Ser        | Met        | ГÀв        | Ser        | Asp<br>455 | Leu        | Ser        | Val        | Ser        | Ser<br>460 | Ala        | Gly        | Pro        | Ile        |
| Ser<br>465 | Gln        | Phe        | Asn        | Tyr        | Lys<br>470 | Gln        | Ser        | Phe        | Ser        | Asn<br>475 | Pro        | Thr        | Cha        | Leu        | Ile<br>480 |
| Leu        | Ala        | Thr        | Val        | Pro<br>485 | His        | Asn        | Leu        | Thr        | Thr<br>490 | Ile        | Thr        | ГÀЗ        | Pro        | Leu<br>495 | Lys        |
| Tyr        | Ser        | Tyr        | Ile<br>500 | Asn        | rys        | CAa        | Ser        | Arg<br>505 | Leu        | Leu        | Ser        | Asp        | Asp<br>510 | Arg        | Thr        |
| Glu        | Val        | Pro<br>515 | Gln        | Leu        | Val        | Asn        | Ala<br>520 | Asn        | Gln        | Tyr        | Ser        | Pro<br>525 | Cha        | Val        | Ser        |
| Ile        | Val<br>530 | Pro        | Ser        | Thr        | Val        | Trp<br>535 | Glu        | Asp        | Gly        | Asp        | Tyr<br>540 | Tyr        | Arg        | Lys        | Gln        |
| Leu<br>545 | Ser        | Pro        | Leu        | Glu        | Gly<br>550 |            | Gly        |            | Leu        | Val<br>555 |            | Ser        | Gly        | Ser        | Thr<br>560 |
| Val        | Ala        | Met        | Thr        | Glu<br>565 | Gln        | Leu        | Gln        | Met        | Gly<br>570 | Phe        | Gly        | Ile        | Thr        | Val<br>575 | Gln        |
| Tyr        | Gly        | Thr        | Asp<br>580 | Thr        | Asn        | Ser        | Val        | Сув<br>585 | Pro        | Lys        | Leu        | Glu        | Phe<br>590 | Ala        | Asn        |
| Asp        | Thr        | Lys<br>595 | Ile        | Ala        | Ser        | Gln        | Leu<br>600 | Gly        | Asn        | CAa        | Val        | Glu<br>605 | Tyr        | Ser        | Leu        |
| Tyr        | Gly<br>610 | Val        | Ser        | Gly        | Arg        | Gly<br>615 | Val        | Phe        | Gln        | Asn        | Сув<br>620 | Thr        | Ala        | Val        | Gly        |
| Val<br>625 | Arg        | Gln        | Gln        | Arg        | Phe<br>630 | Val        | Tyr        | Asp        | Ala        | Tyr<br>635 | Gln        | Asn        | Leu        | Val        | Gly<br>640 |
| Tyr        | Tyr        | Ser        | Asp        | Asp<br>645 | Gly        | Asn        | Tyr        | Tyr        | Сув<br>650 | Leu        | Arg        | Ala        | СЛа        | Val<br>655 | Ser        |
| Val        | Pro        | Val        | Ser<br>660 | Val        | Ile        | Tyr        | Asp        | Lys<br>665 | Glu        | Thr        | ГÀа        | Thr        | His<br>670 | Ala        | Thr        |
| Leu        | Phe        | Gly<br>675 | Ser        | Val        | Ala        | Cys        | Glu<br>680 | His        | Ile        | Ser        | Ser        | Thr<br>685 | Met        | Ser        | Gln        |
|            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |

| Ser<br>690                 | Arg   | Ser  | Thr   | Arg   | Ser<br>695  | Met   | Leu  | Lys   | Arg  | Arg<br>700             | Asp   | Ser   | Thr   | Tyr   |
|----------------------------|---|--|---|---|---|---|--|---|--|------------------------|---|---|---|---|
| Pro                        | Leu   | Gln  | Thr   | Pro<br>710  | Val   | Gly   | Cha  | Val   | Leu<br>715   | Gly                    | Leu   | Val   | Asn   | Ser<br>720  |
| Leu                        | Phe   | Val  | Glu<br>725  | Asp   | CÀa   | Lys   | Leu  | Pro<br>730  | Leu  | Gly                    | Gln   | Ser   | Leu<br>735  | Cys   |
| Leu                        | Pro   | Asp<br>740   | Thr   | Pro   | Ser   | Thr   | Leu<br>745   | Thr   | Pro  | Arg                    | Ser   | Val<br>750  | Arg   | Ser   |
| Pro                        | Gly<br>755  | Glu  | Met   | Arg   | Leu   | Ala<br>760  | Ser  | Ile   | Ala  | Phe                    |   |   | Pro   | Ile   |
| Val<br>770                 | Asp   | Gln  | Leu   | Asn   | Ser<br>775  | Ser   | Tyr  | Phe   | Lys  | Leu<br>780             | Ser   | Ile   | Pro   | Thr   |
| Phe                        | Ser   | Phe  | Gly   | Val<br>790  | Thr   | Gln   | Glu  | Tyr   | Ile<br>795   | Gln                    | Thr   | Thr   | Ile   | Gln<br>800  |
| Val                        | Thr   | Val  | Asp<br>805  | CAa   | Lys   | Gln   | Tyr  | Val<br>810  | Cys  | Asn                    | Gly   | Phe   | Gln<br>815  | Lys   |
| Glu                        | Gln   | Leu<br>820   | Leu   | Arg   | Glu   | Tyr   | Gly<br>825   | Gln   | Phe  | Cys                    | Ser   | Lys   | Ile   | Asn   |
| Ala                        | Leu<br>835  | His  | Gly   | Ala   | Asn   | Leu<br>840  | Arg  | Gln   | Asp  | Asp                    |   |   | Arg   | Asn   |
| Phe<br>850                 | Ala   | Ser  | Val   | rya   | Ser<br>855  | Ser   | Gln  | Ser   | Ser  | Pro<br>860             | Ile   | Ile   | Pro   | Gly   |
| Gly                        | Gly   | Asp  | Phe   | Asn<br>870  | Leu   | Thr   | Leu  | Leu   | Glu<br>875   | Pro                    | Val   | Ser   | Ile   | Ser<br>880  |
| Gly                        | Ser   | Arg  | Ser<br>885  | Ala   | Arg   | Ser   | Ala  | Ile<br>890  | Glu  | Asp                    | Leu   | Leu   | Phe<br>895  | Asp   |
| Val                        | Thr   | Ile<br>900   | Ala   | Asp   | Pro   | Gly   | Tyr<br>905   | Met   | Gln  | Gly                    | Tyr   | Asp<br>910  | Asp   | Cys   |
| Gln                        | Gln<br>915  | Gly  | Pro   | Ala   | Ser   | Ala<br>920  | Arg  | Asp   | Leu  | Ile                    |   |   | Gln   | Tyr   |
| Ala<br>930                 | Gly   | Tyr  | rys   | Val   | Leu<br>935  | Pro   | Pro  | Leu   | Met  | Asp<br>940             | Val   | Asn   | Met   | Glu   |
| Ala                        | Tyr   | Thr  | Ser   | Ser<br>950  | Leu   | Leu   | Gly  | Ser   | Ile<br>955   | Ala                    | Gly   | Val   | Gly   | Trp<br>960  |
| Ala                        | Gly   | Leu  | Ser<br>965  | Ser   | Phe   | Ala   | Ala  | Ile<br>970  | Pro  | Phe                    | Ala   | Gln   | Ser<br>975  | Ile   |
| Tyr                        | Arg   | Leu<br>980   | Asn   | Gly   | Val   | Gly   | Ile<br>985   | Thr   | Gln  | Gln                    | Val   | Leu<br>990  | Ser   | Glu   |
| Gln                        | Lys<br>995  | Leu  | Ile   | Ala   | Asn   |   |  | e Asr   | n Gli  | n Al                   |   |   | ly A  | la Met  |
| Thr<br>1010                |   | Phe  | Thr   | Thr   |   |   | en Gl  | lu Al   | La Pl  |                        |   | Lys 7   | Val (   | Gln   |
|                            |   |  |   |   |   |   |  |   |  |                        |   |   |   |   |
| Ala<br>1025                |   | . Asr  | n Asr   | n Asn   | 103   |   | ln Al  | la Le   | eu Se  |                        | 035<br>9  | Leu i   | Ala :   | Ser   |
|                            | Ser   |  |   | n Asn   | 103   | 30<br>7 Al  |  |   |  | 1<br>la S              | 035   | Leu i   |   |   |
| 1025<br>Leu                | Ser<br>Glr  | : Asr  | n Thr   |   | 103<br>Gly<br>104   | 30<br>/ Al<br>15<br>L Le  |  | Le Se   | er Al  | la S<br>1<br>sp A      | 035<br>er<br>050  |   | Gly A   | Asp   |
| 1025<br>Leu<br>1040<br>Ile | Ser<br>Glr  | : Asr  | n Thr   | r Phe   | 103<br>Gly<br>104<br>Val  | 30<br>/ Al<br>15<br>L Le<br>50  | la Il  | le Se   | er Al  | la S<br>1<br>sp A<br>1 | er<br>050<br>1a<br>065  | Ile (   | Gly A   | Aap   |
|                            | 690 Pro Leu Leu Pro Val Glu Ala Phe 850 Gly Val Gln Ala Ala Tyr Gln Thr | Pro Leu  Leu Phe  Leu Pro  Pro Gly 755  Val Asp 770  Phe Ser  Val Thr  Glu Gln  Ala Leu 835  Phe Ala 850  Gly Gly  Gly Ser  Val Thr  Gln Gln 915  Ala Gly 930  Ala Tyr  Ala Gly Tyr Arg  Gln Lys 995 | 690 Pro Leu Gln Leu Phe Val Leu Pro Asp 740 Pro Gly Glu 755 Val Asp Gln 770 Phe Ser Phe Val Thr Val Glu Gln Leu 820 Ala Leu His 835 Phe Ala Ser 850 Gly Gly Asp Gly Gly Asp Gly Gly Asp Gly Gly Tyr 915 Ala Gly Tyr 930 Ala Gly Tyr Ala Gly Leu Tyr Arg Leu 995 Thr Gly Phe Gly Phe Gly Phe | Pro Leu Gln Thr Leu Phe Val Glu 725 Leu Pro Asp Thr 740 Pro Gly Glu Met 775 Glu Met 775 Gln Leu Phe Ser Phe Gly Val Thr Val Asp 805 Glu Gln Leu Leu 820 Ala Leu His Gly 835 Phe Ala Ser Val 850 Gly Gly Asp Phe Gly Ser Arg Ser 885 Val Thr Ile Ala 900 Gln Gln Gly Pro 915 Ala Gly Tyr Lys 930 Ala Tyr Thr Ser Ala Gly Leu Ser 965 Tyr Arg Leu Asn 980 Gln Lys Leu Ile 995 Thr Gly Phe Thr | 690         Fro         Leu         Gln         Thr         Pro         710           Leu         Phe         Val         Glu         Asp         Asp         Thr         Pro         Asp         Pro         Asp         Pro         Asp         Pro         Asp         Pro         Ala         Asp         Asp | 695 Pro Leu Gln Thr Pro Val 710 Leu Phe Val Glu Asp Cys 725 Leu Pro Asp Thr Pro Ser 740 Pro Gly Glu Met Arg Leu 755 Phe Ser Phe Gly Val Thr 770 Val Thr Val Asp Cys Lys 805 Glu Gln Leu Arg Glu 820 Ala Leu His Gly Ala Asn 835 Phe Ala Ser Val Lys Ser 855 Gly Gly Asp Phe Asn Leu 870 Ala Thr Ile Ala Asp Pro 900 Gln Gln Gly Pro Ala Ser 915 Ala Gly Leu Ser Ser Leu 930 Ala Gly Leu Ser Ser Phe 965 Tyr Arg Leu Asn Gly Val 995 Thr Gly Phe Thr Thr Thr | 690 695  Pro Leu Gln Thr Pro Val Gly 710  Leu Phe Val Glu Asp Cys Lys 725  Leu Pro Asp Thr Pro Ser Thr 740  Pro Gly Glu Met Arg Leu Ala 755  Phe Ser Phe Gly Val Thr Gln 790  Val Asp Gln Leu Arg Glu Tyr 805  Glu Gln Leu Leu Arg Glu Tyr 820  Ala Leu His Gly Ala Asn Leu 835  Gly Gly Asp Phe Asn Leu Thr 870  Gly Gly Asp Phe Asn Leu Thr 870  Gly Ser Arg Ser Ala Arg Ser 885  Gly Gly Asp Phe Asn Leu Thr 870  Gln Gln Gln Gly Pro Ala Ser Ala 915  Ala Gly Tyr Lys Val Leu Pro 930  Ala Gly Tyr Lys Val Leu Pro 930  Ala Gly Leu Ser Ser Leu Leu 950  Ala Gly Leu Ser Ser Phe Ala 965  Tyr Arg Leu Asn Gly Val Gly 995  Cln Gln Lys Leu Ile Ala Asn Lys 995  Thr Gly Phe Thr Thr Thr As | 690 695  Pro Leu Gln Thr Pro Val Gly Cys 710  Leu Phe Val Glu Asp Cys Lys Leu 725  Leu Pro Asp Thr Pro Ser Thr Leu 740  Pro Gly Glu Met Arg Leu Ala Ser 755  Val Asp Gln Leu Asn Ser Ser Tyr 775  Phe Ser Phe Gly Val Thr Gln Glu 790  Val Thr Val Asp Cys Lys Gln Tyr 805  Glu Gln Leu Leu Arg Glu Tyr Gly 820  Ala Leu His Gly Ala Asn Leu Arg 835  Ala Ser Val Lys Ser Ser Gln 855  Gly Gly Asp Phe Asn Leu Thr Leu 870  Gly Gly Asp Phe Asn Leu Thr Leu 885  Gly Gly Asp Phe Asn Leu Thr Leu 885  Gly Gly Asp Phe Asn Leu Thr Leu 870  Gli Gln Gln Gly Pro Ala Ser Ala Arg 905  Ala Gly Tyr Lys Val Leu Pro Pro 935  Ala Tyr Thr Ser Ser Leu Leu Gly 950  Ala Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Ser Ser Phe Ala Ala  Gly Leu Asn Gly Val Gly Ile 980  Gln Lys Leu Ile Ala Asn Lys Phe 1000  Thr Gly Phe Thr Thr Thr Asn G | 690 695  Pro Leu Gln Thr Pro Val Gly Cys Val 710  Leu Phe Val Glu Asp Cys Lys Leu Pro 725  Leu Pro Asp Thr Pro Ser Thr Leu Thr 745  Pro Gly Glu Met Arg Leu Ala Ser Ile 760  Val Asp Gln Leu Asn Ser Ser Tyr Phe 775  Phe Ser Phe Gly Val Thr Gln Glu Tyr 790  Val Thr Val Asp Cys Lys Gln Tyr Val 810  Glu Gln Leu Leu Arg Glu Tyr Gly Gln 825  Ala Leu His Gly Ala Asn Leu Arg Gln 825  Ala Leu His Gly Ala Asn Leu Arg Gln Ser 850  Gly Gly Asp Phe Asn Leu Thr Leu Leu 870  Gly Gly Asp Phe Asn Leu Thr Leu Leu 885  Gly Gly Asp Phe Asn Leu Thr Leu Leu 880  Gli Gln Gln Gly Pro Ala Ser Ala Arg Asp 900  Ala Gly Tyr Lys Val Leu Pro Pro Leu 930  Ala Gly Tyr Lys Val Leu Pro Pro Leu 930  Ala Gly Leu Ser Ser Leu Leu Gly Ser 950  Ala Gly Leu Ser Ser Phe Ala Ala Ile 965  Tyr Arg Leu Asn Gly Val Gly Ile Thr 985  Gln Lys Leu Ile Ala Asn Lys Phe Asn 1000  Thr Gly Phe Thr Thr Thr Asn Glu Asn 1000  Thr Gly Phe Thr Thr Thr Asn Glu Asn 1000  Thr Gly Phe Thr Thr Thr Asn Glu Asn 1000 | 690                    | 690 695 700 Pro Leu Gln Thr Pro Val Gly Cys Val Leu Gly 715 Leu Phe Val Glu Asp Cys Lys Leu Pro Leu Gly 725 Leu Pro Asp Thr Pro Ser Thr Leu Thr Pro Arg 740 Pro Gly Glu Met Arg Leu Ala Ser Ile Ala Phe 755 Val Asp Cys Lys Gln Tyr Phe Lys Leu 770 Phe Ser Phe Gly Val Thr Gln Glu Tyr Ile Gln 790 Val Thr Val Asp Cys Lys Gln Tyr Val Cys Asn 810 Glu Gln Leu Leu Arg Glu Tyr Gly Gln Phe Cys 820 Ala Leu His Gly Ala Asn Leu Arg Gln Asp Asp 835 Gly Gly Asp Phe Asn Leu Thr Leu Glu Pro 870 Gly Gly Asp Phe Asn Leu Thr Leu Glu Pro 870 Gly Ser Arg Ser Ala Arg Ser Ala Ile Glu Asp 885 Val Thr Ile Ala Asp Pro Gly Tyr Met Gln Gly 900 Gln Gln Gly Pro Ala Ser Ala Arg Asp Leu Ile 915 Ala Gly Tyr Lys Val Leu Pro Pro Leu Met Asp 930 Ala Tyr Thr Ser Ser Leu Leu Gly Ser Ile Ala 950 Tyr Arg Leu Asn Gly Val Gly Ile Thr Gln Gln 1985 Tyr Arg Leu Asn Gly Val Gly Ile Thr Gln Gln 1995 Thr Gly Phe Thr Thr Thr Asn Glu Ala Phe G | 690 695 700  Pro Leu Gln Thr Pro Val Gly Cys Val Leu Gly Leu 715  Leu Phe Val Glu Asp Cys Lys Leu Pro Leu Gly Gln 725  Leu Pro Asp Thr Pro Ser Thr Leu Thr Pro Arg Ser 740  Pro Gly Glu Met Arg Leu Ala Ser Ile Ala Phe Asn 765  Val Asp Gln Leu Asn Ser Ser Tyr Phe Lys Leu Ser 770  Phe Ser Phe Gly Val Thr Gln Glu Tyr Ile Gln Thr 790  Val Thr Val Asp Cys Lys Gln Tyr Val Cys Asn Gly 805  Glu Gln Leu Leu Arg Glu Tyr Gly Gln Phe Cys Ser 825  Ala Leu His Gly Ala Asn Leu Arg Gln Asp Asp Ser 840  Phe Ala Ser Val Lys Ser Ser Gln Ser Ser Pro Ile 850  Gly Gly Asp Phe Asn Leu Thr Leu Leu Glu Pro Val 870  Gly Ser Arg Ser Ala Arg Ser Ala Ile Glu Asp Leu 885  Val Thr Ile Ala Asp Pro Gly Tyr Met Gln Gly Tyr 900  Gln Gln Gly Pro Ala Ser Ala Arg Asp Leu Ile Cys 915  Ala Gly Tyr Lys Val Leu Pro Pro Leu Met Asp Val 935  Ala Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala 965  Ala Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala 965  Ala Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala 980  Tyr Arg Leu Asn Gly Val Gly Ile Thr Gln Gln Val 985  Gln Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Le 100  Thr Gly Phe Thr Thr Thr Asn Glu Ala Phe Gln | 690 695 700 697 Pro Leu Gln Thr Pro Val Gly Cys Val Leu Gly Leu Val 710 Leu Phe Val Glu Asp Cys Lys Leu Pro Leu Gly Gln Ser 725 Leu Pro Asp Thr Pro Ser Thr Leu Thr Pro Arg Ser Val 740 Pro Gly Glu Met Arg Leu Ala Ser Ile Ala Phe Asn His 765 Val Asp Gln Leu Asn Ser Ser Tyr Phe Lys Leu Ser Ile 770 Phe Ser Phe Gly Val Thr Gln Glu Tyr Ile Gln Thr Thr 790 Val Thr Val Asp Cys Lys Gln Tyr Val Cys Asn Gly Phe 805 Glu Gln Leu Leu Arg Glu Tyr Gly Gln Phe Cys Ser Lys 825 Ala Leu His Gly Ala Asn Leu Arg Gln Asp Asp Ser Val 835 Gly Gly Asp Phe Asn Leu Thr Leu Leu Glu Pro Val Ser 870 Gly Ser Arg Ser Ala Arg Ser Ala Ile Glu Asp Leu Leu 885 Gly Gly Asp Phe Asn Leu Thr Leu Leu Glu Pro Val Ser 875 Gli Gln Gli Gly Pro Ala Ser Ala Arg Asp Leu Ile Cys Ala 915 Ala Gly Tyr Lys Val Leu Pro Pro Leu Met Asp 940 Ala Gly Tyr Lys Val Leu Pro Pro Leu Met Asp Val Asn 930 Ala Cys Cys Leu Leu Gly Ser Ile Ala Gly Val 955 Ala Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala Gln 965 Clu Gli Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala Gln 965 Clu Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala Gln 965 Clu Gly Leu Ser Ser Phe Ala Ala Ile Pro Phe Ala Gln 965 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly Pro Pro Pro Pro Phe Ala Gln 965 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly 995 Clu Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly Phe Thr Thr Thr Asn Glu Ala Phe Gln Lys Phe | Pro         Leu         Gln         Thr         Pro         Val         Gly         Cys         Val         Leu         Gly         Leu         Gly         Leu         Gly         Leu         Val         Asp         Cys         Lys         Leu         Pro         Leu         Gly         Gln         Ser         Leu         Pro         Arg         Gly         Asp         Thr         Pro         Ser         Thr         Leu         Thr         Pro         Arg         Ser         Val         Arg         Asp         Thr         Pro         Arg         Fro         765           Val         Asp         Glu         Met         Arg         Leu         Ala         Ser         Thr         Pro         Arg         Ser         Val         Arg         Fro         765         765         765         775 |

|                                      |  |                                    |                                     |                   |       |             |       |         |           |            |       | -001        | ILTI        | iuec  | ı     |
|--------------------------------------|--|------------------------------------|-------------------------------------|-------------------|-------|-------------|-------|---------|-----------|------------|-------|-------------|-------------|-------|-------|
| Ala                                  | Lys<br>1100  |                                    | Lys                                 | Val               | Asn   | Glu<br>1105 |       | Va      | 1 I       | Jys        | Ala   | Gln<br>1110 | Ser         | Lys   | Arg   |
| Ser                                  | Gly<br>1115  |                                    | Cys                                 | Gly               | Gln   | Gly<br>1120 |       | Hi      | s I       | [le        | Val   | Ser<br>1125 | Phe         | Val   | Val   |
| Asn                                  | Ala<br>1130  |                                    | Asn                                 | Gly               | Leu   | Tyr<br>1135 |       | e Me    | t F       | lis        | Val   | Gly<br>1140 | Tyr         | Tyr   | Pro   |
| Ser                                  | Asn<br>1145  |                                    | Ile                                 | Glu               | Val   | Val<br>1150 |       | Al      | a T       | Гуr        | Gly   | Leu<br>1155 | Сув         | Asp   | Ala   |
| Ala                                  | Asn<br>1160  |                                    | Thr                                 | Asn               | Cys   | Ile<br>1165 |       | ı Pr    | 7 O       | /al        | Asn   | Gly<br>1170 | Tyr         | Phe   | Ile   |
| ГÀз                                  | Thr<br>1175  |                                    | Asn                                 | Thr               | Arg   | Ile<br>1180 |       | . As    | pc        | 3lu        | Trp   | Ser<br>1185 | Tyr         | Thr   | Gly   |
| Ser                                  | Ser<br>1190  | Phe                                | Tyr                                 | Ala               | Pro   | Glu<br>1195 |       | ) Il    | e T       | Chr        | Ser   | Leu<br>1200 | Asn         | Thr   | Lys   |
| Tyr                                  | Val<br>1205  |                                    | Pro                                 | Gln               | Val   | Thr<br>1210 |       | Gl      | n A       | Asn        | Ile   | Ser<br>1215 | Thr         | Asn   | Leu   |
| Pro                                  | Pro<br>1220  |                                    | Leu                                 | Leu               | Gly   | Asn<br>1225 |       | Th      | ır G      | 3ly        | Ile   | Asp<br>1230 | Phe         | Gln   | Asp   |
| Glu                                  | Leu<br>1235  | _                                  | Glu                                 | Phe               | Phe   | Lys<br>1240 |       | ı Va    | .1 S      | Ser        | Thr   | Ser<br>1245 | Ile         | Pro   | Asn   |
| Phe                                  | Gly<br>1250  |                                    | Leu                                 | Thr               | Gln   | Ile<br>1255 |       | 1 Th    | ır T      | Chr        | Leu   | Leu<br>1260 | Asp         | Leu   | Thr   |
| Tyr                                  | Glu<br>1265  |                                    | Leu                                 | Ser               | Leu   | Gln<br>1270 |       | ı Va    | .1 V      | /al        | Lys   | Ala<br>1275 | Leu         | Asn   | Glu   |
| Ser                                  | Tyr<br>1280  |                                    | Asp                                 | Leu               | Lys   | Glu<br>1285 |       | ı Gl    | y F       | Asn        | Tyr   | Thr<br>1290 | Tyr         | Tyr   | Asn   |
| ГÀв                                  | Trp<br>1295  |                                    | Trp                                 | Tyr               | Ile   | Trp<br>1300 |       | ı Gl    | y F       | Phe        | Ile   | Ala<br>1305 | Gly         | Leu   | Val   |
| Ala                                  | Leu<br>1310  |                                    | Leu                                 | Cys               | Val   | Phe<br>1315 |       | : Il    | e I       | Leu        | Cys   | Сув<br>1320 | Thr         | Gly   | Сув   |
| Gly                                  | Thr<br>1325  |                                    | Cys                                 | Met               | Gly   | Lys         |       | ι Ьу    | s (       | Cys        | Asn   | Arg<br>1335 | Сув         | Cys   | Asp   |
| Arg                                  | Tyr<br>1340  |                                    | Glu                                 | Tyr               | Asp   | Leu<br>1345 |       | ı Pr    | o F       | lis        | Lys   | Val<br>1350 | His         | Val   | His   |
| <211<br><212<br><213<br><220<br><223 | 0> SE<br>L> LE<br>2> TY<br>3> OR<br>0> FE<br>3> OT | NGTH<br>PE:<br>GANI<br>ATUR<br>HER | : 61!<br>PRT<br>SM: 2<br>E:<br>INFO | 5<br>Arti<br>RMAT |       |             | -     |         | oly       | /pep       | otid€ | <b>=</b>    |             |       |       |
| < 400                                | )> SE  | QUEN                               | CE: 2                               | 26                |       |             |       |         |           |            |       |             |             |       |       |
| Met<br>1                             | Ile  | His                                |                                     | Val :             | Phe 1 | Leu I       | eu M  | let     | Phe<br>10 | e Le       | eu Le | eu Thi      | r Pro       | 15    | Glu   |
| Ser                                  | Asp  | _                                  | Lys 1<br>20                         | Leu :             | Pro 1 | Leu G       | _     | ln<br>5 | Ser       | c Le       | eu Cy | ys Ala      | a Let<br>30 | ı Pro | Asp   |
| Thr                                  |  | Ser<br>35                          | Thr 1                               | Leu '             | Thr 1 |             | arg S | er      | Val       | L Aı       | g Se  | er Val      | l Pro       | o Gly | / Glu |
| Met                                  | Arg<br>50  | Leu .                              | Alas                                | Ser               |       | Ala F<br>55 | he A  | sn      | His       | e Pi       | :0 I  | le Gli      | n Vai       | l Asj | Gln   |
| Leu<br>65                            | Asn  | Ser                                | Ser '                               |                   | Phe 1 | Lys I       | eu S  | er      | Il∈       | e Pi<br>75 |       | nr Ası      | n Phe       | e Sei | Phe   |
| Gly                                  | Val  | Thr                                | Gln (                               | Glu '             | Tyr : | Ile G       | 3ln T | hr      | Thr       | : I]       | Le G  | ln Lys      | s Vai       | l Thi | . Val |

| Asp        | Cha        | ГÀв        | Gln<br>100 | Tyr        | Val        | CAa        | Asn        | Gly<br>105 | Phe        | Gln        | ГÀа        | CAa        | Glu<br>110 | Gln        | Leu        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Leu        | Arg        | Glu<br>115 | Tyr        | Gly        | Gln        | Phe        | Cys<br>120 | Ser        | ГÀа        | Ile        | Asn        | Gln<br>125 | Ala        | Leu        | His        |
| Gly        | Ala<br>130 | Asn        | Leu        | Arg        | Gln        | Asp<br>135 | Asp        | Ser        | Val        | Arg        | Asn<br>140 | Leu        | Phe        | Ala        | Ser        |
| Val<br>145 | Lys        | Ser        | Ser        | Gln        | Ser<br>150 | Ser        | Pro        | Ile        | Ile        | Pro<br>155 | Gly        | Phe        | Gly        | Gly        | Asp<br>160 |
| Phe        | Asn        | Leu        | Thr        | Leu<br>165 | Leu        | Glu        | Pro        | Val        | Ser<br>170 | Ile        | Ser        | Thr        | Gly        | Ser<br>175 | Arg        |
| Ser        | Ala        | Arg        | Ser<br>180 | Ala        | Ile        | Glu        | Asp        | Leu<br>185 | Leu        | Phe        | Asp        | Lys        | Val<br>190 | Thr        | Ile        |
| Ala        | Asp        | Pro<br>195 | Gly        | Tyr        | Met        | Gln        | Gly<br>200 | Tyr        | Asp        | Asp        | CÀa        | Met<br>205 | Gln        | Gln        | Gly        |
| Pro        | Ala<br>210 | Ser        | Ala        | Arg        | Asp        | Leu<br>215 | Ile        | Cys        | Ala        | Gln        | Tyr<br>220 | Val        | Ala        | Gly        | Tyr        |
| Lys<br>225 | Val        | Leu        | Pro        | Pro        | Leu<br>230 | Met        | Asp        | Val        | Asn        | Met<br>235 | Glu        | Ala        | Ala        | Tyr        | Thr<br>240 |
| Ser        | Ser        | Leu        | Leu        | Gly<br>245 | Ser        | Ile        | Ala        | Gly        | Val<br>250 | Gly        | Trp        | Thr        | Ala        | Gly<br>255 | Leu        |
| Ser        | Ser        | Phe        | Ala<br>260 | Ala        | Ile        | Pro        | Phe        | Ala<br>265 | Gln        | Ser        | Ile        | Phe        | Tyr<br>270 | Arg        | Leu        |
| Asn        | Gly        | Val<br>275 | Gly        | Ile        | Thr        | Gln        | Gln<br>280 | Val        | Leu        | Ser        | Glu        | Asn<br>285 | Gln        | Lys        | Leu        |
| Ile        | Ala<br>290 | Asn        | Lys        | Phe        | Asn        | Gln<br>295 | Ala        | Leu        | Gly        | Ala        | Met<br>300 | Gln        | Thr        | Gly        | Phe        |
| Thr<br>305 | Thr        | Thr        | Asn        | Glu        | Ala<br>310 | Phe        | Gln        | Lys        | Val        | Gln<br>315 | Asp        | Ala        | Val        | Asn        | Asn<br>320 |
| Asn        | Ala        | Gln        | Ala        | Leu<br>325 | Ser        | Lys        | Leu        | Ala        | Ser<br>330 | Glu        | Leu        | Ser        | Asn        | Thr<br>335 | Phe        |
| Gly        | Ala        | Ile        | Ser<br>340 | Ala        | Ser        | Ile        | Gly        | Asp<br>345 | Ile        | Ile        | Gln        | Arg        | Leu<br>350 | Asp        | Val        |
| Leu        | Glu        | Gln<br>355 | Asp        | Ala        | Gln        | Ile        | Asp<br>360 | Arg        | Leu        | Ile        | Asn        | Gly<br>365 | Arg        | Leu        | Thr        |
| Thr        | Leu<br>370 | Asn        | Ala        | Phe        | Val        | Ala<br>375 | Gln        | Gln        | Leu        | Val        | Arg<br>380 | Ser        | Glu        | Ser        | Ala        |
| Ala<br>385 | Leu        | Ser        | Ala        | Gln        | Leu<br>390 | Ala        | Lys        | Asp        | Lys        | Val<br>395 | Asn        | Glu        | Càa        | Val        | Lys<br>400 |
| Ala        | Gln        | Ser        | Lys        | Arg<br>405 | Ser        | Gly        | Phe        | Сув        | Gly<br>410 | Gln        | Gly        | Thr        | His        | Ile<br>415 | Val        |
| Ser        | Phe        | Val        | Val<br>420 | Asn        | Ala        | Pro        | Asn        | Gly<br>425 | Leu        | Tyr        | Phe        | Met        | His<br>430 | Val        | Gly        |
| Tyr        | Tyr        | Pro<br>435 | Ser        | Asn        | His        | Ile        | Glu<br>440 | Val        | Val        | Ser        | Ala        | Tyr<br>445 | Gly        | Leu        | Cys        |
| Asp        | Ala<br>450 | Ala        | Asn        | Pro        | Thr        | Asn<br>455 | СЛа        | Ile        | Ala        | Pro        | Val<br>460 | Asn        | Gly        | Tyr        | Phe        |
| Ile<br>465 | Lys        | Thr        | Asn        | Asn        | Thr<br>470 | Arg        | Ile        | Val        | Asp        | Glu<br>475 | Trp        | Ser        | Tyr        | Thr        | Gly<br>480 |
| Ser        | Ser        | Phe        | Tyr        | Ala<br>485 | Pro        | Glu        | Pro        | Ile        | Thr<br>490 | Ser        | Leu        | Asn        | Thr        | Lys<br>495 | Tyr        |
| Val        | Ala        | Pro        | Gln<br>500 | Val        | Thr        | Tyr        | Gln        | Asn<br>505 | Ile        | Ser        | Thr        | Asn        | Leu<br>510 | Pro        | Pro        |
| Pro        | Leu        | Leu        | Gly        | Asn        | Ser        | Thr        | Gly        | Ile        | Asp        | Phe        | Gln        | Asp        | Glu        | Leu        | Asp        |

|                              |            | 515                             |                             |             |            |            | 520        |            |            |            |            | 525        |            |            |            |
|------------------------------|------------|---------------------------------|-----------------------------|-------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Glu                          | Phe<br>530 | Phe                             | Lys                         | Asn         | Val        | Ser<br>535 | Thr        | Ser        | Ile        | Pro        | Asn<br>540 | Phe        | Gly        | Ser        | Leu        |
| Thr<br>545                   | Gln        | Ile                             | Asn                         | Thr         | Thr<br>550 | Leu        | Leu        | Asp        | Leu        | Thr<br>555 | Tyr        | Glu        | Met        | Leu        | Ser<br>560 |
| Leu                          | Gln        | Gln                             | Val                         | Val<br>565  | Lys        | Ala        | Leu        | Asn        | Glu<br>570 | Ser        | Tyr        | Ile        | Asp        | Leu<br>575 | Lys        |
| Glu                          | Leu        | Gly                             | Asn<br>580                  | Tyr         | Thr        | Tyr        | Tyr        | Asn<br>585 | Lys        | Trp        | Pro        | Asp        | Lys<br>590 | Ile        | Glu        |
| Glu                          | Ile        | Leu<br>595                      | Ser                         | Lys         | Ile        | Tyr        | His<br>600 | Ile        | Glu        | Asn        | Glu        | Ile<br>605 | Ala        | Arg        | Ile        |
| ràa                          | Lys<br>610 | Leu                             | Ile                         | Gly         | Glu        | Ala<br>615 |            |            |            |            |            |            |            |            |            |
| <211<br><212<br><213<br><220 | )> FI      | ENGTH<br>(PE:<br>RGAN)<br>EATUR | H: 13<br>PRT<br>ISM:<br>RE: | 353<br>Unki |            | : Mi       | ddle       | East       | : re:      | spira      | atory      | i syi      | ndror      | me co      | pronavirus |
| < 400                        | )> SI      | EQUE                            | ICE:                        | 27          |            |            |            |            |            |            |            |            |            |            |            |
| Met<br>1                     | Ile        | His                             | Ser                         | Val<br>5    | Phe        | Leu        | Leu        | Met        | Phe<br>10  | Leu        | Leu        | Thr        | Pro        | Thr<br>15  | Glu        |
| Ser                          | Tyr        | Val                             | Asp<br>20                   | Val         | Gly        | Pro        | Asp        | Ser<br>25  | Val        | Lys        | Ser        | Ala        | Сув<br>30  | Ile        | Glu        |
| Val                          | Asp        | Ile<br>35                       | Gln                         | Gln         | Thr        | Phe        | Phe<br>40  | Asp        | Lys        | Thr        | Trp        | Pro<br>45  | Arg        | Pro        | Ile        |
| Asp                          | Val<br>50  | Ser                             | Lys                         | Ala         | Asp        | Gly<br>55  | Ile        | Ile        | Tyr        | Pro        | Gln<br>60  | Gly        | Arg        | Thr        | Tyr        |
| Ser<br>65                    | Asn        | Ile                             | Thr                         | Ile         | Thr<br>70  | Tyr        | Gln        | Gly        | Leu        | Phe<br>75  | Pro        | Tyr        | Gln        | Gly        | Asp<br>80  |
| His                          | Gly        | Asp                             | Met                         | Tyr<br>85   | Val        | Tyr        | Ser        | Ala        | Gly<br>90  | His        | Ala        | Thr        | Gly        | Thr<br>95  | Thr        |
| Pro                          | Gln        | ГÀа                             | Leu<br>100                  | Phe         | Val        | Ala        | Asn        | Tyr<br>105 | Ser        | Gln        | Asp        | Val        | Lys<br>110 | Gln        | Phe        |
| Ala                          | Asn        | Gly<br>115                      | Phe                         | Val         | Val        | Arg        | Ile<br>120 | Gly        | Ala        | Ala        | Ala        | Asn<br>125 | Ser        | Thr        | Gly        |
| Thr                          | Val<br>130 | Ile                             | Ile                         | Ser         |            | Ser<br>135 |            | Ser        | Ala        |            | Ile<br>140 | Arg        | ГÀа        | Ile        | Tyr        |
| Pro<br>145                   | Ala        | Phe                             | Met                         | Leu         | Gly<br>150 | Ser        | Ser        | Val        | Gly        | Asn<br>155 | Phe        | Ser        | Asp        | Gly        | Lys<br>160 |
| Met                          | Gly        | Arg                             | Phe                         | Phe<br>165  | Asn        | His        | Thr        | Leu        | Val<br>170 | Leu        | Leu        | Pro        | Asp        | Gly<br>175 | CAa        |
| Gly                          | Thr        | Leu                             | Leu<br>180                  | Arg         | Ala        | Phe        | Tyr        | Cys<br>185 | Ile        | Leu        | Glu        | Pro        | Arg<br>190 | Ser        | Gly        |
| Asn                          | His        | Суз<br>195                      | Pro                         | Ala         | Gly        | Asn        | Ser<br>200 | Tyr        | Thr        | Ser        | Phe        | Ala<br>205 | Thr        | Tyr        | His        |
| Thr                          | Pro<br>210 | Ala                             | Thr                         | Asp         | Сув        | Ser<br>215 | Asp        | Gly        | Asn        | Tyr        | Asn<br>220 | Arg        | Asn        | Ala        | Ser        |
| Leu<br>225                   | Asn        | Ser                             | Phe                         | Lys         | Glu<br>230 | Tyr        | Phe        | Asn        | Leu        | Arg<br>235 | Asn        | СЛа        | Thr        | Phe        | Met<br>240 |
| Tyr                          | Thr        | Tyr                             | Asn                         | Ile<br>245  | Thr        | Glu        | Asp        | Glu        | Ile<br>250 | Leu        | Glu        | Trp        | Phe        | Gly<br>255 | Ile        |
| Thr                          | Gln        | Thr                             | Ala                         | Gln         | Gly        | Val        | His        | Leu        | Phe        | Ser        | Ser        | Arg        | Tyr        | Val        | Asp        |

|            |            |            | 260        |            |            |            |            | 265        |            |            |            |            | 270        |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|            |            |            | 200        |            |            |            |            | 203        |            |            |            |            | 2,0        |            |            |
| Leu        | Tyr        | Gly<br>275 | Gly        | Asn        | Met        | Phe        | Gln<br>280 | Phe        | Ala        | Thr        | Leu        | Pro<br>285 | Val        | Tyr        | Asp        |
| Thr        | Ile<br>290 | ГÀз        | Tyr        | Tyr        | Ser        | Ile<br>295 | Ile        | Pro        | His        | Ser        | Ile<br>300 | Arg        | Ser        | Ile        | Gln        |
| Ser<br>305 | Asp        | Arg        | Lys        | Ala        | Trp<br>310 | Ala        | Ala        | Phe        | Tyr        | Val<br>315 | Tyr        | ГÀа        | Leu        | Gln        | Pro<br>320 |
| Leu        | Thr        | Phe        | Leu        | Leu<br>325 | Asp        | Phe        | Ser        | Val        | 330        | Gly        | Tyr        | Ile        | Arg        | Arg<br>335 | Ala        |
| Ile        | Asp        | CÀa        | Gly<br>340 | Phe        | Asn        | Asp        | Leu        | Ser<br>345 | Gln        | Leu        | His        | CAa        | Ser<br>350 | Tyr        | Glu        |
| Ser        | Phe        | Asp<br>355 | Val        | Glu        | Ser        | Gly        | Val<br>360 | Tyr        | Ser        | Val        | Ser        | Ser<br>365 | Phe        | Glu        | Ala        |
| Lys        | Pro<br>370 | Ser        | Gly        | Ser        | Val        | Val<br>375 | Glu        | Gln        | Ala        | Glu        | Gly<br>380 | Val        | Glu        | Cys        | Asp        |
| Phe<br>385 | Ser        | Pro        | Leu        | Leu        | Ser<br>390 | Gly        | Thr        | Pro        | Pro        | Gln<br>395 | Val        | Tyr        | Asn        | Phe        | Lys<br>400 |
| Arg        | Leu        | Val        | Phe        | Thr<br>405 | Asn        | Cys        | Asn        | Tyr        | Asn<br>410 | Leu        | Thr        | Lys        | Leu        | Leu<br>415 | Ser        |
| Leu        | Phe        | Ser        | Val<br>420 | Asn        | Asp        | Phe        | Thr        | Сув<br>425 | Ser        | Gln        | Ile        | Ser        | Pro<br>430 | Ala        | Ala        |
| Ile        | Ala        | Ser<br>435 | Asn        | Cya        | Tyr        | Ser        | Ser<br>440 | Leu        | Ile        | Leu        | Asp        | Tyr<br>445 | Phe        | Ser        | Tyr        |
| Pro        | Leu<br>450 | Ser        | Met        | ГÀа        | Ser        | Asp<br>455 | Leu        | Ser        | Val        | Ser        | Ser<br>460 | Ala        | Gly        | Pro        | Ile        |
| Ser<br>465 | Gln        | Phe        | Asn        | Tyr        | Lys<br>470 | Gln        | Ser        | Phe        | Ser        | Asn<br>475 | Pro        | Thr        | CAa        | Leu        | Ile<br>480 |
| Leu        | Ala        | Thr        | Val        | Pro<br>485 | His        | Asn        | Leu        | Thr        | Thr<br>490 | Ile        | Thr        | Lys        | Pro        | Leu<br>495 | Lys        |
| Tyr        | Ser        | Tyr        | Ile<br>500 | Asn        | ràa        | CÀa        | Ser        | Arg<br>505 | Leu        | Leu        | Ser        | Asp        | Asp<br>510 | Arg        | Thr        |
| Glu        | Val        | Pro<br>515 | Gln        | Leu        | Val        | Asn        | Ala<br>520 | Asn        | Gln        | Tyr        | Ser        | Pro<br>525 | Cha        | Val        | Ser        |
| Ile        | Val<br>530 | Pro        | Ser        | Thr        | Val        | Trp<br>535 | Glu        | Asp        | Gly        | Asp        | Tyr<br>540 | Tyr        | Arg        | ГÀЗ        | Gln        |
| Leu<br>545 | Ser        | Pro        | Leu        | Glu        | Gly<br>550 | Gly        | Gly        | Trp        | Leu        | Val<br>555 | Ala        | Ser        | Gly        | Ser        | Thr<br>560 |
| Val        | Ala        | Met        | Thr        | Glu<br>565 | Gln        | Leu        | Gln        | Met        | Gly<br>570 | Phe        | Gly        | Ile        | Thr        | Val<br>575 | Gln        |
| Tyr        | Gly        | Thr        | Asp<br>580 | Thr        | Asn        | Ser        | Val        | Cys<br>585 | Pro        | Lys        | Leu        | Glu        | Phe<br>590 | Ala        | Asn        |
| Asp        | Thr        | Lys<br>595 | Ile        | Ala        | Ser        | Gln        | Leu<br>600 | Gly        | Asn        | Cys        | Val        | Glu<br>605 | Tyr        | Ser        | Leu        |
| Tyr        | Gly<br>610 | Val        | Ser        | Gly        | Arg        | Gly<br>615 | Val        | Phe        | Gln        | Asn        | Cys<br>620 | Thr        | Ala        | Val        | Gly        |
| Val<br>625 | Arg        | Gln        | Gln        | Arg        | Phe<br>630 | Val        | Tyr        | Asp        | Ala        | Tyr<br>635 | Gln        | Asn        | Leu        | Val        | Gly<br>640 |
| Tyr        | Tyr        | Ser        | Asp        | Asp<br>645 | Gly        | Asn        | Tyr        | Tyr        | Cys<br>650 | Leu        | Arg        | Ala        | Cys        | Val<br>655 | Ser        |
| Val        | Pro        | Val        | Ser<br>660 | Val        | Ile        | Tyr        | Asp        | Lys<br>665 | Glu        | Thr        | Lys        | Thr        | His<br>670 | Ala        | Thr        |
| Leu        | Phe        | Gly<br>675 | Ser        | Val        | Ala        | Cys        | Glu<br>680 | His        | Ile        | Ser        | Ser        | Thr<br>685 | Met        | Ser        | Gln        |

| Tyr        | Ser<br>690  | Arg        | Ser        | Thr        | Arg        | Ser<br>695 | Met         | Leu        | Lys        | Arg        | Arg<br>700 | Asp        | Ser              | Thr        | Tyr        |
|------------|-------------|------------|------------|------------|------------|------------|-------------|------------|------------|------------|------------|------------|------------------|------------|------------|
| Gly<br>705 | Pro         | Leu        | Gln        | Thr        | Pro<br>710 | Val        | Gly         | Сув        | Val        | Leu<br>715 | Gly        | Leu        | Val              | Asn        | Ser<br>720 |
| Ser        | Leu         | Phe        | Val        | Glu<br>725 | Asp        | Cys        | Lys         | Leu        | Pro<br>730 | Leu        | Gly        | Gln        | Ser              | Leu<br>735 | Cha        |
| Ala        | Leu         | Pro        | Asp<br>740 | Thr        | Pro        | Ser        | Thr         | Leu<br>745 | Thr        | Pro        | Arg        | Ser        | Val<br>750       | Arg        | Ser        |
| Val        | Pro         | Gly<br>755 | Glu        | Met        | Arg        | Leu        | Ala<br>760  | Ser        | Ile        | Ala        | Phe        | Asn<br>765 | His              | Pro        | Ile        |
| Gln        | Val<br>770  | Asp        | Gln        | Leu        | Asn        | Ser<br>775 | Ser         | Tyr        | Phe        | Lys        | Leu<br>780 | Ser        | Ile              | Pro        | Thr        |
| Asn<br>785 | Phe         | Ser        | Phe        | Gly        | Val<br>790 | Thr        | Gln         | Glu        | Tyr        | Ile<br>795 | Gln        | Thr        | Thr              | Ile        | Gln<br>800 |
| Lys        | Val         | Thr        | Val        | Asp<br>805 | CAa        | Lys        | Gln         | Tyr        | Val<br>810 | Cys        | Asn        | Gly        | Phe              | Gln<br>815 | Lys        |
| Cys        | Glu         | Gln        | Leu<br>820 | Leu        | Arg        | Glu        | Tyr         | Gly<br>825 | Gln        | Phe        | Cys        | Ser        | Lys              | Ile        | Asn        |
| Gln        | Ala         | Leu<br>835 | His        | Gly        | Ala        | Asn        | Leu<br>840  | Arg        | Gln        | Asp        | Asp        | Ser<br>845 | Val              | Arg        | Asn        |
| Leu        | Phe<br>850  | Ala        | Ser        | Val        | ГÀа        | Ser<br>855 | Ser         | Gln        | Ser        | Ser        | Pro<br>860 | Ile        | Ile              | Pro        | Gly        |
| Phe<br>865 | Gly         | Gly        | Asp        | Phe        | Asn<br>870 | Leu        | Thr         | Leu        | Leu        | Glu<br>875 | Pro        | Val        | Ser              | Ile        | Ser<br>880 |
| Thr        | Gly         | Ser        | Arg        | Ser<br>885 | Ala        | Arg        | Ser         | Ala        | Ile<br>890 | Glu        | Asp        | Leu        | Leu              | Phe<br>895 | Asp        |
| Lys        | Val         | Thr        | Ile<br>900 | Ala        | Asp        | Pro        | Gly         | Tyr<br>905 | Met        | Gln        | Gly        | Tyr        | Asp<br>910       | Asp        | Cya        |
| Met        | Gln         | Gln<br>915 | Gly        | Pro        | Ala        | Ser        | Ala<br>920  | Arg        | Asp        | Leu        | Ile        | Cys<br>925 | Ala              | Gln        | Tyr        |
| Val        | Ala<br>930  | Gly        | Tyr        | Lys        | Val        | Leu<br>935 | Pro         | Pro        | Leu        | Met        | Asp<br>940 | Val        | Asn              | Met        | Glu        |
| Ala<br>945 | Ala         | Tyr        | Thr        | Ser        | Ser<br>950 | Leu        | Leu         | Gly        | Ser        | Ile<br>955 | Ala        | Gly        | Val              | Gly        | Trp<br>960 |
| Thr        | Ala         | Gly        | Leu        | Ser<br>965 | Ser        | Phe        | Ala         | Ala        | Ile<br>970 | Pro        | Phe        | Ala        | Gln              | Ser<br>975 | Ile        |
| Phe        | Tyr         | Arg        | Leu<br>980 | Asn        | Gly        | Val        | Gly         | Ile<br>985 | Thr        | Gln        | Gln        | Val        | Leu<br>990       | Ser        | Glu        |
| Asn        | Gln         | 995<br>995 | Leu        | Ile        | Ala        | Asn        | Lys<br>1000 |            | e Ası      | n Gli      | n Al       |            | u G:<br>05       | ly A       | la Met     |
| Gln        | Thr<br>1010 |            | Phe        | e Thr      | Thr        | Th:        |             | en G       | Lu A       | la Pl      |            | rg<br>020  | Lys V            | Val (      | Gln        |
| Asp        | Ala<br>1025 |            | . Asr      | n Asr      | n Asn      | 103        |             | ln A       | la Le      | eu Se      |            | 035<br>9   | Leu i            | Ala s      | Ser        |
| Glu        | Leu<br>1040 |            | : Asr      | n Thr      | Phe        | Gly        |             | la II      | Le Se      | er Al      |            | er<br>050  | Ile              | Gly A      | Aap.       |
| Ile        | Ile<br>1055 |            | n Arg      | g Leu      | ı Asp      | Va:        |             | eu G       | Lu G       | ln A:      |            | la<br>065  | Gln :            | Ile A      | Asp        |
| Arg        | Leu<br>1070 |            | e Asr      | n Gly      | ⁄ Arg      | Le:        |             | ır Th      | ır Le      | eu Ai      |            | la<br>080  | Phe <sup>v</sup> | Val i      | Ala        |
| Gln        | Gln<br>1085 |            | ı Val      | l Arg      | g Ser      | Gl:        |             | er Al      | La Ai      | la L       |            | er<br>095  | Ala (            | Gln 1      | Leu        |

| Ala Lys  |
|--|
| Asn Ala Pro Asn Gly Leu Tyr 1135 Phe Met His Val Gly Tyr Tyr Pro 1140 Pro 1155 Pro Try Phe Ile 1155 Pro Ile Thr Ser Leu Asn Thr Arg Ile Val Asp Glu Try Ser Tyr Thr Gly 1150 Pro Ile Thr Ser Leu Asn Thr Lys 1195 Pro Ile Thr Ser Leu Asn Thr Lys 1195 Pro Ile Thr Ser Leu Asn Thr Lys 1195 Pro Pro Ile Thr Ser Leu 1215 Pro Pro Pro Leu Leu Gly Asn Ser Thr Gly Ile Asp Phe Gln Asp 1220 Pro Leu Leu Gly Asn Val Ser Thr Ser Ile Pro Asn 1235 Pro Asn 1245 Pro Asn 1245 Pro Ile Thr Leu Leu Asp Leu Thr 1250 Pro Ile Met Leu Ser Leu Gln Gln Val Val Lys Ala Leu Asn Glu 1270 Pro Ile Asp Leu Asn Glu 1270 Pro Ile Asp 1290 Pro Ile Asn Ile Ser Tyr Ile Asp Leu Lys Glu Lys Gla Leu Gly Asn Tyr Thr Tyr Tyr Asn 1280 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1285 Pro Trp Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295 Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1290 Pro His Lys Val His Val His |
| 1130   |
| 1145   |
| 1160   |
| 1175   |
| 1190       1195       1200         Tyr Val 1205       Ala Pro His Val Thr 1210       Tyr Gln Asn Ile Ser Thr Asn Leu 1215       Thr Asn Leu 1215         Pro Pro Pro Pro Leu Leu Gly Asn 1225       Ser Thr Gly Ile Asp 1230       Phe Gln Asp 1230         Glu Leu Asp Glu Phe Phe Lys 1235       Asn Val Ser Thr Ser 1245       Ile Pro Asn 1245         Phe Gly Ser Leu Thr Gln Ile 1255       Asn Thr Thr Leu Leu Leu Asp Leu Thr 1260         Tyr Glu Met Leu Ser Leu Gln 1270       Gln Val Val Lys Ala 1275       Leu Asn Glu 1275         Ser Tyr Ile Asp Leu Lys Glu 1285       Leu Gly Asn Tyr Thr Tyr Tyr Asn 1290         Lys Trp Pro Trp Tyr Ile Trp 1300       Leu Gly Phe Ile Ala 1305       Gly Leu Val 1305         Ala Leu Ala Leu Cys Val Phe 1315       Phe Ile Leu Cys Cys Thr Gly Cys 1320         Gly Thr Asp Cys Met Gly Lys 1330       Leu Lys Cys Asn Arg Cys Cys Asp 1335         Arg Tyr Glu Glu Tyr Asp Leu       Glu Pro His Lys Val His Val His   |
| 1210   1215   1210   1215  |
| 1220   1225   1230   |
| 1235       1240       1245         Phe Gly 1250       Ser Leu Thr Gln Ile 1255       Asn Thr Thr Leu Leu Leu Leu Leu 1260       Asp Leu Thr 1260         Tyr Glu 1265       Met Leu Ser Leu Gln 1270       Gln Val Val Lys Ala 1275       Leu Asn Glu 1275         Ser Tyr 1265       Ile Asp Leu Lys Glu 1285       Leu Gly Asn Tyr Thr 1290       Tyr Tyr Asn 1290         Lys Trp 1295       Pro Trp Tyr Ile Trp 1300       Leu Gly Phe Ile Ala 1305       Gly Leu Val 1305         Ala Leu Ala Leu Cys Val Phe 1315       Phe Ile Leu Cys Cys 1320       Thr Gly Cys 1320         Gly Thr Asn Cys Met Gly Lys 1330       Leu Lys Cys Asn Arg 1335       Cys Cys Asp 1335         Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His   |
| Tyr Glu Met Leu Ser Leu Gln Gln Val Val Lys Ala Leu Asn Glu 1265  Ser Tyr Ile Asp Leu Lys Glu Leu Gly Asn Tyr Thr Tyr Tyr Asn 1280  Lys Trp Pro Trp Tyr Ile Trp 1300  Leu Gly Phe Ile Ala Gly Leu Val 1305  Ala Leu Ala Leu Cys Val Phe Phe Ile Leu Cys Cys Thr Gly Cys 1310  Gly Thr Asn Cys Met Gly Lys Leu Lys Cys Asn Arg Cys Cys Asp 1335  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His  |
| Ser Tyr Ile Asp Leu Lys Glu Leu Gly Asn Tyr Thr 1290  Lys Trp Pro Trp Tyr Ile Trp 1300  Leu Gly Phe Ile Ala Gly Leu Val 1305  Ala Leu Ala Leu Cys Val Phe 1315  Gly Thr Asn Cys Met Gly Lys Leu Lys Cys Asn Arg 1325  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His  |
| Lys Trp Pro Trp Tyr Ile Trp Leu Gly Phe Ile Ala Gly Leu Val 1295  Ala Leu Ala Leu Cys Val Phe Phe Ile Leu Cys Cys Thr Gly Cys 1310  Gly Thr Asn Cys Met Gly Lys Leu Lys Cys Asn Arg Cys Cys Asp 1325  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His  |
| 1295 1300 1305  Ala Leu Ala Leu Cys Val Phe Phe Ile Leu Cys Cys Thr Gly Cys 1310  Gly Thr Asn Cys Met Gly Lys Leu Lys Cys Asn Arg Cys Cys Asp 1325  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His  |
| 1310 1315 1320  Gly Thr Asn Cys Met Gly Lys Leu Lys Cys Asn Arg Cys Cys Asp 1325 1330  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His   |
| 1325 1330 1335  Arg Tyr Glu Glu Tyr Asp Leu Glu Pro His Lys Val His Val His  |
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| Val Asp Ile Gln Gln Thr Phe Phe Asp Lys Thr Trp Pro Arg Pro Ile 35 40 45   |
| Asp Val Ser Lys Ala Asp Gly Ile Ile Tyr Pro Gln Gly Arg Thr Tyr 50 55 60   |
| Ser Asn Ile Thr Ile Thr Tyr Gln Gly Leu Phe Pro Tyr Gln Gly Asp 65 70 75 80  |
| His Gly Asp Met Tyr Val Tyr Ser Ala Gly His Ala Thr Gly Thr Thr<br>85 90 95  |

| Pro        | Gln        | Lys        | Leu<br>100 | Phe        | Val        | Ala        | Asn        | Tyr<br>105 | Ser        | Gln        | Asp        | Val        | Lys<br>110 | Gln        | Phe        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Ala        | Asn        | Gly<br>115 | Phe        | Val        | Val        | Arg        | Ile<br>120 | Gly        | Ala        | Ala        | Ala        | Asn<br>125 | Ser        | Thr        | Gly        |
| Thr        | Val<br>130 | Ile        | Ile        | Ser        | Pro        | Ser<br>135 | Thr        | Ser        | Ala        | Thr        | Ile<br>140 | Arg        | Lys        | Ile        | Tyr        |
| Pro<br>145 | Ala        | Phe        | Met        | Leu        | Gly<br>150 | Ser        | Ser        | Val        | Gly        | Asn<br>155 | Phe        | Ser        | Asp        | Gly        | Lys<br>160 |
| Met        | Gly        | Arg        | Phe        | Phe<br>165 | Asn        | His        | Thr        | Leu        | Val<br>170 | Leu        | Leu        | Pro        | Asp        | Gly<br>175 | Сув        |
| Gly        | Thr        | Leu        | Leu<br>180 | Arg        | Ala        | Phe        | Tyr        | Суз<br>185 | Ile        | Leu        | Glu        | Pro        | Arg<br>190 | Ser        | Gly        |
| Asn        | His        | Сув<br>195 | Pro        | Ala        | Gly        | Asn        | Ser<br>200 | Tyr        | Thr        | Ser        | Phe        | Ala<br>205 | Thr        | Tyr        | His        |
| Thr        | Pro<br>210 | Ala        | Thr        | Asp        | CAa        | Ser<br>215 | Asp        | Gly        | Asn        | Tyr        | Asn<br>220 | Arg        | Asn        | Ala        | Ser        |
| Leu<br>225 | Asn        | Ser        | Phe        | Lys        | Glu<br>230 | Tyr        | Phe        | Asn        | Leu        | Arg<br>235 | Asn        | Cys        | Thr        | Phe        | Met<br>240 |
| Tyr        | Thr        | Tyr        | Asn        | Ile<br>245 | Thr        | Glu        | Asp        | Glu        | Ile<br>250 | Leu        | Glu        | Trp        | Phe        | Gly<br>255 | Ile        |
| Thr        | Gln        | Thr        | Ala<br>260 | Gln        | Gly        | Val        | His        | Leu<br>265 | Phe        | Ser        | Ser        | Arg        | Tyr<br>270 | Val        | Asp        |
| Leu        | Tyr        | Gly<br>275 | Gly        | Asn        | Met        | Phe        | Gln<br>280 | Phe        | Ala        | Thr        | Leu        | Pro<br>285 | Val        | Tyr        | Asp        |
| Thr        | Ile<br>290 | Lys        | Tyr        | Tyr        | Ser        | Ile<br>295 | Ile        | Pro        | His        | Ser        | Ile<br>300 | Arg        | Ser        | Ile        | Gln        |
| Ser<br>305 | Asp        | Arg        | ГÀа        | Ala        | Trp<br>310 | Ala        | Ala        | Phe        | Tyr        | Val<br>315 | Tyr        | ГÀЗ        | Leu        | Gln        | Pro<br>320 |
| Leu        | Thr        | Phe        | Leu        | Leu<br>325 | Asp        | Phe        | Ser        | Val        | 330        | Gly        | Tyr        | Ile        | Arg        | Arg<br>335 | Ala        |
| Ile        | Asp        | Cys        | Gly<br>340 | Phe        | Asn        | Asp        | Leu        | Ser<br>345 | Gln        | Leu        | His        | CAa        | Ser<br>350 | Tyr        | Glu        |
| Ser        | Phe        | Asp<br>355 | Val        | Glu        | Ser        | Gly        | Val<br>360 | Tyr        | Ser        | Val        | Ser        | Ser<br>365 | Phe        | Glu        | Ala        |
| Lys        | Pro<br>370 | Ser        | Gly        | Ser        | Val        | Val<br>375 | Glu        | Gln        | Ala        | Glu        | Gly<br>380 | Val        | Glu        | Cys        | Asp        |
| Phe<br>385 | Ser        | Pro        | Leu        | Leu        | Ser<br>390 |            | Thr        | Pro        | Pro        | Gln<br>395 |            | Tyr        | Asn        | Phe        | Lys<br>400 |
| Arg        | Leu        | Val        | Phe        | Thr<br>405 | Asn        | Cys        | Asn        | Tyr        | Asn<br>410 | Leu        | Thr        | Lys        | Leu        | Leu<br>415 | Ser        |
| Leu        | Phe        | Ser        | Val<br>420 | Asn        | Asp        | Phe        | Thr        | Cys<br>425 | Ser        | Gln        | Ile        | Ser        | Pro<br>430 | Ala        | Ala        |
| Ile        | Ala        | Ser<br>435 | Asn        | Cya        | Tyr        | Ser        | Ser<br>440 | Leu        | Ile        | Leu        | Asp        | Tyr<br>445 | Phe        | Ser        | Tyr        |
| Pro        | Leu<br>450 | Ser        | Met        | Lys        | Ser        | Asp<br>455 | Leu        | Ser        | Val        | Ser        | Ser<br>460 | Ala        | Gly        | Pro        | Ile        |
| Ser<br>465 | Gln        | Phe        | Asn        | Tyr        | Lys<br>470 | Gln        | Ser        | Phe        | Ser        | Asn<br>475 | Pro        | Thr        | Сла        | Leu        | Ile<br>480 |
| Leu        | Ala        | Thr        | Val        | Pro<br>485 | His        | Asn        | Leu        | Thr        | Thr<br>490 | Ile        | Thr        | Lys        | Pro        | Leu<br>495 | Lys        |
| Tyr        | Ser        | Tyr        | Ile<br>500 | Asn        | Lys        | Cys        | Ser        | Arg<br>505 | Leu        | Leu        | Ser        | Asp        | Asp<br>510 | Arg        | Thr        |
| Glu        | Val        | Pro        | Gln        | Leu        | Val        | Asn        | Ala        | Asn        | Gln        | Tyr        | Ser        | Pro        | Cys        | Val        | Ser        |

| _          |            | 515        |            |            |            |            | 520        |            |            |            |            | 525        |            |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Ile        | Val<br>530 | Pro        | Ser        | Thr        | Val        | Trp<br>535 | Glu        | Asp        | Gly        | Asp        | Tyr<br>540 | Tyr        | Arg        | Lys        | Gln        |
| Leu<br>545 | Ser        | Pro        | Leu        | Glu        | Gly<br>550 | Gly        | Gly        | Trp        | Leu        | Val<br>555 | Ala        | Ser        | Gly        | Ser        | Thr<br>560 |
| Val        | Ala        | Met        | Thr        | Glu<br>565 | Gln        | Leu        | Gln        | Met        | Gly<br>570 | Phe        | Gly        | Ile        | Thr        | Val<br>575 | Gln        |
| Tyr        | Gly        | Thr        | Asp<br>580 | Thr        | Asn        | Ser        | Val        | Сув<br>585 | Pro        | Lys        | Leu        | Glu        | Phe<br>590 | Ala        | Asn        |
| Asp        | Thr        | Lys<br>595 | Ile        | Ala        | Ser        | Gln        | Leu<br>600 | Gly        | Asn        | Сув        | Val        | Glu<br>605 | Tyr        | Ser        | Leu        |
| Tyr        | Gly<br>610 | Val        | Ser        | Gly        | Arg        | Gly<br>615 | Val        | Phe        | Gln        | Asn        | Cys<br>620 | Thr        | Ala        | Val        | Gly        |
| Val<br>625 | Arg        | Gln        | Gln        | Arg        | Phe<br>630 | Val        | Tyr        | Asp        | Ala        | Tyr<br>635 | Gln        | Asn        | Leu        | Val        | Gly<br>640 |
| Tyr        | Tyr        | Ser        | Asp        | Asp<br>645 | Gly        | Asn        | Tyr        | Tyr        | Сув<br>650 | Leu        | Arg        | Ala        | Cys        | Val<br>655 | Ser        |
| Val        | Pro        | Val        | Ser<br>660 | Val        | Ile        | Tyr        | Asp        | Lys<br>665 | Glu        | Thr        | Lys        | Thr        | His<br>670 | Ala        | Thr        |
| Leu        | Phe        | Gly<br>675 | Ser        | Val        | Ala        | CAa        | Glu<br>680 | His        | Ile        | Ser        | Ser        | Thr<br>685 | Met        | Ser        | Gln        |
| Tyr        | Ser<br>690 | Arg        | Ser        | Thr        | Arg        | Ser<br>695 | Met        | Leu        | Lys        | Arg        | Arg<br>700 | Asp        | Ser        | Thr        | Tyr        |
| Gly<br>705 | Pro        | Leu        | Gln        | Thr        | Pro<br>710 | Val        | Gly        | Cys        | Val        | Leu<br>715 | Gly        | Leu        | Val        | Asn        | Ser<br>720 |
| Ser        | Leu        | Phe        | Val        | Glu<br>725 | Asp        | GÀa        | Lys        | Leu        | Pro<br>730 | Leu        | Gly        | Gln        | Ser        | Leu<br>735 | CÀa        |
| Ala        | Leu        | Pro        | Asp<br>740 | Thr        | Pro        | Ser        | Thr        | Leu<br>745 | Thr        | Pro        | Arg        | Ser        | Val<br>750 | Arg        | Ser        |
| Val        | Pro        | Gly<br>755 | Glu        | Met        | Arg        | Leu        | Ala<br>760 | Ser        | Ile        | Ala        | Phe        | Asn<br>765 | His        | Pro        | Ile        |
| Gln        | Val<br>770 | Asp        | Gln        | Leu        | Asn        | Ser<br>775 | Ser        | Tyr        | Phe        | Lys        | Leu<br>780 | Ser        | Ile        | Pro        | Thr        |
| Asn<br>785 | Phe        | Ser        | Phe        | Gly        | Val<br>790 | Thr        | Gln        | Glu        | Tyr        | Ile<br>795 | Gln        | Thr        | Thr        | Ile        | Gln<br>800 |
| Lys        | Val        | Thr        | Val        | Asp<br>805 | Cys        | Lys        | Gln        | Tyr        | Val<br>810 | Сув        | Asn        | Gly        | Phe        | Gln<br>815 | ГЛа        |
| Cys        | Glu        | Gln        | Leu<br>820 | Leu        | Arg        | Glu        | Tyr        | Gly<br>825 | Gln        | Phe        | CAa        | Ser        | 830        | Ile        | Asn        |
| Gln        | Ala        | Leu<br>835 | His        | Gly        | Ala        | Asn        | Leu<br>840 | Arg        | Gln        | Asp        | Asp        | Ser<br>845 | Val        | Arg        | Asn        |
| Leu        | Phe<br>850 | Ala        | Ser        | Val        | Lys        | Ser<br>855 | Ser        | Gln        | Ser        | Ser        | Pro<br>860 | Ile        | Ile        | Pro        | Gly        |
| Phe<br>865 | Gly        | Gly        | Asp        | Phe        | Asn<br>870 | Leu        | Thr        | Leu        | Leu        | Glu<br>875 | Pro        | Val        | Ser        | Ile        | Ser<br>880 |
| Thr        | Gly        | Ser        | Arg        | Ser<br>885 | Ala        | Arg        | Ser        | Ala        | Ile<br>890 | Glu        | Asp        | Leu        | Leu        | Phe<br>895 | Asp        |
| Lys        | Val        | Thr        | Ile<br>900 | Ala        | Asp        | Pro        | Gly        | Tyr<br>905 | Met        | Gln        | Gly        | Tyr        | Asp<br>910 | Asp        | Cya        |
| Met        | Gln        | Gln<br>915 | Gly        | Pro        | Ala        | Ser        | Ala<br>920 | Arg        | Asp        | Leu        | Ile        | Сув<br>925 | Ala        | Gln        | Tyr        |
| Val        | Ala<br>930 | Gly        | Tyr        | Lys        | Val        | Leu<br>935 | Pro        | Pro        | Leu        | Met        | Asp<br>940 | Val        | Asn        | Met        | Glu        |

| Ala<br>945 | Ala         | Tyr        | Thr   |       | Ser 1<br>950 | Leu L       | eu G        | ly S       |       | le A:<br>55 | la Gly      | / Val | l Gl  | 7 Trp<br>960 |
|------------|-------------|------------|-------|-------|--------------|-------------|-------------|------------|-------|-------------|-------------|-------|-------|--------------|
|            | Ala         | Gly        | Leu   |       |              | Phe A       | la A        | la I       |       |             | ne Ala      | a Glr | n Sei |              |
| Die e      | Ma          | 3          |       | 965   | G] 1         | 7-1 G       | 1 T         |            | 70    | J G.        | 1 17-1      |       | 975   |              |
| Pne        | ıyr         | Arg        | 980   | ASII  | GIY          | /al G       |             | 1e 1<br>35 | nr G  | III G.      | ln Val      | 990   |       | GIU          |
| Asn        | Gln         | 995<br>995 | Leu   | Ile   | Ala A        |             | 000<br>ys 1 | Phe .      | Asn ( | Gln A       |             | eu (  | Gly A | Ala Met      |
| Gln        | Thr<br>1010 | -          | Phe   | Thr   | Thr          | Thr<br>1015 |             | Glu        | Ala   | Phe         | Arg<br>1020 | Lys   | Val   | Gln          |
| Asp        | Ala<br>1025 |            | . Asn | . Asn | Asn          | Ala<br>1030 |             | Ala        | Leu   | Ser         | Lys<br>1035 | Leu   | Ala   | Ser          |
| Glu        | Leu<br>1040 |            | Asn   | Thr   | Phe          | Gly<br>1045 |             | Ile        | Ser   | Ala         | Ser<br>1050 | Ile   | Gly   | Aap          |
| Ile        | Ile<br>1055 |            | Arg   | Leu   | Asp          | Val<br>1060 |             | Glu        | Gln   | Asp         | Ala<br>1065 | Gln   | Ile   | Aap          |
| Arg        | Leu<br>1070 |            | Asn   | Gly   | Arg          | Leu<br>1075 |             | Thr        | Leu   | Asn         | Ala<br>1080 | Phe   | Val   | Ala          |
| Gln        | Gln<br>1085 |            | Val   | Arg   | Ser          | Glu<br>1090 |             | Ala        | Ala   | Leu         | Ser<br>1095 | Ala   | Gln   | Leu          |
| Ala        | Lys<br>1100 |            | Lys   | Val   | Asn          | Glu<br>1105 |             | Val        | Lys   | Ala         | Gln<br>1110 | Ser   | Lys   | Arg          |
| Ser        | Gly<br>1115 |            | Cys   | Gly   | Gln          | Gly<br>1120 |             | His        | Ile   | Val         | Ser<br>1125 | Phe   | Val   | Val          |
| Asn        | Ala<br>1130 |            | Asn   | . Gly | Leu          | Tyr<br>1135 |             | Met        | His   | Val         | Gly<br>1140 | Tyr   | Tyr   | Pro          |
| Ser        | Asn<br>1145 |            | Ile   | Glu   | Val          | Val<br>1150 |             | Ala        | Tyr   | Gly         | Leu<br>1155 | Сув   | Asp   | Ala          |
| Ala        | Asn<br>1160 |            | Thr   | Asn   | . Сув        | Ile<br>1165 |             | Pro        | Val   | Asn         | Gly<br>1170 | Tyr   | Phe   | Ile          |
| Lys        | Thr<br>1175 |            | . Asn | Thr   | Arg          | Ile<br>1180 |             | Asp        | Glu   | Trp         | Ser<br>1185 | Tyr   | Thr   | Gly          |
| Ser        | Ser<br>1190 |            | Tyr   | Ala   | Pro          | Glu<br>1195 |             | Ile        | Thr   | Ser         | Leu<br>1200 | Asn   | Thr   | Lys          |
| Tyr        | Val<br>1205 |            | Pro   | His   | Val          | Thr<br>1210 |             | Gln        | Asn   | Ile         | Ser<br>1215 | Thr   | Asn   | Leu          |
| Pro        | Pro<br>1220 |            | Leu   | Leu   | Gly          | Asn<br>1225 | Ser         | Thr        | Gly   | Ile         | Asp<br>1230 | Phe   | Gln   | Asp          |
| Glu        | Leu<br>1235 |            | Glu   | Phe   | Phe          | Lys<br>1240 |             | Val        | Ser   | Thr         | Ser<br>1245 | Ile   | Pro   | Asn          |
| Phe        | Gly<br>1250 |            | Leu   | Thr   | Gln          | Ile<br>1255 |             | Thr        | Thr   | Leu         | Leu<br>1260 | Asp   | Leu   | Thr          |
| Tyr        | Glu<br>1265 |            | Leu   | . Ser | Leu          | Gln<br>1270 |             | Val        | Val   | ГЛа         | Ala<br>1275 | Leu   | Asn   | Glu          |
| Ser        | Tyr<br>1280 |            | Asp   | Leu   | Lys          | Glu<br>1285 |             | Gly        | Asn   | Tyr         | Thr<br>1290 | Tyr   | Tyr   | Asn          |
| Lys        | Trp<br>1295 |            | Trp   | Tyr   | Ile          | Trp<br>1300 |             | Gly        | Phe   | Ile         | Ala<br>1305 | Gly   | Leu   | Val          |
| Ala        | Leu<br>1310 |            | Leu   | . Сув | Val          | Phe<br>1315 |             | Ile        | Leu   | Cys         | Cys<br>1320 | Thr   | Gly   | Cys          |
| Gly        | Thr<br>1325 |            | . CAa | Met   | Gly          | Lys<br>1330 |             | Lys        | CÀa   | Asn         | Arg<br>1335 | Сла   | СЛа   | Asp          |

| Arg                          | Tyr<br>1340                      |  | ı Glı                       | и Ту:       | r Asl        | 2 Let<br>134 |            | lu Pi      | 10 H:      | is Ly      |            | al I<br>350 | His V      | /al I      | lis        |
|------------------------------|----------------------------------|--|-----------------------------|-------------|--------------|--------------|------------|------------|------------|------------|------------|-------------|------------|------------|------------|
| <211<br><212<br><213<br><220 | L> LE<br>2> TY<br>3> OF<br>0> FE | EQ II<br>ENGTI<br>YPE:<br>RGANI<br>EATUI<br>THER | H: 12<br>PRT<br>ISM:<br>RE: | 255<br>Unki | nown<br>FION | : Hur        | man S      | SARS       | cor        | onav:      | irus       |             |            |            |            |
| < 400                        | )> SI                            | EQUEI  | ICE :                       | 29          |              |              |            |            |            |            |            |             |            |            |            |
| Met<br>1                     | Phe                              | Ile  | Phe                         | Leu<br>5    | Leu          | Phe          | Leu        | Thr        | Leu<br>10  | Thr        | Ser        | Gly         | Ser        | Asp<br>15  | Leu        |
| Asp                          | Arg                              | Cys  | Thr<br>20                   | Thr         | Phe          | Asp          | Asp        | Val<br>25  | Gln        | Ala        | Pro        | Asn         | Tyr<br>30  | Thr        | Gln        |
| His                          | Thr                              | Ser<br>35  | Ser                         | Met         | Arg          | Gly          | Val<br>40  | Tyr        | Tyr        | Pro        | Asp        | Glu<br>45   | Ile        | Phe        | Arg        |
| Ser                          | 50                               | Thr  | Leu                         | Tyr         | Leu          | Thr<br>55    | Gln        | Asp        | Leu        | Phe        | Leu<br>60  | Pro         | Phe        | Tyr        | Ser        |
| Asn<br>65                    | Val                              | Thr  | Gly                         | Phe         | His<br>70    | Thr          | Ile        | Asn        | His        | Thr<br>75  | Phe        | Gly         | Asn        | Pro        | Val<br>80  |
| Ile                          | Pro                              | Phe  | Lys                         | Asp<br>85   | Gly          | Ile          | Tyr        | Phe        | Ala<br>90  | Ala        | Thr        | Glu         | Lys        | Ser<br>95  | Asn        |
| Val                          | Val                              | Arg  | Gly<br>100                  | Trp         | Val          | Phe          | Gly        | Ser<br>105 | Thr        | Met        | Asn        | Asn         | Lys<br>110 | Ser        | Gln        |
| Ser                          | Val                              | Ile<br>115                                       | Ile                         | Ile         | Asn          | Asn          | Ser<br>120 | Thr        | Asn        | Val        | Val        | Ile<br>125  | Arg        | Ala        | CÀa        |
| Asn                          | Phe<br>130                       | Glu  | Leu                         | CAa         | Asp          | Asn<br>135   | Pro        | Phe        | Phe        | Ala        | Val<br>140 | Ser         | Lys        | Pro        | Met        |
| Gly<br>145                   | Thr                              | Gln  | Thr                         | His         | Thr<br>150   | Met          | Ile        | Phe        | Asp        | Asn<br>155 | Ala        | Phe         | Asn        | Сув        | Thr<br>160 |
| Phe                          | Glu                              | Tyr  | Ile                         | Ser<br>165  | Asp          | Ala          | Phe        | Ser        | Leu<br>170 | Asp        | Val        | Ser         | Glu        | Lys<br>175 | Ser        |
| Gly                          | Asn                              | Phe  | Lys<br>180                  | His         | Leu          | Arg          | Glu        | Phe<br>185 | Val        | Phe        | Lys        | Asn         | Lys<br>190 | Asp        | Gly        |
| Phe                          | Leu                              | Tyr<br>195                                       | Val                         | Tyr         | ràa          | Gly          | Tyr<br>200 | Gln        | Pro        | Ile        | Asp        | Val<br>205  | Val        | Arg        | Asp        |
| Leu                          | Pro<br>210                       | Ser  | Gly                         | Phe         | Asn          | Thr<br>215   | Leu        | Lys        | Pro        | Ile        | Phe<br>220 | ГÀа         | Leu        | Pro        | Leu        |
| Gly<br>225                   | Ile                              | Asn  | Ile                         | Thr         | Asn<br>230   | Phe          | Arg        | Ala        | Ile        | Leu<br>235 | Thr        | Ala         | Phe        | Ser        | Pro<br>240 |
| Ala                          | Gln                              | Asp  | Ile                         | Trp<br>245  | Gly          | Thr          | Ser        | Ala        | Ala<br>250 | Ala        | Tyr        | Phe         | Val        | Gly<br>255 | Tyr        |
| Leu                          | Lys                              | Pro  | Thr<br>260                  | Thr         | Phe          | Met          | Leu        | Lys<br>265 | Tyr        | Asp        | Glu        | Asn         | Gly<br>270 | Thr        | Ile        |
| Thr                          | Asp                              | Ala<br>275                                       | Val                         | Asp         | CAa          | Ser          | Gln<br>280 | Asn        | Pro        | Leu        | Ala        | Glu<br>285  | Leu        | Lys        | Cys        |
| Ser                          | Val<br>290                       | ГЛа  | Ser                         | Phe         | Glu          | Ile<br>295   | Asp        | Lys        | Gly        | Ile        | Tyr<br>300 | Gln         | Thr        | Ser        | Asn        |
| Phe<br>305                   | Arg                              | Val  | Val                         | Pro         | Ser<br>310   | Gly          | Asp        | Val        | Val        | Arg<br>315 | Phe        | Pro         | Asn        | Ile        | Thr<br>320 |
| Asn                          | Leu                              | Сув  | Pro                         | Phe<br>325  | Gly          | Glu          | Val        | Phe        | Asn<br>330 | Ala        | Thr        | Lys         | Phe        | Pro<br>335 | Ser        |
| Val                          | Tyr                              | Ala  | Trp<br>340                  | Glu         | Arg          | Lys          | Lys        | Ile<br>345 | Ser        | Asn        | Сув        | Val         | Ala<br>350 | Asp        | Tyr        |
|                              |                                  |  |                             |             |              |              |            |            |            |            |            |             |            |            |            |

| Ser        | Val        | Leu<br>355 | Tyr        | Asn        | Ser        | Thr        | Phe<br>360 | Phe        | Ser        | Thr        | Phe        | Lys<br>365 | CÀa        | Tyr        | Gly        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Val        | Ser<br>370 | Ala        | Thr        | Lys        | Leu        | Asn<br>375 | Asp        | Leu        | CAa        | Phe        | Ser<br>380 | Asn        | Val        | Tyr        | Ala        |
| Asp<br>385 | Ser        | Phe        | Val        | Val        | 390<br>Lys | Gly        | Asp        | Asp        | Val        | Arg<br>395 | Gln        | Ile        | Ala        | Pro        | Gly<br>400 |
| Gln        | Thr        | Gly        | Val        | Ile<br>405 | Ala        | Asp        | Tyr        | Asn        | Tyr<br>410 | Lys        | Leu        | Pro        | Asp        | Asp<br>415 | Phe        |
| Met        | Gly        | Cys        | Val<br>420 | Leu        | Ala        | Trp        | Asn        | Thr<br>425 | Arg        | Asn        | Ile        | Asp        | Ala<br>430 | Thr        | Ser        |
| Thr        | Gly        | Asn<br>435 | Tyr        | Asn        | Tyr        | Lys        | Tyr<br>440 | Arg        | Tyr        | Leu        | Arg        | His<br>445 | Gly        | Lys        | Leu        |
| Arg        | Pro<br>450 | Phe        | Glu        | Arg        | Asp        | Ile<br>455 | Ser        | Asn        | Val        | Pro        | Phe<br>460 | Ser        | Pro        | Asp        | Gly        |
| Lys<br>465 | Pro        | Cha        | Thr        | Pro        | Pro<br>470 | Ala        | Leu        | Asn        | Cys        | Tyr<br>475 | Trp        | Pro        | Leu        | Asn        | Asp<br>480 |
| Tyr        | Gly        | Phe        | Tyr        | Thr<br>485 | Thr        | Thr        | Gly        | Ile        | Gly<br>490 | Tyr        | Gln        | Pro        | Tyr        | Arg<br>495 | Val        |
| Val        | Val        | Leu        | Ser<br>500 | Phe        | Glu        | Leu        | Leu        | Asn<br>505 | Ala        | Pro        | Ala        | Thr        | Val<br>510 | Cys        | Gly        |
| Pro        | Lys        | Leu<br>515 | Ser        | Thr        | Asp        | Leu        | Ile<br>520 | Lys        | Asn        | Gln        | Cys        | Val<br>525 | Asn        | Phe        | Asn        |
| Phe        | Asn<br>530 | Gly        | Leu        | Thr        | Gly        | Thr<br>535 | Gly        | Val        | Leu        | Thr        | Pro<br>540 | Ser        | Ser        | Lys        | Arg        |
| Phe<br>545 | Gln        | Pro        | Phe        | Gln        | Gln<br>550 | Phe        | Gly        | Arg        | Asp        | Val<br>555 | Ser        | Asp        | Phe        | Thr        | Asp<br>560 |
| Ser        | Val        | Arg        | Asp        | Pro<br>565 | ГÀз        | Thr        | Ser        | Glu        | Ile<br>570 | Leu        | Asp        | Ile        | Ser        | Pro<br>575 | Сув        |
| Ser        | Phe        | Gly        | Gly<br>580 | Val        | Ser        | Val        | Ile        | Thr<br>585 | Pro        | Gly        | Thr        | Asn        | Ala<br>590 | Ser        | Ser        |
| Glu        | Val        | Ala<br>595 | Val        | Leu        | Tyr        | Gln        | Asp<br>600 | Val        | Asn        | CAa        | Thr        | Asp<br>605 | Val        | Ser        | Thr        |
| Ala        | Ile<br>610 | His        | Ala        | Asp        | Gln        | Leu<br>615 | Thr        | Pro        | Ala        | Trp        | Arg<br>620 | Ile        | Tyr        | Ser        | Thr        |
| Gly<br>625 | Asn        | Asn        | Val        | Phe        | Gln<br>630 | Thr        | Gln        | Ala        | Gly        | Сув<br>635 | Leu        | Ile        | Gly        | Ala        | Glu<br>640 |
| His        | Val        | Asp        | Thr        | Ser<br>645 | Tyr        | Glu        | Cys        | Asp        | Ile<br>650 | Pro        | Ile        | Gly        | Ala        | Gly<br>655 | Ile        |
| CÀa        | Ala        | Ser        | Tyr<br>660 | His        | Thr        | Val        | Ser        | Leu<br>665 | Leu        | Arg        | Ser        | Thr        | Ser<br>670 | Gln        | Lys        |
| Ser        | Ile        | Val<br>675 | Ala        | Tyr        | Thr        | Met        | Ser<br>680 | Leu        | Gly        | Ala        | Asp        | Ser<br>685 | Ser        | Ile        | Ala        |
| Tyr        | Ser<br>690 | Asn        | Asn        | Thr        | Ile        | Ala<br>695 | Ile        | Pro        | Thr        | Asn        | Phe<br>700 | Ser        | Ile        | Ser        | Ile        |
| Thr<br>705 | Thr        | Glu        | Val        | Met        | Pro<br>710 | Val        | Ser        | Met        | Ala        | Lys<br>715 | Thr        | Ser        | Val        | Asp        | Cys<br>720 |
| Asn        | Met        | Tyr        | Ile        | Сув<br>725 | Gly        | Asp        | Ser        | Thr        | Glu<br>730 | Сув        | Ala        | Asn        | Leu        | Leu<br>735 | Leu        |
| Gln        | Tyr        | Gly        | Ser<br>740 | Phe        | СЛа        | Thr        | Gln        | Leu<br>745 | Asn        | Arg        | Ala        | Leu        | Ser<br>750 | Gly        | Ile        |
| Ala        | Ala        | Glu<br>755 | Gln        | Asp        | Arg        | Asn        | Thr<br>760 | Arg        | Glu        | Val        | Phe        | Ala<br>765 | Gln        | Val        | Lys        |
| Gln        | Met        | Tyr        | Lys        | Thr        | Pro        | Thr        | Leu        | Lys        | Tyr        | Phe        | Gly        | Gly        | Phe        | Asn        | Phe        |

| -continued |
|------------|
| -concinued |

|            | 770         |            |            |            |            | 775          |             |            |            |            | 780         |             |            |            |            |
|------------|-------------|------------|------------|------------|------------|--------------|-------------|------------|------------|------------|-------------|-------------|------------|------------|------------|
| Ser<br>785 |             | Ile        | Leu        | Pro        | Asp<br>790 |              | Leu         | Lys        | Pro        | Thr        |             | Arg         | Ser        | Phe        | Ile<br>800 |
| Glu        | Asp         | Leu        | Leu        | Phe<br>805 | Asn        | Lys          | Val         | Thr        | Leu<br>810 | Ala        | Asp         | Ala         | Gly        | Phe        |            |
| Lys        | Gln         | Tyr        | Gly<br>820 | Glu        | Cys        | Leu          | Gly         | Asp<br>825 | Ile        | Asn        | Ala         | Arg         | Asp<br>830 | Leu        | Ile        |
| Cys        | Ala         | Gln<br>835 | Lys        | Phe        | Asn        | Gly          | Leu<br>840  | Thr        | Val        | Leu        | Pro         | Pro<br>845  | Leu        | Leu        | Thr        |
| Asp        | Asp<br>850  | Met        | Ile        | Ala        | Ala        | Tyr<br>855   | Thr         | Ala        | Ala        | Leu        | Val<br>860  | Ser         | Gly        | Thr        | Ala        |
| Thr<br>865 | Ala         | Gly        | Trp        | Thr        | Phe<br>870 | Gly          | Ala         | Gly        | Ala        | Ala<br>875 | Leu         | Gln         | Ile        | Pro        | Phe<br>880 |
| Ala        | Met         | Gln        | Met        | Ala<br>885 | Tyr        | Arg          | Phe         | Asn        | Gly<br>890 | Ile        | Gly         | Val         | Thr        | Gln<br>895 | Asn        |
| Val        | Leu         | Tyr        | Glu<br>900 | Asn        | Gln        | Lys          | Gln         | Ile<br>905 | Ala        | Asn        | Gln         | Phe         | Asn<br>910 | ГÀа        | Ala        |
| Ile        | Ser         | Gln<br>915 | Ile        | Gln        | Glu        | Ser          | Leu<br>920  | Thr        | Thr        | Thr        | Ser         | Thr<br>925  | Ala        | Leu        | Gly        |
| Lys        | Leu<br>930  | Gln        | Asp        | Val        | Val        | Asn<br>935   | Gln         | Asn        | Ala        | Gln        | Ala<br>940  | Leu         | Asn        | Thr        | Leu        |
| Val<br>945 | Lys         | Gln        | Leu        | Ser        | Ser<br>950 | Asn          | Phe         | Gly        | Ala        | Ile<br>955 | Ser         | Ser         | Val        | Leu        | Asn<br>960 |
| Asp        | Ile         | Leu        | Ser        | Arg<br>965 | Leu        | Asp          | Lys         | Val        | Glu<br>970 | Ala        | Glu         | Val         | Gln        | Ile<br>975 | Aap        |
| Arg        | Leu         | Ile        | Thr<br>980 | Gly        | Arg        | Leu          | Gln         | Ser<br>985 | Leu        | Gln        | Thr         | Tyr         | Val<br>990 | Thr        | Gln        |
| Gln        | Leu         | Ile<br>995 | Arg        | Ala        | Ala        | Glu          | Ile<br>1000 |            | g Al       | a Se       | r Ala       | a As        |            | eu A       | la Ala     |
| Thr        | Lys<br>1010 |            | Sei        | Glu        | ı Cys      | Val<br>101   |             | eu Gi      | ly G       | ln S       | er Ly<br>10 | ys<br>020   | Arg '      | Val        | Asp        |
| Phe        | Cys<br>1025 | -          | / Lys      | Gly        | / Tyr      | His<br>103   |             | eu Me      | et S       | er Pl      | he P:       | ro<br>035   | Gln .      | Ala        | Ala        |
| Pro        | His<br>1040 |            | / Val      | l Val      | l Phe      | Let<br>104   |             | Ls Va      | al T       | hr T       |             | al<br>050   | Pro        | Ser        | Gln        |
| Glu        | Arg<br>1055 |            | n Phe      | e Thi      | Thr        | 106          |             | ro Ai      | la I       | le C       | ys H:<br>10 | is<br>065   | Glu        | Gly        | Lys        |
| Ala        | Tyr<br>1070 |            | e Pro      | Arg        | g Glu      | 1 Gly        |             | al Pl      | ne V       | al P       |             | sn<br>080   | Gly        | Thr        | Ser        |
| Trp        | Phe<br>1085 |            | e Thi      | Glr        | n Arg      | J Asr<br>109 |             | ne Pl      | ne S       | er P       |             | ln<br>095   | Ile        | Ile        | Thr        |
| Thr        | Asp<br>1100 |            | 1 Thi      | : Phe      | e Val      | Sei<br>110   |             | Ly As      | ≅n C       | ys A       |             | al<br>110   | Val        | Ile        | Gly        |
| Ile        | Ile<br>1115 |            | n Asr      | n Thi      | . Val      | 1 Ty:        |             | sp Pi      | ro L       | eu G       |             | ro<br>125   | Glu :      | Leu        | Asp        |
| Ser        | Phe<br>1130 | _          | g Glu      | ı Glu      | ı Lev      | 1 Ası<br>113 | _           | /s Ty      | yr Pl      | he L       | _           | sn<br>140   | His        | Thr        | Ser        |
| Pro        | Asp<br>1145 |            | l Asp      | ) Let      | ı Gly      | / Asp        |             | Le Se      | ∍r G       | ly I       |             | sn .<br>155 | Ala        | Ser        | Val        |
| Val        | Asn<br>1160 |            | e Glr      | ı Lys      | s Glu      | 1 Ile<br>116 |             | ep Ai      | rg L       | eu A       |             | lu<br>170   | Val .      | Ala        | Lys        |
| Asn        | Leu<br>1175 |            | ı Glu      | ı Sei      | . Leu      | 1 Ile<br>118 |             | sp Le      | ∋u G       | ln G       |             | eu<br>185   | Gly :      | Lys        | Tyr        |

#### -continued

Glu Gln Tyr Ile Lys Trp Pro Trp Tyr Val Trp Leu Gly Phe Ile 1190 1195 1200 Ala Gly Leu Ile Ala Ile Val Met Val Thr Ile Leu Leu Cys Cys 1210 Met Thr Ser Cys Cys Ser Cys Leu Lys Gly Ala Cys Ser Cys Gly 1225 Ser Cys Cys Lys Phe Asp Glu Asp Asp Ser Glu Pro Val Leu Lys 1240 Gly Val Lys Leu His Tyr Thr <210> SEQ ID NO 30 <211> LENGTH: 1353 <212> TYPE: PRT <213 > ORGANISM: Human coronavirus <400> SEQUENCE: 30 Met Phe Leu Ile Leu Leu Ile Ser Leu Pro Thr Ala Phe Ala Val Ile Gly Asp Leu Lys Cys Thr Ser Asp Asn Ile Asn Asp Lys Asp Thr Gly Pro Pro Pro Ile Ser Thr Asp Thr Val Asp Val Thr Asn Gly Leu Gly 40 Thr Tyr Tyr Val Leu Asp Arg Val Tyr Leu Asn Thr Thr Leu Phe Leu Asn Gly Tyr Tyr Pro Thr Ser Gly Ser Thr Tyr Arg Asn Met Ala Leu Lys Gly Ser Val Leu Leu Ser Arg Leu Trp Phe Lys Pro Pro Phe Leu Ser Asp Phe Ile Asn Gly Ile Phe Ala Lys Val Lys Asn Thr Lys Val 105 Ile Lys Asp Arg Val Met Tyr Ser Glu Phe Pro Ala Ile Thr Ile Gly 120 Ser Thr Phe Val Asn Thr Ser Tyr Ser Val Val Val Gln Pro Arg Thr Ile Asn Ser Thr Gln Asp Gly Asp Asn Lys Leu Gln Gly Leu Leu Glu 155 Val Ser Val Cys Gln Tyr Asn Met Cys Glu Tyr Pro Gln Thr Ile Cys His Pro Asn Leu Gly Asn His Arg Lys Glu Leu Trp His Leu Asp Thr Gly Val Val Ser Cys Leu Tyr Lys Arg Asn Phe Thr Tyr Asp Val Asn Ala Asp Tyr Leu Tyr Phe His Phe Tyr Gln Glu Gly Gly Thr Phe Tyr Ala Tyr Phe Thr Asp Thr Gly Val Val Thr Lys Phe Leu Phe Asn Val 230 Tyr Leu Gly Met Ala Leu Ser His Tyr Tyr Val Met Pro Leu Thr Cys 250 Asn Ser Lys Leu Thr Leu Glu Tyr Trp Val Thr Pro Leu Thr Ser Arg Gln Tyr Leu Leu Ala Phe Asn Gln Asp Gly Ile Ile Phe Asn Ala Glu Asp Cys Met Ser Asp Phe Met Ser Glu Ile Lys Cys Lys Thr Gln Ser

| _          | 290        |            |            |            |            | 295        |            |            |            |            | 300        |            |            |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Ile<br>305 |            | Pro        | Pro        | Thr        | Gly<br>310 |            | Tyr        | Glu        | Leu        | Asn<br>315 |            | Tyr        | Thr        | Val        | Gln<br>320 |
| Pro        | Ile        | Ala        | Asp        | Val<br>325 | Tyr        | Arg        | Arg        | Lys        | Pro<br>330 | Asn        | Leu        | Pro        | Asn        | Сув<br>335 | Asn        |
| Ile        | Glu        | Ala        | Trp<br>340 | Leu        | Asn        | Asp        | Lys        | Ser<br>345 | Val        | Pro        | Ser        | Pro        | Leu<br>350 | Asn        | Trp        |
| Glu        | Arg        | Lys<br>355 | Thr        | Phe        | Ser        | Asn        | Сув<br>360 | Asn        | Phe        | Asn        | Met        | Ser<br>365 | Ser        | Leu        | Met        |
| Ser        | Phe<br>370 | Ile        | Gln        | Ala        | Asp        | Ser<br>375 | Phe        | Thr        | Cys        | Asn        | Asn<br>380 | Ile        | Asp        | Ala        | Ala        |
| 385<br>Lys | Ile        | Tyr        | Gly        | Met        | Cys        | Phe        | Ser        | Ser        | Ile        | Thr<br>395 | Ile        | Asp        | Lys        | Phe        | Ala<br>400 |
| Ile        | Pro        | Asn        | Gly        | Arg<br>405 | Lys        | Val        | Asp        | Leu        | Gln<br>410 | Leu        | Gly        | Asn        | Leu        | Gly<br>415 | Tyr        |
| Leu        | Gln        | Ser        | Phe<br>420 | Asn        | Tyr        | Arg        | Ile        | Asp<br>425 | Thr        | Thr        | Ala        | Thr        | Ser<br>430 | Cys        | Gln        |
| Leu        | Tyr        | Tyr<br>435 | Asn        | Leu        | Pro        | Ala        | Ala<br>440 | Asn        | Val        | Ser        | Val        | Ser<br>445 | Arg        | Phe        | Asn        |
| Pro        | Ser<br>450 | Thr        | Trp        | Asn        | Lys        | Arg<br>455 | Phe        | Gly        | Phe        | Ile        | Glu<br>460 | Asp        | Ser        | Val        | Phe        |
| Lys<br>465 | Pro        | Arg        | Pro        | Ala        | Gly<br>470 | Val        | Leu        | Thr        | Asn        | His<br>475 | Asp        | Val        | Val        | Tyr        | Ala<br>480 |
| Gln        | His        | Сув        | Phe        | Lys<br>485 | Ala        | Pro        | Lys        | Asn        | Phe<br>490 | Сув        | Pro        | CAa        | Lys        | Leu<br>495 | Asn        |
| Gly        | Ser        | Сув        | Val<br>500 | Gly        | Ser        | Gly        | Pro        | Gly<br>505 | Lys        | Asn        | Asn        | Gly        | Ile<br>510 | Gly        | Thr        |
| CAa        | Pro        | Ala<br>515 | Gly        | Thr        | Asn        | Tyr        | Leu<br>520 | Thr        | Cys        | Asp        | Asn        | Leu<br>525 | Cys        | Thr        | Pro        |
| Asp        | Pro<br>530 | Ile        | Thr        | Phe        | Thr        | Gly<br>535 | Thr        | Tyr        | Lys        | Сув        | Pro<br>540 | Gln        | Thr        | Lys        | Ser        |
| Leu<br>545 | Val        | Gly        | Ile        | Gly        | Glu<br>550 | His        | Cys        | Ser        | Gly        | Leu<br>555 | Ala        | Val        | Lys        | Ser        | Asp<br>560 |
| Tyr        | Сув        | Gly        | Gly        | Asn<br>565 | Ser        | Cys        | Thr        | Cys        | Arg<br>570 | Pro        | Gln        | Ala        | Phe        | Leu<br>575 | Gly        |
| Trp        | Ser        | Ala        | Asp<br>580 | Ser        | Cys        | Leu        | Gln        | Gly<br>585 | Asp        | Lys        | Cys        | Asn        | Ile<br>590 | Phe        | Ala        |
| Asn        | Phe        | Ile<br>595 | Leu        | His        | Asp        | Val        | Asn<br>600 | Ser        | Gly        | Leu        | Thr        | 605        | Ser        | Thr        | Asp        |
| Leu        | Gln<br>610 | Lys        | Ala        | Asn        | Thr        | Asp<br>615 | Ile        | Ile        | Leu        | Gly        | Val<br>620 | CAa        | Val        | Asn        | Tyr        |
| Asp<br>625 | Leu        | Tyr        | Gly        | Ile        | Leu<br>630 | Gly        | Gln        | Gly        | Ile        | Phe<br>635 | Val        | Glu        | Val        | Asn        | Ala<br>640 |
| Thr        | Tyr        | Tyr        | Asn        | Ser<br>645 | Trp        | Gln        | Asn        | Leu        | Leu<br>650 | Tyr        | Asp        | Ser        | Asn        | Gly<br>655 | Asn        |
| Leu        | Tyr        | Gly        | Phe<br>660 | Arg        | Asp        | Tyr        | Ile        | Ile<br>665 | Asn        | Arg        | Thr        | Phe        | Met<br>670 | Ile        | Arg        |
| Ser        | Сув        | Tyr<br>675 | Ser        | Gly        | Arg        | Val        | Ser<br>680 | Ala        | Ala        | Phe        | His        | Ala<br>685 | Asn        | Ser        | Ser        |
| Glu        | Pro<br>690 | Ala        | Leu        | Leu        | Phe        | Arg<br>695 | Asn        | Ile        | Lys        | Сув        | Asn<br>700 | Tyr        | Val        | Phe        | Asn        |
| Asn<br>705 | Ser        | Leu        | Thr        | Arg        | Gln<br>710 | Leu        | Gln        | Pro        | Ile        | Asn<br>715 | Tyr        | Phe        | Asp        | Ser        | Tyr<br>720 |

| Leu        | Gly         | Cys        | Val        | Val<br>725 | Asn        | Ala        | Tyr         | Asn        | Ser<br>730 | Thr        | Ala        | Ile        | Ser        | Val<br>735       | Gln        |
|------------|-------------|------------|------------|------------|------------|------------|-------------|------------|------------|------------|------------|------------|------------|------------------|------------|
| Thr        | Cya         | Asp        | Leu<br>740 |            | Val        | Gly        | Ser         | Gly<br>745 |            | Сла        | Val        | Asp        | Tyr<br>750 |                  | ГЛа        |
| Asn        | Arg         | Arg<br>755 | Ser        | Arg        | Gly        | Ala        | Ile<br>760  | Thr        | Thr        | Gly        | Tyr        | Arg        |            | Thr              | Asn        |
| Phe        | Glu<br>770  | Pro        | Phe        | Thr        | Val        | Asn<br>775 | Ser         | Val        | Asn        | Asp        | Ser<br>780 | Leu        | Glu        | Pro              | Val        |
| Gly<br>785 | Gly         | Leu        | Tyr        | Glu        | Ile<br>790 | Gln        | Ile         | Pro        | Ser        | Glu<br>795 | Phe        | Thr        | Ile        | Gly              | Asn<br>800 |
| Met        | Val         | Glu        | Phe        | Ile<br>805 | Gln        | Thr        | Ser         | Ser        | Pro<br>810 | Lys        | Val        | Thr        | Ile        | Asp<br>815       | Cys        |
| Ala        | Ala         | Phe        | Val<br>820 | CÀa        | Gly        | Asp        | Tyr         | Ala<br>825 | Ala        | СЛа        | rys        | Ser        | Gln<br>830 | Leu              | Val        |
| Glu        | Tyr         | Gly<br>835 | Ser        | Phe        | CAa        | Asp        | Asn<br>840  | Ile        | Asn        | Ala        | Ile        | Leu<br>845 |            | Glu              | Val        |
| Asn        | Glu<br>850  | Leu        | Leu        | Asp        | Thr        | Thr<br>855 | Gln         | Leu        | Gln        | Val        | Ala<br>860 | Asn        | Ser        | Leu              | Met        |
| Asn<br>865 | Gly         | Val        | Thr        | Leu        | Ser<br>870 | Thr        | Lys         | Leu        | Lys        | Asp<br>875 | Gly        | Val        | Asn        | Phe              | Asn<br>880 |
| Val        | Asp         | Asp        | Ile        | Asn<br>885 | Phe        | Ser        | Pro         | Val        | Leu<br>890 | Gly        | CÀa        | Leu        | Gly        | Ser<br>895       | Glu        |
| CÀa        | Ser         | ГÀа        | Ala<br>900 | Ser        | Ser        | Arg        | Ser         | Ala<br>905 | Ile        | Glu        | Asp        | Leu        | Leu<br>910 | Phe              | Asp        |
| ГÀз        | Val         | Lys<br>915 | Leu        | Ser        | Asp        | Val        | Gly<br>920  | Phe        | Val        | Glu        | Ala        | Tyr<br>925 |            | Asn              | СЛа        |
| Thr        | Gly<br>930  | Gly        | Ala        | Glu        | Ile        | Arg<br>935 | Asp         | Leu        | Ile        | СЛа        | Val<br>940 | Gln        | Ser        | Tyr              | ГЛЗ        |
| Gly<br>945 | Ile         | ГÀз        | Val        | Leu        | Pro<br>950 | Pro        | Leu         | Leu        | Ser        | Glu<br>955 | Asn        | Gln        | Ile        | Ser              | Gly<br>960 |
| Tyr        | Thr         | Leu        | Ala        | Ala<br>965 | Thr        | Ser        | Ala         | Ser        | Leu<br>970 | Phe        | Pro        | Pro        | Trp        | Thr<br>975       | Ala        |
| Ala        | Ala         | Gly        | Val<br>980 | Pro        | Phe        | Tyr        | Leu         | Asn<br>985 | Val        | Gln        | Tyr        | Arg        | Ile<br>990 | Asn              | Gly        |
| Leu        | Gly         | Val<br>995 | Thr        | Met        | Asp        | Val        | Leu<br>1000 |            | Glı        | n Ası      | n Gl       | n Ly<br>10 |            | eu I             | le Ala     |
| Asn        | Ala<br>1010 |            | e Asr      | n Asr      | n Ala      | Let<br>101 |             | /r A       | La I       | le G       |            | lu<br>020  | Gly        | Phe I            | Aap        |
| Ala        | Thr<br>1025 |            | n Ser      | Ala        | ı Lev      | 103        | _           | /s I.      | Le G       | ln A       |            | al<br>035  | Val .      | Asn i            | Ala        |
| Asn        | Ala<br>1040 |            | ı Ala      | ı Lev      | ı Asr      | Asr<br>104 |             | eu Le      | eu G       | ln G       |            | eu<br>050  | Ser .      | Asn i            | Arg        |
| Phe        | Gly<br>1055 |            | a Il∈      | e Ser      | Ala        | Sei<br>106 |             | eu G       | ln G       | lu I       |            | eu<br>065  | Ser .      | Arg 1            | Leu        |
| Asp        | Ala<br>1070 |            | ı Glu      | ı Ala      | a Glu      | 107        |             | ln I       | Le As      | ep A:      | -          | eu<br>080  | Ile .      | Asn (            | Gly        |
| Arg        | Leu<br>1085 |            | Ala        | ı Lev      | ı Asr      | 109        | -           | yr Va      | al Se      | er G       |            | ln<br>095  | Leu        | Ser A            | Aap        |
| Ser        | Thr         |            | ı Val      | . Lys      | Phe        | Se:        |             | La A       | La G       | ln A       |            | et<br>110  | Glu :      | Lys <sup>1</sup> | Val        |
| Asn        | Glu<br>1115 |            | val        | . Lys      | Ser        | Glr<br>112 |             | er Se      | er Ai      | rg I       |            | sn<br>125  | Phe        | Cys (            | Gly        |

| Asn          | Gly<br>1130                      |             | His         | Il∈       | · Ile     | Ser<br>113 |            | ∋u V       | al (      | Gln        | Asn  | Ala<br>1140 |      | ro         | Tyr       | Gly       |
|--------------|----------------------------------|-------------|-------------|-----------|-----------|------------|------------|------------|-----------|------------|------|-------------|------|------------|-----------|-----------|
| Leu          | Tyr<br>1145                      |             | lle         | His       | Phe       | Ser<br>115 |            | yr V       | al I      | Pro        | Thr  | Lys<br>1155 |      | yr         | Val       | Thr       |
| Ala          | Arg<br>1160                      |             | Ser         | Pro       | Gly       | Leu<br>116 |            | ys I       | le A      | Ala        | Gly  | Asp<br>1170 |      | rg         | Gly       | Ile       |
| Ala          | Pro<br>1175                      |             | Ser         | Gly       | Tyr       | Ph∈        |            | al A       | sn 7      | Val        | Asn  | Asn<br>1185 |      | hr         | Trp       | Met       |
| Tyr          | Thr<br>1190                      |             | Ser         | Gly       | Tyr       | Туг<br>119 |            | yr P       | ro (      | Glu        | Pro  | Ile<br>1200 |      | hr         | Glu       | Asn       |
| Asn          | Val<br>1205                      |             | . Val       | Met       | Ser       | Thr<br>121 |            | ys A       | la V      | Val        | Asn  | Tyr<br>1215 |      | hr         | Lys       | Ala       |
| Pro          | Tyr<br>1220                      |             | . Met       | Leu       | . Asn     | Thr<br>122 |            | ∍r I       | le 1      | Pro        | Asn  | Leu<br>1230 |      | ro         | Asp       | Phe       |
| ГÀа          | Glu<br>1235                      |             | Leu         | Asp       | Gln       | Trp<br>124 |            | ne L       | Aa 1      | Asn        | Gln  | Thr<br>1245 |      | er         | Val       | Ala       |
| Pro          | Asp<br>1250                      |             | Ser         | Leu       | . Asp     | Tyr<br>125 |            | le A       | sn \      | Val        | Thr  | Phe<br>1260 |      | eu         | Asp       | Leu       |
| Gln          | Val<br>1265                      |             | Met         | Asr       | Arg       | Leu<br>127 |            | ln G       | lu 2      | Ala        | Ile  | Lys<br>1275 |      | al         | Leu       | Asn       |
| Gln          | Ser<br>1280                      |             | Ile         | Asr       | Leu       | Lys<br>128 |            | ab I       | le (      | Gly        | Thr  | Tyr<br>1290 |      | lu         | Tyr       | Tyr       |
| Val          | Lys<br>1295                      |             | Pro         | Trp       | Tyr       | Val        |            | rp L       | eu 1      | Leu        | Ile  | 1305<br>Cys |      | eu         | Ala       | Gly       |
| Val          | Ala<br>1310                      |             | Leu         | Val       | . Leu     | Leu<br>131 |            | ne P       | he :      | Ile        | CAa  | Cys<br>1320 |      | уs         | Thr       | Gly       |
| CAa          | Gly<br>1325                      |             | Ser         | Cys       | Phe       | Lys<br>133 |            | Aa C       | ys (      | Gly        | Gly  | 1335<br>Cys |      | ys         | Asp       | Asp       |
| Tyr          | Thr<br>1340                      |             | Tyr         | Gln       | ı Glu     | Leu<br>134 |            | al I       | le 1      | ГÀа        | Thr  | Ser<br>1350 |      | is         | Asp       | Asp       |
| <211<br><212 | )> SE<br>L> LE<br>2> TY<br>3> OR | NGTH<br>PE: | : 13<br>PRT | 51        | ın cc     | rona       | ıvirı      | ıs         |           |            |      |             |      |            |           |           |
| < 400        | )> SE                            | QUEN        | ICE:        | 31        |           |            |            |            |           |            |      |             |      |            |           |           |
| Met<br>1     | Phe                              | Leu         | Ile         | Ile<br>5  | Phe       | Ile        | Leu        | Pro        | Th:<br>10 | r Th       | ır L | eu Al       | la ' | Val        | Ile<br>15 | Gly       |
| Asp          | Phe                              | Asn         | Cys<br>20   | Thr       | Asn       | Ser        | Phe        | Ile<br>25  | Ası       | n As       | эр Т | yr As       |      | go<br>Pàs  | Thr       | Ile       |
| Pro          | Arg                              | Ile<br>35   | Ser         | Glu       | Asp       | Val        | Val<br>40  | Asp        | Va:       | l S∈       | er L | eu G]<br>45 |      | Leu        | Gly       | Thr       |
| Tyr          | Tyr<br>50                        | Val         | Leu         | Asn       | Arg       | Val<br>55  | Tyr        | Leu        | Ası       | n Th       | nr T |             | eu I | Leu        | Phe       | Thr       |
| Gly<br>65    | Tyr                              | Phe         | Pro         | Lys       | Ser<br>70 | Gly        | Ala        | Asn        | Phe       | e Ar<br>75 |      | sp Le       | eu I | Ala        | Leu       | Lys<br>80 |
| Gly          | Ser                              | Ile         | Tyr         | Leu<br>85 | Ser       | Thr        | Leu        | Trp        | Ту:<br>90 | r Ly       | s P  | ro Pı       | ro 1 | Phe        | Leu<br>95 | . Ser     |
| Asp          | Phe                              | Asn         | Asn<br>100  | Gly       | Ile       | Phe        | Ser        | Lys<br>105 | Va:       | l Ly       | /s A | sn Th       |      | նչs<br>110 |           | ı Tyr     |
| Val          | Asn                              | Asn<br>115  | Thr         | Leu       | Tyr       | Ser        | Glu<br>120 | Phe        | Sei       | r Th       | nr I | le Va       |      | Ile        | Gly       | Ser       |
| Val          | Phe<br>130                       | Val         | Asn         | Thr       | Ser       | Tyr<br>135 | Thr        | Ile        | Va:       | l Va       |      | ln Pi<br>40 | :o 1 | His        | Asr       | Gly       |

| Ile<br>145 | Leu        | Glu        | Ile        | Thr        | Ala<br>150 | Сув        | Gln        | Tyr        | Thr        | Met<br>155 | СЛа        | Glu        | Tyr        | Pro        | His<br>160 |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Thr        | Val        | Сув        | Lys        | Ser<br>165 | Lys        | Gly        | Ser        | Ile        | Arg<br>170 | Asn        | Glu        | Ser        | Trp        | His<br>175 | Ile        |
| Asp        | Ser        | Ser        | Glu<br>180 | Pro        | Leu        | Cys        | Leu        | Phe<br>185 | Lys        | Lys        | Asn        | Phe        | Thr<br>190 | Tyr        | Asn        |
| Val        | Ser        | Ala<br>195 | Asp        | Trp        | Leu        | Tyr        | Phe<br>200 | His        | Phe        | Tyr        | Gln        | Glu<br>205 | Arg        | Gly        | Val        |
| Phe        | Tyr<br>210 | Ala        | Tyr        | Tyr        | Ala        | Asp<br>215 | Val        | Gly        | Met        | Pro        | Thr<br>220 | Thr        | Phe        | Leu        | Phe        |
| Ser<br>225 | Leu        | Tyr        | Leu        | Gly        | Thr<br>230 | Ile        | Leu        | Ser        | His        | Tyr<br>235 | Tyr        | Val        | Met        | Pro        | Leu<br>240 |
| Thr        | Сув        | Asn        | Ala        | Ile<br>245 | Ser        | Ser        | Asn        | Thr        | Asp<br>250 | Asn        | Glu        | Thr        | Leu        | Glu<br>255 | Tyr        |
| Trp        | Val        | Thr        | Pro<br>260 | Leu        | Ser        | Arg        | Arg        | Gln<br>265 | Tyr        | Leu        | Leu        | Asn        | Phe<br>270 | Asp        | Glu        |
| His        | Gly        | Val<br>275 | Ile        | Thr        | Asn        | Ala        | Val<br>280 | Asp        | Càa        | Ser        | Ser        | Ser<br>285 | Phe        | Leu        | Ser        |
| Glu        | Ile<br>290 | Gln        | CÀa        | Lys        | Thr        | Gln<br>295 | Ser        | Phe        | Ala        | Pro        | Asn<br>300 | Thr        | Gly        | Val        | Tyr        |
| Asp<br>305 | Leu        | Ser        | Gly        | Phe        | Thr<br>310 | Val        | Lys        | Pro        | Val        | Ala<br>315 | Thr        | Val        | Tyr        | Arg        | Arg<br>320 |
| Ile        | Pro        | Asn        | Leu        | Pro<br>325 | Asp        | CÀa        | Aap        | Ile        | 330<br>Asp | Asn        | Trp        | Leu        | Asn        | Asn<br>335 | Val        |
| Ser        | Val        | Pro        | Ser<br>340 | Pro        | Leu        | Asn        | Trp        | Glu<br>345 | Arg        | Arg        | Ile        | Phe        | Ser<br>350 | Asn        | Cys        |
| Asn        | Phe        | Asn<br>355 | Leu        | Ser        | Thr        | Leu        | Leu<br>360 | Arg        | Leu        | Val        | His        | Val<br>365 | Asp        | Ser        | Phe        |
| Ser        | Cys<br>370 | Asn        | Asn        | Leu        | Asp        | Lys<br>375 | Ser        | Lys        | Ile        | Phe        | Gly<br>380 | Ser        | Cya        | Phe        | Asn        |
| Ser<br>385 | Ile        | Thr        | Val        | Asp        | 390<br>Lys | Phe        | Ala        | Ile        | Pro        | Asn<br>395 | Arg        | Arg        | Arg        | Asp        | Asp<br>400 |
| Leu        | Gln        | Leu        | Gly        | Ser<br>405 | Ser        | Gly        | Phe        | Leu        | Gln<br>410 | Ser        | Ser        | Asn        | Tyr        | Lys<br>415 | Ile        |
| Asp        | Ile        | Ser        | Ser<br>420 | Ser        | Ser        | Cys        | Gln        | Leu<br>425 | Tyr        | Tyr        | Ser        | Leu        | Pro<br>430 | Leu        | Val        |
| Asn        | Val        | Thr<br>435 | Ile        | Asn        | Asn        | Phe        | Asn<br>440 | Pro        | Ser        | Ser        | Trp        | Asn<br>445 | Arg        | Arg        | Tyr        |
| Gly        | Phe<br>450 | Gly        | Ser        | Phe        | Asn        | Leu<br>455 | Ser        | Ser        | Tyr        | Asp        | Val<br>460 | Val        | Tyr        | Ser        | Asp        |
| His<br>465 | Cya        | Phe        | Ser        | Val        | Asn<br>470 | Ser        | Aap        | Phe        | Cys        | Pro<br>475 | CÀa        | Ala        | Asp        | Pro        | Ser<br>480 |
| Val        | Val        | Asn        | Ser        | Cys<br>485 | Ala        | Lys        | Ser        | Lys        | Pro<br>490 | Pro        | Ser        | Ala        | Ile        | Cys<br>495 | Pro        |
| Ala        | Gly        | Thr        | 500        | Tyr        | Arg        | His        | Cys        | Asp<br>505 | Leu        | Asp        | Thr        | Thr        | Leu<br>510 | Tyr        | Val        |
| ГЛа        | Asn        | Trp<br>515 | CÀa        | Arg        | CAa        | Ser        | Cys<br>520 | Leu        | Pro        | Asp        | Pro        | Ile<br>525 | Ser        | Thr        | Tyr        |
| Ser        | Pro<br>530 | Asn        | Thr        | Сла        | Pro        | Gln<br>535 | Lys        | Lys        | Val        | Val        | Val<br>540 | Gly        | Ile        | Gly        | Glu        |
| His<br>545 | Сув        | Pro        | Gly        | Leu        | Gly<br>550 | Ile        | Asn        | Glu        | Glu        | Lуз<br>555 | СЛа        | Gly        | Thr        | Gln        | Leu<br>560 |

| Asn        | His        | Ser        | Ser        | Сув<br>565 | Phe        | Cys        | Ser        | Pro        | Asp<br>570 | Ala        | Phe        | Leu        | Gly        | Trp<br>575 | Ser        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Phe        | Asp        | Ser        | 580        | Ile        | Ser        | Asn        | Asn        | Arg<br>585 | Cys        | Asn        | Ile        | Phe        | Ser<br>590 | Asn        | Phe        |
| Ile        | Phe        | Asn<br>595 | Gly        | Ile        | Asn        | Ser        | Gly<br>600 | Thr        | Thr        | Суз        | Ser        | Asn<br>605 | Asp        | Leu        | Leu        |
| Tyr        | Ser<br>610 | Asn        | Thr        | Glu        | Ile        | Ser<br>615 | Thr        | Gly        | Val        | Суз        | Val<br>620 | Asn        | Tyr        | Asp        | Leu        |
| Tyr<br>625 | Gly        | Ile        | Thr        | Gly        | Gln<br>630 | Gly        | Ile        | Phe        | Lys        | Glu<br>635 | Val        | Ser        | Ala        | Ala        | Tyr<br>640 |
| Tyr        | Asn        | Asn        | Trp        | Gln<br>645 | Asn        | Leu        | Leu        | Tyr        | Asp<br>650 | Ser        | Asn        | Gly        | Asn        | Ile<br>655 | Ile        |
| Gly        | Phe        | ГÀа        | Asp        | Phe        | Leu        | Thr        | Asn        | Lys        | Thr        | Tyr        | Thr        | Ile        | Leu<br>670 | Pro        | CAa        |
| Tyr        | Ser        | Gly<br>675 | Arg        | Val        | Ser        | Ala        | Ala<br>680 | Phe        | Tyr        | Gln        | Asn        | Ser<br>685 | Ser        | Ser        | Pro        |
| Ala        | Leu<br>690 | Leu        | Tyr        | Arg        | Asn        | Leu<br>695 | ГЛа        | Cys        | Ser        | Tyr        | Val<br>700 | Leu        | Asn        | Asn        | Ile        |
| Ser<br>705 | Phe        | Ile        | Ser        | Gln        | Pro<br>710 | Phe        | Tyr        | Phe        | Asp        | Ser<br>715 | Tyr        | Leu        | Gly        | CÀa        | Val<br>720 |
| Leu        | Asn        | Ala        | Val        | Asn<br>725 | Leu        | Thr        | Ser        | Tyr        | Ser<br>730 | Val        | Ser        | Ser        | Cha        | Asp<br>735 | Leu        |
| Arg        | Met        | Gly        | Ser<br>740 | Gly        | Phe        | Cys        | Ile        | Asp<br>745 | Tyr        | Ala        | Leu        | Pro        | Ser<br>750 | Ser        | Arg        |
| Arg        | Lys        | Arg<br>755 | Arg        | Gly        | Ile        | Ser        | Ser<br>760 | Pro        | Tyr        | Arg        | Phe        | Val<br>765 | Thr        | Phe        | Glu        |
| Pro        | Phe<br>770 | Asn        | Val        | Ser        | Phe        | Val<br>775 | Asn        | Asp        | Ser        | Val        | Glu<br>780 | Thr        | Val        | Gly        | Gly        |
| Leu<br>785 | Phe        | Glu        | Ile        | Gln        | Ile<br>790 | Pro        | Thr        | Asn        | Phe        | Thr<br>795 | Ile        | Ala        | Gly        | His        | Glu<br>800 |
| Glu        | Phe        | Ile        | Gln        | Thr<br>805 | Ser        | Ser        | Pro        | Lys        | Val<br>810 | Thr        | Ile        | Asp        | Cys        | Ser<br>815 | Ala        |
| Phe        | Val        | Cys        | Ser<br>820 | Asn        | Tyr        | Ala        | Ala        | Сув<br>825 | His        | Asp        | Leu        | Leu        | Ser<br>830 | Glu        | Tyr        |
| Gly        | Thr        | Phe<br>835 | Cys        | Asp        | Asn        | Ile        | Asn<br>840 | Ser        | Ile        | Leu        | Asn        | Glu<br>845 | Val        | Asn        | Asp        |
| Leu        | Leu<br>850 | Asp        | Ile        | Thr        | Gln        | Leu<br>855 |            | Val        | Ala        | Asn        | Ala<br>860 | Leu        | Met        | Gln        | Gly        |
| Val<br>865 | Thr        | Leu        | Ser        | Ser        | Asn<br>870 | Leu        | Asn        | Thr        | Asn        | Leu<br>875 | His        | Ser        | Asp        | Val        | Asp        |
| Asn        | Ile        | Asp        | Phe        | Lys<br>885 | Ser        | Leu        | Leu        | Gly        | 690        | Leu        | Gly        | Ser        | Gln        | Cys<br>895 | Gly        |
| Ser        | Ser        | Ser        | Arg<br>900 | Ser        | Leu        | Leu        | Glu        | Asp<br>905 | Leu        | Leu        | Phe        | Asn        | Lys<br>910 | Val        | ГÀа        |
| Leu        | Ser        | Asp<br>915 | Val        | Gly        | Phe        | Val        | Glu<br>920 | Ala        | Tyr        | Asn        | Asn        | Сув<br>925 | Thr        | Gly        | Gly        |
| Ser        | Glu<br>930 | Ile        | Arg        | Asp        | Leu        | Leu<br>935 | Cys        | Val        | Gln        | Ser        | Phe<br>940 | Asn        | Gly        | Ile        | ГХз        |
| Val<br>945 | Leu        | Pro        | Pro        | Ile        | Leu<br>950 | Ser        | Glu        | Thr        | Gln        | Ile<br>955 | Ser        | Gly        | Tyr        | Thr        | Thr<br>960 |
| Ala        | Ala        | Thr        | Val        | Ala<br>965 | Ala        | Met        | Phe        | Pro        | Pro<br>970 | Trp        | Ser        | Ala        | Ala        | Ala<br>975 | Gly        |
| Val        | Pro        | Phe        | Ser        | Leu        | Asn        | Val        | Gln        | Tyr        | Arg        | Ile        | Asn        | Gly        | Leu        | Gly        | Val        |

|                 | 980         | 985                   | 990                                 |
|-----------------|-------------|-----------------------|-------------------------------------|
| Thr Met Asp 995 | Val Leu Asn | Lys Asn Gln L<br>1000 | Lys Leu Ile Ala Asn Ala Phe<br>1005 |
| Asn Lys Ala     | Leu Leu Ser | Ile Gln Asn           | Gly Phe Thr Ala Thr Asn             |
| 1010            |             | 1015                  | 1020                                |
| Ser Ala Leu     | Ala Lys Ile | Gln Ser Val           | Val Asn Ala Asn Ala Gln             |
| 1025            |             | 1030                  | 1035                                |
| Ala Leu Asn     | Ser Leu Leu | Gln Gln Leu           | Phe Asn Lys Phe Gly Ala             |
| 1040            |             | 1045                  | 1050                                |
| Ile Ser Ser     | Ser Leu Gln | Glu Ile Leu           | Ser Arg Leu Asp Asn Leu             |
| 1055            |             | 1060                  | 1065                                |
| Glu Ala Gln     | Val Gln Ile | Asp Arg Leu           | Ile Asn Gly Arg Leu Thr             |
| 1070            |             | 1075                  | 1080                                |
| Ala Leu Asn     | Ala Tyr Val | Ser Gln Gln           | Leu Ser Asp Ile Thr Leu             |
| 1085            |             | 1090                  | 1095                                |
| Ile Lys Ala     | Gly Ala Ser | Arg Ala Ile           | Glu Lys Val Asn Glu Cys             |
| 1100            |             | 1105                  | 1110                                |
| Val Lys Ser     | Gln Ser Pro | Arg Ile Asn           | Phe Cys Gly Asn Gly Asn             |
| 1115            |             | 1120                  | 1125                                |
| His Ile Leu     | Ser Leu Val | Gln Asn Ala           | Pro Tyr Gly Leu Leu Phe             |
| 1130            |             | 1135                  | 1140                                |
| Ile His Phe     | Ser Tyr Lys | Pro Thr Ser           | Phe Lys Thr Val Leu Val             |
| 1145            |             | 1150                  | 1155                                |
| Ser Pro Gly     | Leu Cys Leu | Ser Gly Asp           | Arg Gly Ile Ala Pro Lys             |
| 1160            |             | 1165                  | 1170                                |
| Gln Gly Tyr     | Phe Ile Lys | Gln Asn Asp           | Ser Trp Met Phe Thr Gly             |
| 1175            |             | 1180                  | 1185                                |
| Ser Ser Tyr     | Tyr Tyr Pro | Glu Pro Ile           | Ser Asp Lys Asn Val Val             |
| 1190            |             | 1195                  | 1200                                |
| Phe Met Asn     | Ser Cys Ser | Val Asn Phe           | Thr Lys Ala Pro Phe Ile             |
| 1205            |             | 1210                  | 1215                                |
| Tyr Leu Asn     | Asn Ser Ile | Pro Asn Leu           | Ser Asp Phe Glu Ala Glu             |
| 1220            |             | 1225                  | 1230                                |
| Leu Ser Leu     | Trp Phe Lys | Asn His Thr           | Ser Ile Ala Pro Asn Leu             |
| 1235            |             | 1240                  | 1245                                |
| Thr Phe Asn     | Ser His Ile | Asn Ala Thr           | Phe Leu Asp Leu Tyr Tyr             |
| 1250            |             | 1255                  | 1260                                |
| Glu Met Asn     | Val Ile Gln | Glu Ser Ile           | Lys Ser Leu Asn Ser Ser             |
| 1265            |             | 1270                  | 1275                                |
| Phe Ile Asn     | Leu Lys Glu | Ile Gly Thr           | Tyr Glu Met Tyr Val Lys             |
| 1280            |             | 1285                  | 1290                                |
| Trp Pro Trp     | Tyr Ile Trp | Leu Leu Ile           | Val Ile Leu Phe Ile Ile             |
| 1295            |             | 1300                  | 1305                                |
| Phe Leu Met     | Ile Leu Phe | Phe Ile Cys           | Cys Cys Thr Gly Cys Gly             |
| 1310            |             | 1315                  | 1320                                |
| Ser Ala Cys     | Phe Ser Lys | Cys His Asn           | Cys Cys Asp Glu Tyr Gly             |
| 1325            |             | 1330                  | 1335                                |
| Gly His Asn     | Asp Phe Val | Ile Lys Ala           | Ser His Asp Asp                     |
| 1340            |             | 1345                  | 1350                                |
| <210> SEQ ID    | NO 32       |                       |                                     |

<sup>&</sup>lt;210> SEQ ID NO 32 <211> LENGTH: 526 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence

|            | -continued   |            |            |            |            |            |            |            |            |            |            |            |            |            |            |
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|            | <pre>&lt;220&gt; FEATURE:<br/>&lt;223&gt; OTHER INFORMATION: Synthetic Polypeptide</pre> |            |            |            |            |            |            |            |            |            |            |            |            |            |            |
| < 40       | 0 > SI   | EQUEI      | ICE:       | 32         |            |            |            |            |            |            |            |            |            |            |            |
| Met<br>1   | Phe  | Ile        | Phe        | Leu<br>5   | Leu        | Phe        | Leu        | Thr        | Leu<br>10  | Thr        | Ser        | Gly        | Ser        | Asp<br>15  | Leu        |
| Asp        | Arg  | Ala        | Leu<br>20  | Ser        | Gly        | Ile        | Ala        | Ala<br>25  | Glu        | Gln        | Asp        | Arg        | Asn<br>30  | Thr        | Arg        |
| Glu        | Val  | Phe<br>35  | Ala        | Gln        | Val        | Lys        | Gln<br>40  | Met        | Tyr        | Lys        | Thr        | Pro<br>45  | Thr        | Leu        | Lys        |
| Tyr        | Phe<br>50  | Gly        | Gly        | Phe        | Asn        | Phe<br>55  | Ser        | Gln        | Ile        | Leu        | Pro<br>60  | Asp        | Pro        | Leu        | Lys        |
| Pro<br>65  | Thr  | Lys        | Arg        | Ser        | Phe<br>70  | Ile        | Glu        | Asp        | Leu        | Leu<br>75  | Phe        | Asn        | Lys        | Val        | Thr<br>80  |
| Leu        | Ala  | Asp        | Ala        | Gly<br>85  | Phe        | Met        | Lys        | Gln        | Tyr<br>90  | Gly        | Glu        | CAa        | Leu        | Gly<br>95  | Asp        |
| Ile        | Asn  | Ala        | Arg<br>100 | Asp        | Leu        | Ile        | CÀa        | Ala<br>105 | Gln        | Lys        | Phe        | Asn        | Gly<br>110 | Leu        | Thr        |
| Val        | Leu  | Pro<br>115 | Pro        | Leu        | Leu        | Thr        | Asp<br>120 | Asp        | Met        | Ile        | Ala        | Ala<br>125 | Tyr        | Thr        | Ala        |
| Ala        | Leu<br>130   | Val        | Ser        | Gly        | Thr        | Ala<br>135 | Thr        | Ala        | Gly        | Trp        | Thr<br>140 | Phe        | Gly        | Ala        | Gly        |
| Ala<br>145 | Ala  | Leu        | Gln        | Ile        | Pro<br>150 | Phe        | Ala        | Met        | Gln        | Met<br>155 | Ala        | Tyr        | Arg        | Phe        | Asn<br>160 |
| Gly        | Ile  | Gly        | Val        | Thr<br>165 | Gln        | Asn        | Val        | Leu        | Tyr<br>170 | Glu        | Asn        | Gln        | Lys        | Gln<br>175 | Ile        |
| Ala        | Asn  | Gln        | Phe<br>180 | Asn        | rAa        | Ala        | Ile        | Ser<br>185 | Gln        | Ile        | Gln        | Glu        | Ser<br>190 | Leu        | Thr        |
| Thr        | Thr  | Ser<br>195 | Thr        | Ala        | Leu        | Gly        | Lys<br>200 | Leu        | Gln        | Asp        | Val        | Val<br>205 | Asn        | Gln        | Asn        |
| Ala        | Gln<br>210   | Ala        | Leu        | Asn        | Thr        | Leu<br>215 | Val        | ГÀЗ        | Gln        | Leu        | Ser<br>220 | Ser        | Asn        | Phe        | Gly        |
| Ala<br>225 | Ile  | Ser        | Ser        | Val        | Leu<br>230 | Asn        | Asp        | Ile        | Leu        | Ser<br>235 | Arg        | Leu        | Asp        | Lys        | Val<br>240 |
| Glu        | Ala  | Glu        | Val        | Gln<br>245 | Ile        | Asp        | Arg        | Leu        | Ile<br>250 | Thr        | Gly        | Arg        | Leu        | Gln<br>255 | Ser        |
| Leu        | Gln  |            | Tyr<br>260 |            | Thr        | Gln        |            | Leu<br>265 |            |            | Ala        |            | Glu<br>270 |            | Arg        |
| Ala        | Ser  | Ala<br>275 | Asn        | Leu        | Ala        | Ala        | Thr<br>280 | Lys        | Met        | Ser        | Glu        | Сув<br>285 | Val        | Leu        | Gly        |
| Gln        | Ser<br>290   | Lys        | Arg        | Val        | Asp        | Phe<br>295 | Cys        | Gly        | Lys        | Gly        | Tyr<br>300 | His        | Leu        | Met        | Ser        |
| Phe<br>305 | Pro  | Gln        | Ala        | Ala        | Pro<br>310 | His        | Gly        | Val        | Val        | Phe<br>315 | Leu        | His        | Val        | Thr        | Tyr<br>320 |
| Val        | Pro  | Ser        | Gln        | Glu<br>325 | Arg        | Asn        | Phe        | Thr        | Thr<br>330 | Ala        | Pro        | Ala        | Ile        | Сув<br>335 | His        |
| Glu        | Gly  | Lys        | Ala<br>340 | Tyr        | Phe        | Pro        | Arg        | Glu<br>345 | Gly        | Val        | Phe        | Val        | Phe<br>350 | Asn        | Gly        |
| Thr        | Ser  | Trp<br>355 | Phe        | Ile        | Thr        | Gln        | Arg<br>360 | Asn        | Phe        | Phe        | Ser        | Pro<br>365 | Gln        | Ile        | Ile        |
| Thr        | Thr<br>370   | Asp        | Asn        | Thr        | Phe        | Val<br>375 | Ser        | Gly        | Asn        | Сув        | Asp        | Val        | Val        | Ile        | Gly        |
| Ile<br>385 | Ile  | Asn        | Asn        | Thr        | Val<br>390 | Tyr        | Asp        | Pro        | Leu        | Gln<br>395 | Pro        | Glu        | Leu        | Asp        | Ser<br>400 |
|            |  |            |            |            |            |            |            |            |            |            |            |            |            |            |            |

Phe Lys Glu Glu Leu Asp Lys Tyr Phe Lys Asn His Thr Ser Pro Asp 410 Val Asp Leu Gly Asp Ile Ser Gly Ile Asn Ala Ser Val Val Asn Ile 425 Gln Lys Glu Ile Asp Arg Leu Asn Glu Val Ala Lys Asn Leu Asn Glu Ser Leu Ile Asp Leu Gln Glu Leu Gly Lys Tyr Glu Gln Tyr Ile Lys Trp Pro Trp Tyr Val Trp Leu Gly Phe Ile Ala Gly Leu Ile Ala Ile Val Met Val Thr Ile Leu Leu Cys Cys Met Thr Ser Cys Cys Ser Cys Leu Lys Gly Ala Cys Ser Cys Gly Ser Cys Cys Lys Phe Asp Glu Asp Asp Ser Glu Pro Val Leu Lys Gly Val Lys Leu His Tyr Thr <210> SEO ID NO 33 <211> LENGTH: 588 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223 > OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 33 Met Ile His Ser Val Phe Leu Leu Met Phe Leu Leu Thr Pro Thr Glu 10 Ser Asp Cys Lys Leu Pro Leu Gly Gln Ser Leu Cys Ala Leu Pro Asp 25 Thr Pro Ser Thr Leu Thr Pro Arg Ser Val Arg Ser Val Pro Gly Glu Met Arg Leu Ala Ser Ile Ala Phe Asn His Pro Ile Gln Val Asp Gln Leu Asn Ser Ser Tyr Phe Lys Leu Ser Ile Pro Thr Asn Phe Ser Phe 75 Gly Val Thr Gln Glu Tyr Ile Gln Thr Thr Ile Gln Lys Val Thr Val Asp Cys Lys Gln Tyr Val Cys Asn Gly Phe Gln Lys Cys Glu Gln Leu Leu Arg Glu Tyr Gly Gln Phe Cys Ser Lys Ile Asn Gln Ala Leu His Gly Ala Asn Leu Arg Gln Asp Asp Ser Val Arg Asn Leu Phe Ala Ser Val Lys Ser Ser Gln Ser Ser Pro Ile Ile Pro Gly Phe Gly Gly Asp 155 Phe Asn Leu Thr Leu Leu Glu Pro Val Ser Ile Ser Thr Gly Ser Arg Ser Ala Arg Ser Ala Ile Glu Asp Leu Leu Phe Asp Lys Val Thr Ile 185 Ala Asp Pro Gly Tyr Met Gln Gly Tyr Asp Asp Cys Met Gln Gly 200 Pro Ala Ser Ala Arg Asp Leu Ile Cys Ala Gln Tyr Val Ala Gly Tyr 215 220 Lys Val Leu Pro Pro Leu Met Asp Val Asn Met Glu Ala Ala Tyr Thr 230

Ser Ser Leu Leu Gly Ser Ile Ala Gly Val Gly Trp Thr Ala Gly Leu 250 Ser Ser Phe Ala Ala Ile Pro Phe Ala Gln Ser Ile Phe Tyr Arg Leu 265 Asn Gly Val Gly Ile Thr Gln Gln Val Leu Ser Glu Asn Gln Lys Leu Ile Ala Asn Lys Phe Asn Gln Ala Leu Gly Ala Met Gln Thr Gly Phe Thr Thr Thr Asn Glu Ala Phe Gln Lys Val Gln Asp Ala Val Asn Asn Asn Ala Gln Ala Leu Ser Lys Leu Ala Ser Glu Leu Ser Asn Thr Phe Gly Ala Ile Ser Ala Ser Ile Gly Asp Ile Ile Gln Arg Leu Asp Val Leu Glu Gln Asp Ala Gln Ile Asp Arg Leu Ile Asn Gly Arg Leu Thr 360 Thr Leu Asn Ala Phe Val Ala Gln Gln Leu Val Arg Ser Glu Ser Ala 375 Ala Leu Ser Ala Gln Leu Ala Lys Asp Lys Val Asn Glu Cys Val Lys Ala Gln Ser Lys Arg Ser Gly Phe Cys Gly Gln Gly Thr His Ile Val 410 Ser Phe Val Val Asn Ala Pro Asn Gly Leu Tyr Phe Met His Val Gly 425 Tyr Tyr Pro Ser Asn His Ile Glu Val Val Ser Ala Tyr Gly Leu Cys 440 Asp Ala Ala Asn Pro Thr Asn Cys Ile Ala Pro Val Asn Gly Tyr Phe Ile Lys Thr Asn Asn Thr Arg Ile Val Asp Glu Trp Ser Tyr Thr Gly 470 475 Ser Ser Phe Tyr Ala Pro Glu Pro Ile Thr Ser Leu Asn Thr Lys Tyr Val Ala Pro Gln Val Thr Tyr Gln Asn Ile Ser Thr Asn Leu Pro Pro 505 Pro Leu Leu Gly Asn Ser Thr Gly Ile Asp Phe Gln Asp Glu Leu Asp Glu Phe Phe Lys Asn Val Ser Thr Ser Ile Pro Asn Phe Gly Ser Leu 535 Thr Gln Ile Asn Thr Thr Leu Leu Asp Leu Thr Tyr Glu Met Leu Ser Leu Gln Gln Val Val Lys Ala Leu Asn Glu Ser Tyr Ile Asp Leu Lys Glu Leu Gly Asn Tyr Thr Tyr Tyr Asn Lys Trp Pro 580 <210> SEQ ID NO 34 <211> LENGTH: 526 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 34 Met Phe Ile Phe Leu Leu Phe Leu Thr Leu Thr Ser Gly Ser Asp Leu

5

| Asp        | Arg        | Ala        | Leu<br>20  | Ser        | Gly        | Ile        | Ala        | Ala<br>25  | Glu        | Gln        | Asp        | Arg        | Asn<br>30  | Thr        | Arg        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Glu        | Val        | Phe<br>35  | Ala        | Gln        | Val        | Lys        | Gln<br>40  | Met        | Tyr        | Lys        | Thr        | Pro<br>45  | Thr        | Leu        | Lys        |
| Tyr        | Phe<br>50  | Gly        | Gly        | Phe        | Asn        | Phe<br>55  | Ser        | Gln        | Ile        | Leu        | Pro<br>60  | Asp        | Pro        | Leu        | Lys        |
| Pro<br>65  | Thr        | Lys        | Arg        | Ser        | Phe<br>70  | Ile        | Glu        | Asp        | Leu        | Leu<br>75  | Phe        | Asn        | Lys        | Val        | Thr<br>80  |
| Leu        | Ala        | Asp        | Ala        | Gly<br>85  | Phe        | Met        | Lys        | Gln        | Tyr<br>90  | Gly        | Glu        | Cys        | Leu        | Gly<br>95  | Asp        |
| Ile        | Asn        | Ala        | Arg<br>100 | Asp        | Leu        | Ile        | Cys        | Ala<br>105 | Gln        | Lys        | Phe        | Asn        | Gly<br>110 | Leu        | Thr        |
| Val        | Leu        | Pro<br>115 | Pro        | Leu        | Leu        | Thr        | Asp<br>120 | Asp        | Met        | Ile        | Ala        | Ala<br>125 | Tyr        | Thr        | Ala        |
| Ala        | Leu<br>130 | Val        | Ser        | Gly        | Thr        | Ala<br>135 | Thr        | Ala        | Gly        | Trp        | Thr<br>140 | Phe        | Gly        | Ala        | Gly        |
| Ala<br>145 | Ala        | Leu        | Gln        | Ile        | Pro<br>150 | Phe        | Ala        | Met        | Gln        | Met<br>155 | Ala        | Tyr        | Arg        | Phe        | Asn<br>160 |
| Gly        | Ile        | Gly        | Val        | Thr<br>165 | Gln        | Asn        | Val        | Leu        | Tyr<br>170 | Glu        | Asn        | Gln        | ГЛа        | Gln<br>175 | Ile        |
| Ala        | Asn        | Gln        | Phe<br>180 | Asn        | Lys        | Ala        | Ile        | Ser<br>185 | Gln        | Ile        | Gln        | Glu        | Ser<br>190 | Leu        | Thr        |
| Thr        | Thr        | Ser<br>195 | Thr        | Ala        | Leu        | Gly        | Lys<br>200 | Leu        | Gln        | Asp        | Val        | Val<br>205 | Asn        | Gln        | Asn        |
| Ala        | Gln<br>210 | Ala        | Leu        | Asn        | Thr        | Leu<br>215 | Val        | Lys        | Gln        | Leu        | Ser<br>220 | Ser        | Asn        | Phe        | Gly        |
| Ala<br>225 | Ile        | Ser        | Ser        | Val        | Leu<br>230 | Asn        | Asp        | Ile        | Leu        | Ser<br>235 | Arg        | Leu        | Asp        | Lys        | Val<br>240 |
| Glu        | Ala        | Glu        | Val        | Gln<br>245 | Ile        | Asp        | Arg        | Leu        | Ile<br>250 | Thr        | Gly        | Arg        | Leu        | Gln<br>255 | Ser        |
| Leu        | Gln        | Thr        | Tyr<br>260 | Val        | Thr        | Gln        | Gln        | Leu<br>265 | Ile        | Arg        | Ala        | Ala        | Glu<br>270 | Ile        | Arg        |
| Ala        | Ser        | Ala<br>275 | Asn        | Leu        | Ala        | Ala        | Thr<br>280 | Lys        | Met        | Ser        | Glu        | Сув<br>285 | Val        | Leu        | Gly        |
| Gln        | Ser<br>290 | Lys        | Arg        | Val        | Asp        | Phe<br>295 | Cys        | Gly        | Lys        | Gly        | Tyr<br>300 | His        | Leu        | Met        | Ser        |
| Phe<br>305 | Pro        | Gln        | Ala        | Ala        | Pro<br>310 | His        | Gly        | Val        | Val        | Phe<br>315 | Leu        | His        | Val        | Thr        | Tyr<br>320 |
| Val        | Pro        | Ser        | Gln        | Glu<br>325 | Arg        | Asn        | Phe        | Thr        | Thr<br>330 | Ala        | Pro        | Ala        | Ile        | Сув<br>335 | His        |
| Glu        | Gly        | Lys        | Ala<br>340 | Tyr        | Phe        | Pro        | Arg        | Glu<br>345 | Gly        | Val        | Phe        | Val        | Phe<br>350 | Asn        | Gly        |
| Thr        | Ser        | Trp<br>355 | Phe        | Ile        | Thr        | Gln        | Arg<br>360 | Asn        | Phe        | Phe        | Ser        | Pro<br>365 | Gln        | Ile        | Ile        |
| Thr        | Thr<br>370 | Asp        | Asn        | Thr        | Phe        | Val<br>375 | Ser        | Gly        | Asn        | Cys        | Asp<br>380 | Val        | Val        | Ile        | Gly        |
| Ile<br>385 | Ile        | Asn        | Asn        | Thr        | Val<br>390 | Tyr        | Asp        | Pro        | Leu        | Gln<br>395 | Pro        | Glu        | Leu        | Asp        | Ser<br>400 |
| Phe        | Lys        | Glu        | Glu        | Leu<br>405 | Asp        | Lys        | Tyr        | Phe        | Lys<br>410 | Asn        | His        | Thr        | Ser        | Pro<br>415 | Asp        |
| Val        | Asp        | Leu        | Gly<br>420 | Asp        | Ile        | Ser        | Gly        | Ile<br>425 | Asn        | Ala        | Ser        | Val        | Val<br>430 | Asn        | Ile        |
|            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |

| Gln Lys Gl<br>43  |   | rg Leu Asn<br>440 | Glu Val Ala          | Lys Asn Let<br>445 | ı Asn Glu        |      |
|-------------------|---|-------------------|----------------------|--------------------|------------------|------|
| Ser Leu Il<br>450 | e Asp Leu G   | ln Glu Leu<br>455 | Gly Lys Tyr          | Glu Gln Tyr<br>460 | r Ile Lys        |      |
| Trp Pro Tr        |   | rp Leu Gly<br>70  | Phe Ile Ala          | _                  | e Ala Ile<br>480 |      |
| Val Met Va        | l Thr Ile L<br>485                                      | eu Leu Cys        | Cys Met Thr          | Ser Cys Cys        | s Ser Cys<br>495 |      |
| Leu Lys Gl        | y Ala Cys S<br>500                                      | er Cys Gly        | Ser Cys Cys          | Lys Phe Asp        |                  |      |
| Asp Ser Gl        |   | eu Lys Gly<br>520 | Val Lys Leu          | His Tyr Thi        | r                |      |
| <220> FEAT        | TH: 1864<br>: DNA<br>NISM: Artif<br>URE:<br>R INFORMATI | -                 | ence<br>tic Polynucl | eotide             |                  |      |
|                   |   | acagaagct         | a atacgactca         | ctatagggaa         | ataagagaga       | 60   |
|                   |   |                   | a ccatgggtct         |                    |                  | 120  |
|                   |   |                   | a cacccgccgg         |                    |                  | 180  |
| tctctaagat        | aggggtagta  | ggaatagga         | a gtgcaagcta         | caaagttatg         | actcgttcca       | 240  |
| gccatcaatc        | attagtcata  | aaattaatg         | c ccaatataac         | tctcctcaat         | aactgcacga       | 300  |
| gggtagagat        | tgcagaatac  | aggagacta         | c taagaacagt         | tttggaacca         | attagggatg       | 360  |
| cacttaatgc        | aatgacccag  | aacataagg         | c cggttcagag         | cgtagcttca         | agtaggagac       | 420  |
| acaagagatt        | tgcgggagta  | gteetggea         | g gtgcggccct         | aggtgttgcc         | acagctgctc       | 480  |
| agataacagc        | cggcattgca  | cttcaccgg         | t ccatgctgaa         | ctctcaggcc         | atcgacaatc       | 540  |
| tgagagcgag        | cctggaaact  | actaatcag         | g caattgaggo         | aatcagacaa         | gcagggcagg       | 600  |
| agatgatatt        | ggctgttcag  | ggtgtccaa         | g actacatcaa         | taatgagctg         | ataccgtcta       | 660  |
| tgaaccagct        | atcttgtgat  | ctaatcggt         | c agaagctcgg         | gctcaaattg         | cttagatact       | 720  |
| atacagaaat        | cctgtcatta  | tttggcccc         | a geetaeggga         | ccccatatct         | gcggagatat       | 780  |
| ctatccaggc        | tttgagttat  | gcacttgga         | g gagatatcaa         | taaggtgtta         | gaaaagctcg       | 840  |
| gatacagtgg        | aggcgattta  | ctaggcatc         | t tagagagcag         | aggaataaag         | gctcggataa       | 900  |
| ctcacgtcga        | cacagagtco  | tacttcata         | g teeteagtat         | agcctatccg         | acgctgtccg       | 960  |
| agattaaggg        | ggtgattgto  | caccggcta         | g agggggtctc         | gtacaacata         | ggctctcaag       | 1020 |
| agtggtatac        | cactgtgccc  | aagtatgtt         | g caacccaagg         | gtaccttatc         | tcgaattttg       | 1080 |
| atgagtcatc        | atgtactttc  | atgccagag         | g ggactgtgtg         | cagccaaaat         | gccttgtacc       | 1140 |
| cgatgagtcc        | tctgctccaa  | gaatgeete         | c gggggtccac         | caagtcctgt         | gctcgtacac       | 1200 |
| tcgtatccgg        | gtcttttggg  | aaccggttc         | a ttttatcaca         | agggaaccta         | atagccaatt       | 1260 |
| gtgcatcaat        | tetttgtaag  | tgttacaca         | a caggtacgat         | tattaatcaa         | gaccctgaca       | 1320 |
| agatcctaac        | atacattgct  | geegatege         | t gcccggtagt         | cgaggtgaac         | ggcgtgacca       | 1380 |
| tccaagtcgg        | gagcaggagg  | tatccagac         | g ctgtgtactt         | gcacagaatt         | gacctcggtc       | 1440 |
| ctcccatatc        | attggagagg  | ttggacgta         | g ggacaaatct         | ggggaatgca         | attgccaaat       | 1500 |
| tggaggatgc        | caaggaattg  | ttggaatca         | t cggaccagat         | attgagaagt         | atgaaaggtt       | 1560 |

| tatcgagcac t   | agcatagtc                            | tacatcctga | ttgcagtgtg | tcttggaggg | ttgataggga | 1620 |
|--|--------------------------------------|------------|------------|------------|------------|------|
| tccccacttt a   | aatatgttgc                           | tgcagggggc | gttgtaacaa | aaagggagaa | caagttggta | 1680 |
| tgtcaagacc a   | aggcctaaag                           | cctgacctta | caggaacatc | aaaatcctat | gtaagatcgc | 1740 |
| tttgatgata a   | ataggctgga                           | gcctcggtgg | ccaagcttct | tgccccttgg | gcctccccc  | 1800 |
| ageceeteet e   | cccttcctg                            | cacccgtacc | cccgtggtct | ttgaataaag | tctgagtggg | 1860 |
| cggc   |                                      |            |            |            |            | 1864 |
| <210> SEQ II<br><211> LENGTH<br><212> TYPE:<br><213> ORGANI<br><220> FEATUR<br><223> OTHER | H: 1653<br>DNA<br>ISM: Artifi<br>RE: |            |            | eotide     |            |      |
| <400> SEQUEN   | NCE: 36                              |            |            |            |            |      |
| atgggtctca a   | aggtgaacgt                           | ctctgccgta | ttcatggcag | tactgttaac | tctccaaaca | 60   |
| cccgccggtc a   | aaattcattg                           | gggcaatctc | tctaagatag | gggtagtagg | aataggaagt | 120  |
| gcaagctaca a   | aagttatgac                           | tcgttccagc | catcaatcat | tagtcataaa | attaatgccc | 180  |
| aatataactc t   | cctcaataa                            | ctgcacgagg | gtagagattg | cagaatacag | gagactacta | 240  |
| agaacagttt t   | ggaaccaat                            | tagggatgca | cttaatgcaa | tgacccagaa | cataaggccg | 300  |
| gttcagagcg t   | agcttcaag                            | taggagacac | aagagatttg | cgggagtagt | cctggcaggt | 360  |
| geggeeetag g   | gtgttgccac                           | agctgctcag | ataacagccg | gcattgcact | tcaccggtcc | 420  |
| atgctgaact c   | ctcaggccat                           | cgacaatctg | agagcgagcc | tggaaactac | taatcaggca | 480  |
| attgaggcaa t   | cagacaagc                            | agggcaggag | atgatattgg | ctgttcaggg | tgtccaagac | 540  |
| tacatcaata a   | atgagctgat                           | accgtctatg | aaccagctat | cttgtgatct | aatcggtcag | 600  |
| aagctcgggc t   | caaattgct                            | tagatactat | acagaaatcc | tgtcattatt | tggccccagc | 660  |
| ctacgggacc c   | ccatatctgc                           | ggagatatct | atccaggctt | tgagttatgc | acttggagga | 720  |
| gatatcaata a   | aggtgttaga                           | aaagctcgga | tacagtggag | gcgatttact | aggcatctta | 780  |
| gagagcagag g   | gaataaaggc                           | tcggataact | cacgtcgaca | cagagtccta | cttcatagtc | 840  |
| ctcagtatag c   | ectatecgae                           | gctgtccgag | attaaggggg | tgattgtcca | ccggctagag | 900  |
| ggggtetegt a   | acaacatagg                           | ctctcaagag | tggtatacca | ctgtgcccaa | gtatgttgca | 960  |
| acccaagggt a   | accttatctc                           | gaattttgat | gagtcatcat | gtactttcat | gccagagggg | 1020 |
| actgtgtgca g   | gccaaaatgc                           | cttgtacccg | atgagtcctc | tgctccaaga | atgcctccgg | 1080 |
| gggtccacca a   | agtcctgtgc                           | tcgtacactc | gtatccgggt | cttttgggaa | ccggttcatt | 1140 |
| ttatcacaag g   | ggaacctaat                           | agccaattgt | gcatcaattc | tttgtaagtg | ttacacaaca | 1200 |
| ggtacgatta t   | taatcaaga                            | ccctgacaag | atcctaacat | acattgctgc | cgatcgctgc | 1260 |
| ccggtagtcg a   | aggtgaacgg                           | cgtgaccatc | caagteggga | gcaggaggta | tccagacgct | 1320 |
| gtgtacttgc a   | acagaattga                           | cctcggtcct | cccatatcat | tggagaggtt | ggacgtaggg | 1380 |
| acaaatctgg g   | ggaatgcaat                           | tgccaaattg | gaggatgcca | aggaattgtt | ggaatcatcg | 1440 |
| gaccagatat t   | gagaagtat                            | gaaaggttta | tcgagcacta | gcatagtcta | catcctgatt | 1500 |
| gcagtgtgtc t   | tggagggtt                            | gatagggatc | cccactttaa | tatgttgctg | cagggggcgt | 1560 |
| tgtaacaaaa a   |                                      |            |            |            |            | 1620 |
| ggaacatcaa a   |                                      |            |            | -          | -          | 1653 |
| Januara Coud C   |                                      |            | - 5~       |            |            | 1000 |

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<210> SEQ ID NO 37 <211> LENGTH: 1925 <212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEQUENCE: 37

ggggaaataa gagagaaaag aagagtaaga agaaatataa gagccaccat gggtctcaag 60 gtgaacgtct ctgccgtatt catggcagta ctgttaactc tccaaacacc cgccggtcaa 120 attcattggg gcaatctctc taagataggg gtagtaggaa taggaagtgc aagctacaaa 180 gttatgactc gttccagcca tcaatcatta gtcataaaat taatgcccaa tataactctc ctcaataact gcacgagggt agagattgca gaatacagga gactactaag aacagttttg gaaccaatta gggatgcact taatgcaatg acccagaaca taaggccggt tcagagcgta 360 gcttcaagta ggagacacaa gagatttgcg ggagtagtcc tggcaggtgc ggccctaggt 420 qttqccacaq ctqctcaqat aacaqccqqc attqcacttc accqqtccat qctqaactct 480 caggicateg acaatetgag agegageetg gaaactaeta ateaggeaat tgaggeaate 540 agacaagcag ggcaggagat gatattggct gttcagggtg tccaagacta catcaataat 600 gagetgatae egtetatgaa eeagetatet tgtgatetaa teggteagaa getegggete 660 aaattqctta qatactatac aqaaatcctq tcattatttq qccccaqcct acqqqacccc 720 atatctgcgg agatatctat ccaggctttg agttatgcac ttggaggaga tatcaataag 780 840 gtgttagaaa agctcggata cagtggaggc gatttactag gcatcttaga gagcagagga ataaaggete ggataaetea egtegaeaca gagteetaet teatagteet eagtatagee 900 tatccgacgc tgtccgagat taagggggtg attgtccacc ggctagaggg ggtctcgtac 960 aacatagget eteaagagtg gtataceaet gtgeecaagt atgttgeaae eeaagggtae 1020 cttatctcga attttgatga gtcatcatgt actttcatgc cagaggggac tgtgtgcagc 1080 caaaatgcct tgtacccgat gagtcctctg ctccaagaat gcctccgggg gtccaccaag 1140 teetgtgete gtacaetegt atcegggtet tttgggaace ggtteatttt atcacaaggg 1200 aacctaatag ccaattgtgc atcaattctt tgtaagtgtt acacaacagg tacgattatt 1260 aatcaagacc ctgacaagat cctaacatac attgctgccg atcgctgccc ggtagtcgag 1320 1380 gtgaacggcg tgaccatcca agtcgggagc aggaggtatc cagacgctgt gtacttgcac agaattgacc tcggtcctcc catatcattg gagaggttgg acgtagggac aaatctgggg 1440 aatgcaattg ccaaattgga ggatgccaag gaattgttgg aatcatcgga ccagatattg 1500 agaagtatga aaggtttatc gagcactagc atagtctaca tcctgattgc agtgtgtctt ggagggttga tagggatccc cactttaata tgttgctgca gggggcgttg taacaaaaag 1620 qqaqaacaaq ttqqtatqtc aaqaccaqqc ctaaaqcctq accttacaqq aacatcaaaa 1680 tectatgtaa gategetttg atgataatag getggageet eggtggeeaa gettettgee 1740 cettgggeet eccecagee ectecteece tteetgeace egtaceeceg tggtetttga 1860 1920 1925 tctaq

<212> TYPE: DNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEOUENCE: 38

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<sup>&</sup>lt;210> SEQ ID NO 39

<sup>&</sup>lt;211> LENGTH: 1653

<sup>&</sup>lt;212> TYPE: DNA

<sup>&</sup>lt;213> ORGANISM: Artificial Sequence

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Synthetic Polynucleotide

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| <400> SEQUENCE: 39   |            |            |                    |            |      |  |
|--|------------|------------|--------------------|------------|------|--|
| atgggtctca aggtgaacgt  | ctctgtcata | ttcatggcag | tactgttaac         | tcttcaaaca | 60   |  |
| cccaccggtc aaatccattg  | gggcaatctc | tctaagatag | gggtggtagg         | ggtaggaagt | 120  |  |
| gcaagctaca aagttatgac  | tegttecage | catcaatcat | tagtcataaa         | gttaatgccc | 180  |  |
| aatataactc tcctcaacaa  | ttgcacgagg | gtagggattg | cagaatacag         | gagactactg | 240  |  |
| agaacagttc tggaaccaat  | tagagatgca | cttaatgcaa | tgacccagaa         | tataagaccg | 300  |  |
| gttcagagtg tagcttcaag  | taggagacac | aagagatttg | cgggagttgt         | cctggcaggt | 360  |  |
| geggeeetag gegttgeeae  | agctgctcaa | ataacagccg | gtattgcact         | tcaccagtcc | 420  |  |
| atgctgaact ctcaagccat  | cgacaatctg | agagcgagcc | tagaaactac         | taatcaggca | 480  |  |
| attgaggcaa tcagacaagc  | agggcaggag | atgatattgg | ctgttcaggg         | tgtccaagac | 540  |  |
| tacatcaata atgagetgat  | accgtctatg | aatcaactat | cttgtgattt         | aatcggccag | 600  |  |
| aagctagggc tcaaattgct  | cagatactat | acagaaatcc | tgtcattatt         | tggccccagc | 660  |  |
| ttacgggacc ccatatctgc  | ggagatatct | atccaggctt | tgagctatgc         | gcttggagga | 720  |  |
| gatatcaata aggtgttgga  | aaagctcgga | tacagtggag | gtgatctact         | gggcatctta | 780  |  |
| gagagcagag gaataaaggc  | ccggataact | cacgtcgaca | cagagtccta         | cttcattgta | 840  |  |
| ctcagtatag cctatccgac  | gctatccgag | attaaggggg | tgattgtcca         | ccggctagag | 900  |  |
| ggggtctcgt acaacatagg  | ctctcaagag | tggtatacca | ctgtgcccaa         | gtatgttgca | 960  |  |
| acccaagggt accttatctc  | gaattttgat | gagtcatcat | gcactttcat         | gccagagggg | 1020 |  |
| actgtgtgca gccagaatgc  | cttgtacccg | atgagtcctc | tgctccaaga         | atgcctccgg | 1080 |  |
| gggtccacta agtcctgtgc  | tcgtacactc | gtatccgggt | ctttcgggaa         | ccggttcatt | 1140 |  |
| ttatcacagg ggaacctaat  | agccaattgt | gcatcaatcc | tttgcaagtg         | ttacacaaca | 1200 |  |
| ggaacaatca ttaatcaaga  | ccctgacaag | atcctaacat | acattgctgc         | cgatcactgc | 1260 |  |
| ccggtggtcg aggtgaatgg  | cgtgaccatc | caagtcggga | gcaggaggta         | tccggacgct | 1320 |  |
| gtgtacttgc acaggattga  | cctcggtcct | cccatatctt | tggagaggtt         | ggacgtaggg | 1380 |  |
| acaaatctgg ggaatgcaat  | tgctaagttg | gaggatgcca | aggaattgtt         | ggagtcatcg | 1440 |  |
| gaccagatat tgaggagtat  | gaaaggttta | tcgagcacta | gtatagttta         | catcctgatt | 1500 |  |
| gcagtgtgtc ttggaggatt  | gatagggatc | cccgctttaa | tatgttgctg         | cagggggcgt | 1560 |  |
| tgtaacaaga agggagaaca  | agttggtatg | tcaagaccag | gcctaaagcc         | tgatcttaca | 1620 |  |
| ggaacatcaa aatcctatgt  | aaggtcactc | tga        |                    |            | 1653 |  |
| <pre>&lt;210&gt; SEQ ID NO 40 &lt;211&gt; LENGTH: 1925 &lt;212&gt; TYPE: DNA &lt;213&gt; ORGANISM: Artifi &lt;220&gt; FEATURE: &lt;223&gt; OTHER INFORMATIO</pre> <400> SEQUENCE: 40   | _          |            | <del>c</del> otide |            |      |  |
| -  | 2202012200 | 20222+0+0  | anaganant.         | agatatass= | 60   |  |
| ggggaaataa gagagaaaag  |            |            |                    |            |      |  |
| gtgaacgtct ctgtcatatt  |            |            |                    |            | 120  |  |
| atccattggg gcaatctctc  |            |            |                    |            | 180  |  |
| gttatgactc gttccagcca  | tcaatcatta | gtcataaagt | taatgcccaa         | tataactctc | 240  |  |
| ctcaacaatt gcacgagggt  | agggattgca | gaatacagga | gactactgag         | aacagttctg | 300  |  |
| and the second s |            |            |                    |            | 260  |  |

gaaccaatta gagatgcact taatgcaatg acccagaata taagaccggt tcagagtgta

| getteaagta ggagacacaa gagatttgeg   | ggagttgtcc | tggcaggtgc | ggccctaggc | 420  |  |  |  |  |  |
|--|------------|------------|------------|------|--|--|--|--|--|
| gttgccacag ctgctcaaat aacagccggt   | attgcacttc | accagtccat | gctgaactct | 480  |  |  |  |  |  |
| caagccatcg acaatctgag agcgagccta   | gaaactacta | atcaggcaat | tgaggcaatc | 540  |  |  |  |  |  |
| agacaagcag ggcaggagat gatattggct   | gttcagggtg | tccaagacta | catcaataat | 600  |  |  |  |  |  |
| gagetgatae egtetatgaa teaactatet   | tgtgatttaa | teggeeagaa | gctagggctc | 660  |  |  |  |  |  |
| aaattgctca gatactatac agaaatcctg   | tcattatttg | gccccagctt | acgggacccc | 720  |  |  |  |  |  |
| atatetgegg agatatetat ceaggetttg   | agctatgcgc | ttggaggaga | tatcaataag | 780  |  |  |  |  |  |
| gtgttggaaa agctcggata cagtggaggt   | gatctactgg | gcatcttaga | gagcagagga | 840  |  |  |  |  |  |
| ataaaggccc ggataactca cgtcgacaca   | gagtcctact | tcattgtact | cagtatagcc | 900  |  |  |  |  |  |
| tatccgacgc tatccgagat taagggggtg   | attgtccacc | ggctagaggg | ggtctcgtac | 960  |  |  |  |  |  |
| aacataggct ctcaagagtg gtataccact   | gtgcccaagt | atgttgcaac | ccaagggtac | 1020 |  |  |  |  |  |
| cttatctcga attttgatga gtcatcatgc   | actttcatgc | cagaggggac | tgtgtgcagc | 1080 |  |  |  |  |  |
| cagaatgeet tgtaccegat gagteetetg   | ctccaagaat | gcctccgggg | gtccactaag | 1140 |  |  |  |  |  |
| teetgtgete gtacaetegt ateegggtet   | ttcgggaacc | ggttcatttt | atcacagggg | 1200 |  |  |  |  |  |
| aacctaatag ccaattgtgc atcaatcctt   | tgcaagtgtt | acacaacagg | aacaatcatt | 1260 |  |  |  |  |  |
| aatcaagacc ctgacaagat cctaacatac   | attgctgccg | atcactgccc | ggtggtcgag | 1320 |  |  |  |  |  |
| gtgaatggcg tgaccatcca agtcgggagc   | aggaggtatc | cggacgctgt | gtacttgcac | 1380 |  |  |  |  |  |
| aggattgacc tcggtcctcc catatctttg   | gagaggttgg | acgtagggac | aaatctgggg | 1440 |  |  |  |  |  |
| aatgcaattg ctaagttgga ggatgccaag   | gaattgttgg | agtcatcgga | ccagatattg | 1500 |  |  |  |  |  |
| aggagtatga aaggtttatc gagcactagt   | atagtttaca | tcctgattgc | agtgtgtctt | 1560 |  |  |  |  |  |
| ggaggattga tagggatccc cgctttaata   | tgttgctgca | gggggcgttg | taacaagaag | 1620 |  |  |  |  |  |
| ggagaacaag ttggtatgtc aagaccaggc   | ctaaagcctg | atcttacagg | aacatcaaaa | 1680 |  |  |  |  |  |
| teetatgtaa ggteaetetg atgataatag   | gctggagcct | cggtggccaa | gcttcttgcc | 1740 |  |  |  |  |  |
| cettgggeet ecceecagee cetecteece   | ttcctgcacc | cgtacccccg | tggtctttga | 1800 |  |  |  |  |  |
| ataaagtctg agtgggcggc aaaaaaaaaa   | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | 1860 |  |  |  |  |  |
| aaaaaaaaaa aaaaaaaaaaaaaaaaaaaaaaaaaaa   | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | 1920 |  |  |  |  |  |
| tctag  |            |            |            | 1925 |  |  |  |  |  |
| <210> SEQ ID NO 41 <211> LENGTH: 2065 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |            |            |            |      |  |  |  |  |  |
| <400> SEQUENCE: 41   |            |            |            | 6.0  |  |  |  |  |  |
| tcaagctttt ggaccctcgt acagaagcta   | _          |            |            | 60   |  |  |  |  |  |
| aaagaagagt aagaagaaat ataagagcca   |            |            |            | 120  |  |  |  |  |  |
| ccttctacaa agataaccct tatcccaagg   | gaagtaggat | agttattaac | agagaacatc | 180  |  |  |  |  |  |
| ttatgattga cagaccctat gttctgctgg   | ctgttctgtt | cgtcatgttt | ctgagcttga | 240  |  |  |  |  |  |
| teggattget ggeaattgea ggeattagae   | ttcatcgggc | agccatctac | accgcggaga | 300  |  |  |  |  |  |
| tccataaaag cctcagtacc aatctggatg   | tgactaactc | catcgagcat | caggtcaagg | 360  |  |  |  |  |  |

acgtgctgac accactcttt aaaatcatcg gggatgaagt gggcctgaga acacctcaga

-continued

| -concinued   |      |  |  |  |  |  |  |  |
|--|------|--|--|--|--|--|--|--|
| gattcactga cctagtgaaa ttcatctcgg acaagattaa attccttaat ccggataggg  | 480  |  |  |  |  |  |  |  |
| agtacgactt cagagatete aettggtgea teaaceegee agagaggate aaactagatt  | 540  |  |  |  |  |  |  |  |
| atgatcaata ctgtgcagat gtggctgctg aagagctcat gaatgcattg gtgaactcaa  | 600  |  |  |  |  |  |  |  |
| ctctactgga gaccagaaca accactcagt tcctagctgt ctcaaaggga aactgctcag  | 660  |  |  |  |  |  |  |  |
| ggcccactac aatcagaggt caattetcaa acatgteget gteettgttg gaettgtact  | 720  |  |  |  |  |  |  |  |
| taggtcgagg ttacaatgtg tcatctatag tcactatgac atcccaggga atgtatgggg  | 780  |  |  |  |  |  |  |  |
| gaacctacct agttgaaaag cctaatctga acagcaaagg gtcagagttg tcacaactga  | 840  |  |  |  |  |  |  |  |
| gcatgtaccg agtgtttgaa gtaggtgtga tcagaaaccc gggtttgggg gctccggtgt  | 900  |  |  |  |  |  |  |  |
| tccatatgac aaactatttt gagcaaccag tcagtaatgg tctcggcaac tgtatggtgg  | 960  |  |  |  |  |  |  |  |
| ctttggggga geteaaacte geageeettt gteaegggga egattetate ataatteeet  | 1020 |  |  |  |  |  |  |  |
| atcagggatc agggaaaggt gtcagcttcc agctcgtcaa gctgggtgtc tggaaatccc  | 1080 |  |  |  |  |  |  |  |
| caaccgacat gcaatcctgg gtccccttat caacggatga tccagtggta gacaggcttt  | 1140 |  |  |  |  |  |  |  |
| accteteate teacagaggt gteategetg acaateaage aaaatggget gteeegacaa  | 1200 |  |  |  |  |  |  |  |
| cacgaacaga tgacaagttg cgaatggaga catgcttcca gcaggcgtgt aaaggtaaaa  | 1260 |  |  |  |  |  |  |  |
| tccaagcact ctgcgagaat cccgagtggg taccattgaa ggataacagg attccttcat  | 1320 |  |  |  |  |  |  |  |
| acggggtcct gtctgttgat ctgagtctga cggttgagct taaaatcaaa attgcttcgg  | 1380 |  |  |  |  |  |  |  |
| gattcgggcc attgatcaca cacggctcag ggatggacct atacaaatcc aactgcaaca  | 1440 |  |  |  |  |  |  |  |
| atgtgtattg gctgactatt ccgccaatga gaaatctagc cttaggcgta atcaacacat  | 1500 |  |  |  |  |  |  |  |
| tggagtggat accgagattc aaggttagtc ccaacctctt cactgtccca attaaggaag  | 1560 |  |  |  |  |  |  |  |
| caggcgaaga ctgccatgcc ccaacatacc tacctgcgga ggtggacggt gatgtcaaac  | 1620 |  |  |  |  |  |  |  |
| tcagttccaa cctggtgatt ctacctggtc aagatctcca atatgttttg gcaacctacg  | 1680 |  |  |  |  |  |  |  |
| atacctccag ggttgagcat gctgtggttt attacgttta cagcccaagc cgctcatttt  | 1740 |  |  |  |  |  |  |  |
| cttactttta tccttttagg ttgcctataa agggggtccc aatcgaacta caagtggaat  | 1800 |  |  |  |  |  |  |  |
| getteacatg ggateaaaaa etetggtgee gteacttetg tgtgettgeg gaeteagaat  | 1860 |  |  |  |  |  |  |  |
| ccggtggact tatcactcac tctgggatgg tgggcatggg agtcagctgc acagctaccc  | 1920 |  |  |  |  |  |  |  |
| gggaagatgg aaccaatcgc agataatgat aataggctgg agcctcggtg gccaagcttc  | 1980 |  |  |  |  |  |  |  |
| ttgccccttg ggcctccccc cagcccctcc tccccttcct gcacccgtac ccccgtggtc  | 2040 |  |  |  |  |  |  |  |
| tttgaataaa gtctgagtgg gcggc  | 2065 |  |  |  |  |  |  |  |
| <210> SEQ ID NO 42 <211> LENGTH: 1854 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |      |  |  |  |  |  |  |  |
| <400> SEQUENCE: 42   |      |  |  |  |  |  |  |  |
| atgtcaccgc aacgagaccg gataaatgcc ttctacaaag ataaccctta tcccaaggga  | 60   |  |  |  |  |  |  |  |
| agtaggatag ttattaacag agaacatett atgattgaca gaccetatgt tetgetgget  | 120  |  |  |  |  |  |  |  |
| gttetgtteg teatgtttet gagettgate ggattgetgg caattgeagg cattagaett  | 180  |  |  |  |  |  |  |  |
| catcgggcag ccatctacac cgcggagatc cataaaagcc tcagtaccaa tctggatgtg  | 240  |  |  |  |  |  |  |  |
| actaactcca tcgagcatca ggtcaaggac gtgctgacac cactctttaa aatcatcggg  | 300  |  |  |  |  |  |  |  |
| gatgaagtgg geetgagaae aeeteagaga tteaetgaee tagtgaaatt eateteggae  | 360  |  |  |  |  |  |  |  |
|  |      |  |  |  |  |  |  |  |

aagattaaat teettaatee ggatagggag taegaettea gagateteae ttggtgeate

| aacccgccag   | agaggatcaa | actagattat | gatcaatact | gtgcagatgt | ggctgctgaa | 480  |  |  |
|--|------------|------------|------------|------------|------------|------|--|--|
| gagctcatga   | atgcattggt | gaactcaact | ctactggaga | ccagaacaac | cactcagttc | 540  |  |  |
| ctagctgtct   | caaagggaaa | ctgctcaggg | cccactacaa | tcagaggtca | attctcaaac | 600  |  |  |
| atgtcgctgt   | ccttgttgga | cttgtactta | ggtcgaggtt | acaatgtgtc | atctatagtc | 660  |  |  |
| actatgacat   | cccagggaat | gtatggggga | acctacctag | ttgaaaagcc | taatctgaac | 720  |  |  |
| agcaaagggt   | cagagttgtc | acaactgagc | atgtaccgag | tgtttgaagt | aggtgtgatc | 780  |  |  |
| agaaacccgg   | gtttgggggc | teeggtgtte | catatgacaa | actattttga | gcaaccagtc | 840  |  |  |
| agtaatggtc   | tcggcaactg | tatggtggct | ttgggggagc | tcaaactcgc | agccctttgt | 900  |  |  |
| cacggggacg   | attctatcat | aattccctat | cagggatcag | ggaaaggtgt | cagcttccag | 960  |  |  |
| ctcgtcaagc   | tgggtgtctg | gaaatcccca | accgacatgc | aatcctgggt | ccccttatca | 1020 |  |  |
| acggatgatc   | cagtggtaga | caggctttac | ctctcatctc | acagaggtgt | catcgctgac | 1080 |  |  |
| aatcaagcaa   | aatgggctgt | cccgacaaca | cgaacagatg | acaagttgcg | aatggagaca | 1140 |  |  |
| tgcttccagc   | aggcgtgtaa | aggtaaaatc | caagcactct | gcgagaatcc | cgagtgggta | 1200 |  |  |
| ccattgaagg   | ataacaggat | tccttcatac | ggggtcctgt | ctgttgatct | gagtctgacg | 1260 |  |  |
| gttgagctta   | aaatcaaaat | tgcttcggga | ttcgggccat | tgatcacaca | cggctcaggg | 1320 |  |  |
| atggacctat   | acaaatccaa | ctgcaacaat | gtgtattggc | tgactattcc | gccaatgaga | 1380 |  |  |
| aatctagcct   | taggcgtaat | caacacattg | gagtggatac | cgagattcaa | ggttagtccc | 1440 |  |  |
| aacctcttca   | ctgtcccaat | taaggaagca | ggcgaagact | gccatgcccc | aacataccta | 1500 |  |  |
| cctgcggagg   | tggacggtga | tgtcaaactc | agttccaacc | tggtgattct | acctggtcaa | 1560 |  |  |
| gatctccaat   | atgttttggc | aacctacgat | acctccaggg | ttgagcatgc | tgtggtttat | 1620 |  |  |
| tacgtttaca   | gcccaagccg | ctcattttct | tacttttatc | cttttaggtt | gcctataaag | 1680 |  |  |
| ggggtcccaa   | tcgaactaca | agtggaatgc | ttcacatggg | atcaaaaact | ctggtgccgt | 1740 |  |  |
| cacttctgtg   | tgcttgcgga | ctcagaatcc | ggtggactta | tcactcactc | tgggatggtg | 1800 |  |  |
| ggcatgggag   | tcagctgcac | agctacccgg | gaagatggaa | ccaatcgcag | ataa       | 1854 |  |  |
| <210> SEQ ID NO 43 <211> LENGTH: 2126 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |            |            |            |            |            |      |  |  |
| <400> SEQUE  |            | aagagtaaga | agaaatataa | gagccaccat | gtcaccgcaa | 60   |  |  |
|  |            |            |            | ccaagggaag |            | 120  |  |  |
|  |            |            |            | tgctggctgt |            | 180  |  |  |
|  |            |            |            |            |            | 240  |  |  |
|  |            |            |            | tagactica  |            | 300  |  |  |
|  |            |            |            | tggatgtgac |            |      |  |  |
|  |            |            |            | tcatcgggga |            | 360  |  |  |
|  |            |            |            | tctcggacaa |            | 420  |  |  |
| cttaatccgg   | atagggagta | cgacttcaga | gatctcactt | ggtgcatcaa | cccgccagag | 480  |  |  |
| aggatcaaac   | tagattatga | tcaatactgt | gcagatgtgg | ctgctgaaga | gctcatgaat | 540  |  |  |

gcattggtga actcaactct actggagacc agaacaacca ctcagttcct agctgtctca

| -continued   |      |  |  |  |  |  |  |  |
|--|------|--|--|--|--|--|--|--|
| aagggaaact gctcagggcc cactacaatc agaggtcaat tctcaaacat gtcgctgtcc  | 660  |  |  |  |  |  |  |  |
| ttgttggact tgtacttagg tcgaggttac aatgtgtcat ctatagtcac tatgacatcc  | 720  |  |  |  |  |  |  |  |
| cagggaatgt atgggggaac ctacctagtt gaaaagccta atctgaacag caaagggtca  | 780  |  |  |  |  |  |  |  |
| gagttgtcac aactgagcat gtaccgagtg tttgaagtag gtgtgatcag aaacccgggt  | 840  |  |  |  |  |  |  |  |
| ttgggggctc cggtgttcca tatgacaaac tattttgagc aaccagtcag taatggtctc  | 900  |  |  |  |  |  |  |  |
| ggcaactgta tggtggcttt gggggagctc aaactcgcag ccctttgtca cggggacgat  | 960  |  |  |  |  |  |  |  |
| tctatcataa ttccctatca gggatcaggg aaaggtgtca gcttccagct cgtcaagctg  | 1020 |  |  |  |  |  |  |  |
| ggtgtctgga aatccccaac cgacatgcaa tcctgggtcc ccttatcaac ggatgatcca  | 1080 |  |  |  |  |  |  |  |
| gtggtagaca ggctttacct ctcatctcac agaggtgtca tcgctgacaa tcaagcaaaa  | 1140 |  |  |  |  |  |  |  |
| tgggctgtcc cgacaacacg aacagatgac aagttgcgaa tggagacatg cttccagcag  | 1200 |  |  |  |  |  |  |  |
| gcgtgtaaag gtaaaatcca agcactctgc gagaatcccg agtgggtacc attgaaggat  | 1260 |  |  |  |  |  |  |  |
| aacaggattc cttcatacgg ggtcctgtct gttgatctga gtctgacggt tgagcttaaa  | 1320 |  |  |  |  |  |  |  |
| atcaaaattg cttcgggatt cgggccattg atcacacacg gctcagggat ggacctatac  | 1380 |  |  |  |  |  |  |  |
| aaatccaact gcaacaatgt gtattggctg actattccgc caatgagaaa tctagcctta  | 1440 |  |  |  |  |  |  |  |
| ggcgtaatca acacattgga gtggataccg agattcaagg ttagtcccaa cctcttcact  | 1500 |  |  |  |  |  |  |  |
| gtcccaatta aggaagcagg cgaagactgc catgccccaa catacctacc tgcggaggtg  | 1560 |  |  |  |  |  |  |  |
| gacggtgatg tcaaactcag ttccaacctg gtgattctac ctggtcaaga tctccaatat  | 1620 |  |  |  |  |  |  |  |
| gttttggcaa cctacgatac ctccagggtt gagcatgctg tggtttatta cgtttacagc  | 1680 |  |  |  |  |  |  |  |
| ccaagccgct cattttctta cttttatcct tttaggttgc ctataaaggg ggtcccaatc  | 1740 |  |  |  |  |  |  |  |
| gaactacaag tggaatgott cacatgggat caaaaactot ggtgccgtca cttctgtgtg  | 1800 |  |  |  |  |  |  |  |
| cttgcggact cagaatccgg tggacttatc actcactctg ggatggtggg catgggagtc  | 1860 |  |  |  |  |  |  |  |
| agctgcacag ctacccggga agatggaacc aatcgcagat aatgataata ggctggagcc  | 1920 |  |  |  |  |  |  |  |
| teggtggeca agettettge ecettgggee tecececage ecetectece etteetgeae  | 1980 |  |  |  |  |  |  |  |
| ccgtaccccc gtggtctttg aataaagtct gagtgggcgg caaaaaaaaaa  | 2040 |  |  |  |  |  |  |  |
| аааааааааа аааааааааа аааааааааа аааааа  | 2100 |  |  |  |  |  |  |  |
| aaaaaaaaaa aaaaaaaaaa atctag   | 2126 |  |  |  |  |  |  |  |
| <210> SEQ ID NO 44 <211> LENGTH: 2065 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |      |  |  |  |  |  |  |  |
| <400> SEQUENCE: 44   |      |  |  |  |  |  |  |  |
| tcaagctttt ggaccctcgt acagaagcta atacgactca ctatagggaa ataagagaga  | 60   |  |  |  |  |  |  |  |
| aaagaagagt aagaagaaat ataagagcca ccatgtcacc acaacgagac cggataaatg  | 120  |  |  |  |  |  |  |  |
| ccttctacaa agacaacccc catcctaagg gaagtaggat agttattaac agagaacatc  | 180  |  |  |  |  |  |  |  |
| ttatgattga tagaccttat gttttgctgg ctgttctatt cgtcatgttt ctgagcttga  | 240  |  |  |  |  |  |  |  |
| tegggttget agecattgea ggeattagae tteateggge agecatetae acegeagaga  | 300  |  |  |  |  |  |  |  |
| tocataaaag cotcagcaco aatotggatg taactaacto aatogagcat caggttaagg  | 360  |  |  |  |  |  |  |  |
| acgtgctgac accactcttc aagatcatcg gtgatgaagt gggcttgagg acacctcaga  | 420  |  |  |  |  |  |  |  |
| gattcactga cctagtgaag ttcatctctg acaagattaa attccttaat ccggacaggg  | 480  |  |  |  |  |  |  |  |

540

aatacgactt cagagatete acttggtgta teaaccegee agagagaate aaattggatt

| atgatcaata   | ctgtgcagat | gtggctgctg | aagaactcat | gaatgcattg | gtgaactcaa | 600  |  |  |
|--|------------|------------|------------|------------|------------|------|--|--|
| ctctactgga   | gaccagggca | accaatcagt | tcctagctgt | ctcaaaggga | aactgctcag | 660  |  |  |
| ggcccactac   | aatcagaggc | caattctcaa | acatgtcgct | gtccctgttg | gacttgtatt | 720  |  |  |
| taagtcgagg   | ttacaatgtg | tcatctatag | tcactatgac | atcccaggga | atgtacgggg | 780  |  |  |
| gaacttacct   | agtggaaaag | cctaatctga | gcagcaaagg | gtcagagttg | tcacaactga | 840  |  |  |
| gcatgcaccg   | agtgtttgaa | gtaggtgtta | tcagaaatcc | gggtttgggg | gctccggtat | 900  |  |  |
| tccatatgac   | aaactatctt | gagcaaccag | tcagtaatga | tttcagcaac | tgcatggtgg | 960  |  |  |
| ctttggggga   | gctcaagttc | gcagccctct | gtcacaggga | agattctatc | acaattccct | 1020 |  |  |
| atcagggatc   | agggaaaggt | gtcagcttcc | agcttgtcaa | gctaggtgtc | tggaaatccc | 1080 |  |  |
| caaccgacat   | gcaatcctgg | gtccccctat | caacggatga | tccagtgata | gacaggettt | 1140 |  |  |
| acctctcatc   | tcacagaggc | gttatcgctg | acaatcaagc | aaaatgggct | gtcccgacaa | 1200 |  |  |
| cacggacaga   | tgacaagttg | cgaatggaga | catgcttcca | gcaggcgtgt | aagggtaaaa | 1260 |  |  |
| tccaagcact   | ttgcgagaat | cccgagtgga | caccattgaa | ggataacagg | attccttcat | 1320 |  |  |
| acggggtctt   | gtctgttgat | ctgagtctga | cagttgagct | taaaatcaaa | attgtttcag | 1380 |  |  |
| gattcgggcc   | attgatcaca | cacggttcag | ggatggacct | atacaaatcc | aaccacaaca | 1440 |  |  |
| atatgtattg   | gctgactatc | ccgccaatga | agaacctggc | cttaggtgta | atcaacacat | 1500 |  |  |
| tggagtggat   | accgagattc | aaggttagtc | ccaacctctt | cactgttcca | attaaggaag | 1560 |  |  |
| caggcgagga   | ctgccatgcc | ccaacatacc | tacctgcgga | ggtggatggt | gatgtcaaac | 1620 |  |  |
| tcagttccaa   | tctggtgatt | ctacctggtc | aagatctcca | atatgttctg | gcaacctacg | 1680 |  |  |
| atacttccag   | agttgaacat | gctgtagttt | attacgttta | cagcccaagc | cgctcatttt | 1740 |  |  |
| cttactttta   | tccttttagg | ttgcctgtaa | ggggggtccc | cattgaatta | caagtggaat | 1800 |  |  |
| gcttcacatg   | ggaccaaaaa | ctctggtgcc | gtcacttctg | tgtgcttgcg | gactcagaat | 1860 |  |  |
| ctggtggaca   | tatcactcac | tctgggatgg | tgggcatggg | agtcagctgc | acagccactc | 1920 |  |  |
| gggaagatgg   | aaccagccgc | agatagtgat | aataggctgg | agcctcggtg | gccaagcttc | 1980 |  |  |
| ttgccccttg   | ggcctccccc | cagcccctcc | tccccttcct | gcacccgtac | ccccgtggtc | 2040 |  |  |
| tttgaataaa   | gtctgagtgg | gegge      |            |            |            | 2065 |  |  |
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| <400> SEQUI  | ENCE: 45   |            |            |            |            |      |  |  |
|  |            | gataaatgcc |            |            |            | 60   |  |  |
| agtaggatag   | ttattaacag | agaacatctt | atgattgata | gaccttatgt | tttgctggct | 120  |  |  |
| gttctattcg   | tcatgtttct | gagcttgatc | gggttgctag | ccattgcagg | cattagactt | 180  |  |  |
| catcgggcag   | ccatctacac | cgcagagatc | cataaaagcc | tcagcaccaa | tctggatgta | 240  |  |  |
| actaactcaa   | tcgagcatca | ggttaaggac | gtgctgacac | cactcttcaa | gatcatcggt | 300  |  |  |
| gatgaagtgg   | gcttgaggac | acctcagaga | ttcactgacc | tagtgaagtt | catctctgac | 360  |  |  |
| aagattaaat   | tccttaatcc | ggacagggaa | tacgacttca | gagateteae | ttggtgtatc | 420  |  |  |

aacccgccag agagaatcaa attggattat gatcaatact gtgcagatgt ggctgctgaa

|   |             |            |            |            | -0011011   | iueu<br>   |      |  |
|---|-------------|------------|------------|------------|------------|------------|------|--|
|   | gaactcatga  | atgcattggt | gaactcaact | ctactggaga | ccagggcaac | caatcagttc | 540  |  |
|   | ctagctgtct  | caaagggaaa | ctgctcaggg | cccactacaa | tcagaggcca | attctcaaac | 600  |  |
|   | atgtcgctgt  | ccctgttgga | cttgtattta | agtcgaggtt | acaatgtgtc | atctatagtc | 660  |  |
|   | actatgacat  | cccagggaat | gtacggggga | acttacctag | tggaaaagcc | taatctgagc | 720  |  |
|   | agcaaagggt  | cagagttgtc | acaactgagc | atgcaccgag | tgtttgaagt | aggtgttatc | 780  |  |
|   | agaaatccgg  | gtttgggggc | tccggtattc | catatgacaa | actatcttga | gcaaccagtc | 840  |  |
|   | agtaatgatt  | tcagcaactg | catggtggct | ttgggggagc | tcaagttcgc | agccctctgt | 900  |  |
|   | cacagggaag  | attctatcac | aattccctat | cagggatcag | ggaaaggtgt | cagcttccag | 960  |  |
|   | cttgtcaagc  | taggtgtctg | gaaatcccca | accgacatgc | aatcctgggt | ccccctatca | 1020 |  |
|   | acggatgatc  | cagtgataga | caggctttac | ctctcatctc | acagaggcgt | tatcgctgac | 1080 |  |
|   | aatcaagcaa  | aatgggctgt | cccgacaaca | cggacagatg | acaagttgcg | aatggagaca | 1140 |  |
|   | tgcttccagc  | aggcgtgtaa | gggtaaaatc | caagcacttt | gcgagaatcc | cgagtggaca | 1200 |  |
|   | ccattgaagg  | ataacaggat | tccttcatac | ggggtcttgt | ctgttgatct | gagtctgaca | 1260 |  |
|   | gttgagctta  | aaatcaaaat | tgtttcagga | ttcgggccat | tgatcacaca | cggttcaggg | 1320 |  |
|   | atggacctat  | acaaatccaa | ccacaacaat | atgtattggc | tgactatccc | gccaatgaag | 1380 |  |
|   | aacctggcct  | taggtgtaat | caacacattg | gagtggatac | cgagattcaa | ggttagtccc | 1440 |  |
|   | aacctcttca  | ctgttccaat | taaggaagca | ggcgaggact | gccatgcccc | aacataccta | 1500 |  |
|   | cctgcggagg  | tggatggtga | tgtcaaactc | agttccaatc | tggtgattct | acctggtcaa | 1560 |  |
|   | gatctccaat  | atgttctggc | aacctacgat | acttccagag | ttgaacatgc | tgtagtttat | 1620 |  |
|   | tacgtttaca  | gcccaagccg | ctcattttct | tacttttatc | cttttaggtt | gcctgtaagg | 1680 |  |
|   | ggggtcccca  | ttgaattaca | agtggaatgc | ttcacatggg | accaaaaact | ctggtgccgt | 1740 |  |
|   | cacttctgtg  | tgcttgcgga | ctcagaatct | ggtggacata | tcactcactc | tgggatggtg | 1800 |  |
|   | ggcatgggag  | tcagctgcac | agccactcgg | gaagatggaa | ccagccgcag | atag       | 1854 |  |
| <pre>ggcatgggag tcagctgcac agccactcgg gaagatggaa ccagccgcag atag 1854  &lt;210&gt; SEQ ID NO 46 &lt;211&gt; LENGTH: 2126 &lt;212&gt; TYPE: DNA &lt;213&gt; ORGANISM: Artificial Sequence &lt;220&gt; FEATURE: &lt;223&gt; OTHER INFORMATION: Synthetic Polynucleotide</pre> |             |            |            |            |            |            |      |  |
|   | <400> SEQUI | ENCE: 46   |            |            |            |            |      |  |
|   | ggggaaataa  | gagagaaaag | aagagtaaga | agaaatataa | gagccaccat | gtcaccacaa | 60   |  |
|   | cgagaccgga  | taaatgcctt | ctacaaagac | aacccccatc | ctaagggaag | taggatagtt | 120  |  |
|   | attaacagag  | aacatcttat | gattgataga | ccttatgttt | tgctggctgt | tctattcgtc | 180  |  |
|   | atgtttctga  | gcttgatcgg | gttgctagcc | attgcaggca | ttagacttca | tegggeagee | 240  |  |
|   | atctacaccg  | cagagatcca | taaaagcctc | agcaccaatc | tggatgtaac | taactcaatc | 300  |  |
|   | gagcatcagg  | ttaaggacgt | gctgacacca | ctcttcaaga | tcatcggtga | tgaagtgggc | 360  |  |
|   | ttgaggacac  | ctcagagatt | cactgaccta | gtgaagttca | tctctgacaa | gattaaattc | 420  |  |
|   | cttaatccgg  | acagggaata | cgacttcaga | gatctcactt | ggtgtatcaa | cccgccagag | 480  |  |
|   | agaatcaaat  | tggattatga | tcaatactgt | gcagatgtgg | ctgctgaaga | actcatgaat | 540  |  |
|   | gcattggtga  | actcaactct | actggagacc | agggcaacca | atcagttcct | agctgtctca | 600  |  |
|   | aagggaaact  | gctcagggcc | cactacaatc | agaggccaat | tctcaaacat | gtegetgtee | 660  |  |
|   |             |            |            |            |            |            |      |  |

720

ctgttggact tgtatttaag tcgaggttac aatgtgtcat ctatagtcac tatgacatcc

| cagggaatgt        | acgggggaac                       | ttacctagtg       | gaaaagccta        | atctgagcag        | caaagggtca      | 780  |
|-------------------|----------------------------------|------------------|-------------------|-------------------|-----------------|------|
| gagttgtcac        | aactgagcat                       | gcaccgagtg       | tttgaagtag        | gtgttatcag        | aaatccgggt      | 840  |
| ttgggggctc        | cggtattcca                       | tatgacaaac       | tatcttgago        | aaccagtcag        | taatgatttc      | 900  |
| agcaactgca        | tggtggcttt                       | gggggagctc       | aagttcgcag        | ccctctgtca        | cagggaagat      | 960  |
| tctatcacaa        | ttccctatca                       | gggatcaggg       | aaaggtgtca        | gcttccagct        | tgtcaagcta      | 1020 |
| ggtgtctgga        | aatccccaac                       | cgacatgcaa       | teetgggtee        | ccctatcaac        | ggatgatcca      | 1080 |
| gtgatagaca        | ggctttacct                       | ctcatctcac       | agaggcgtta        | . tcgctgacaa      | tcaagcaaaa      | 1140 |
| tgggctgtcc        | cgacaacacg                       | gacagatgac       | aagttgcgaa        | . tggagacatg      | cttccagcag      | 1200 |
| gcgtgtaagg        | gtaaaatcca                       | agcactttgc       | gagaatcccg        | agtggacacc        | attgaaggat      | 1260 |
| aacaggattc        | cttcatacgg                       | ggtcttgtct       | gttgatctga        | gtctgacagt        | tgagcttaaa      | 1320 |
| atcaaaattg        | tttcaggatt                       | cgggccattg       | atcacacaco        | gttcagggat        | ggacctatac      | 1380 |
| aaatccaacc        | acaacaatat                       | gtattggctg       | actatecego        | caatgaagaa        | cctggcctta      | 1440 |
| ggtgtaatca        | acacattgga                       | gtggataccg       | agattcaagg        | ttagtcccaa        | cctcttcact      | 1500 |
| gttccaatta        | aggaagcagg                       | cgaggactgc       | catgccccaa        | catacctacc        | tgcggaggtg      | 1560 |
| gatggtgatg        | tcaaactcag                       | ttccaatctg       | gtgattctac        | ctggtcaaga        | tctccaatat      | 1620 |
| gttctggcaa        | cctacgatac                       | ttccagagtt       | gaacatgcto        | tagtttatta        | cgtttacagc      | 1680 |
| ccaagccgct        | cattttctta                       | cttttatcct       | tttaggttgc        | ctgtaagggg        | ggtccccatt      | 1740 |
| gaattacaag        | tggaatgctt                       | cacatgggac       | caaaaactct        | ggtgccgtca        | cttctgtgtg      | 1800 |
| cttgcggact        | cagaatctgg                       | tggacatatc       | actcactctc        | ggatggtggg        | catgggagtc      | 1860 |
| agctgcacag        | ccactcggga                       | agatggaacc       | agccgcagat        | agtgataata        | ggctggagcc      | 1920 |
| tcggtggcca        | agcttcttgc                       | cccttgggcc       | tcccccago         | ccctcctccc        | cttcctgcac      | 1980 |
| ccgtaccccc        | gtggtctttg                       | aataaagtct       | gagtgggcgg        | caaaaaaaaa        | aaaaaaaaa       | 2040 |
| aaaaaaaaa         | aaaaaaaaa                        | aaaaaaaaa        | aaaaaaaaa         | aaaaaaaaa         | aaaaaaaaa       | 2100 |
| aaaaaaaaa         | aaaaaaaaa                        | atctag           |                   |                   |                 | 2126 |
| <220> FEATU       | TH: 550<br>: PRT<br>NISM: Artif: | _                |                   | iđe               |                 |      |
| <400> SEQUI       | ENCE: 47                         |                  |                   |                   |                 |      |
| Met Gly Lev<br>1  | ı Lys Val A:<br>5                | en Val Ser       | Ala Val Phe<br>10 | Met Ala Val       | Leu Leu<br>15   |      |
| Thr Leu Gli       | n Thr Pro A                      |                  | Ile His Trp<br>25 | Gly Asn Let<br>30 | ı Ser Lys       |      |
| Ile Gly Val       | l Val Gly I                      | le Gly Ser<br>40 | Ala Ser Tyr       | Lys Val Met<br>45 | Thr Arg         |      |
| Ser Ser His       | s Gln Ser Le                     | eu Val Ile<br>55 | Lys Leu Met       | Pro Asn Ile       | Thr Leu         |      |
| Leu Asn Ası<br>65 | n Cys Thr A:                     | -                | Ile Ala Glu<br>75 | . Tyr Arg Arg     | g Leu Leu<br>80 |      |
| Arg Thr Val       | l Leu Glu P:<br>85               | ro Ile Arg       | Asp Ala Leu<br>90 | . Asn Ala Met     | Thr Gln<br>95   |      |

Asn Ile Arg Pro Val Gln Ser Val Ala Ser Ser Arg Arg His Lys Arg 100 105 110

| Phe        | Ala        | Gly<br>115 | Val        | Val        | Leu        | Ala        | Gly<br>120 | Ala        | Ala        | Leu        | Gly        | Val<br>125 | Ala        | Thr        | Ala        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Ala        | Gln<br>130 | Ile        | Thr        | Ala        | Gly        | Ile<br>135 | Ala        | Leu        | His        | Arg        | Ser<br>140 | Met        | Leu        | Asn        | Ser        |
| Gln<br>145 | Ala        | Ile        | Asp        | Asn        | Leu<br>150 | Arg        | Ala        | Ser        | Leu        | Glu<br>155 | Thr        | Thr        | Asn        | Gln        | Ala<br>160 |
| Ile        | Glu        | Ala        | Ile        | Arg<br>165 | Gln        | Ala        | Gly        | Gln        | Glu<br>170 | Met        | Ile        | Leu        | Ala        | Val<br>175 | Gln        |
| Gly        | Val        | Gln        | Asp<br>180 | Tyr        | Ile        | Asn        | Asn        | Glu<br>185 | Leu        | Ile        | Pro        | Ser        | Met<br>190 | Asn        | Gln        |
| Leu        | Ser        | Суз<br>195 | Asp        | Leu        | Ile        | Gly        | Gln<br>200 | Lys        | Leu        | Gly        | Leu        | Lys<br>205 | Leu        | Leu        | Arg        |
| Tyr        | Tyr<br>210 | Thr        | Glu        | Ile        | Leu        | Ser<br>215 | Leu        | Phe        | Gly        | Pro        | Ser<br>220 | Leu        | Arg        | Asp        | Pro        |
| Ile<br>225 | Ser        | Ala        | Glu        | Ile        | Ser<br>230 | Ile        | Gln        | Ala        | Leu        | Ser<br>235 | Tyr        | Ala        | Leu        | Gly        | Gly<br>240 |
| Asp        | Ile        | Asn        | Lys        | Val<br>245 | Leu        | Glu        | Lys        | Leu        | Gly<br>250 | Tyr        | Ser        | Gly        | Gly        | Asp<br>255 | Leu        |
| Leu        | Gly        | Ile        | Leu<br>260 | Glu        | Ser        | Arg        | Gly        | Ile<br>265 | Lys        | Ala        | Arg        | Ile        | Thr<br>270 | His        | Val        |
| Asp        | Thr        | Glu<br>275 | Ser        | Tyr        | Phe        | Ile        | Val<br>280 | Leu        | Ser        | Ile        | Ala        | Tyr<br>285 | Pro        | Thr        | Leu        |
| Ser        | Glu<br>290 | Ile        | Lys        | Gly        | Val        | Ile<br>295 | Val        | His        | Arg        | Leu        | Glu<br>300 | Gly        | Val        | Ser        | Tyr        |
| Asn<br>305 | Ile        | Gly        | Ser        | Gln        | Glu<br>310 | Trp        | Tyr        | Thr        | Thr        | Val<br>315 | Pro        | ГÀа        | Tyr        | Val        | Ala<br>320 |
| Thr        | Gln        | Gly        | Tyr        | Leu<br>325 | Ile        | Ser        | Asn        | Phe        | 330        | Glu        | Ser        | Ser        | CÀa        | Thr<br>335 | Phe        |
| Met        | Pro        | Glu        | Gly<br>340 | Thr        | Val        | CÀa        | Ser        | Gln<br>345 | Asn        | Ala        | Leu        | Tyr        | Pro<br>350 | Met        | Ser        |
| Pro        | Leu        | Leu<br>355 | Gln        | Glu        | CAa        | Leu        | Arg<br>360 | Gly        | Ser        | Thr        | Lys        | Ser<br>365 | CÀa        | Ala        | Arg        |
| Thr        | Leu<br>370 | Val        | Ser        | Gly        | Ser        | Phe<br>375 | Gly        | Asn        | Arg        | Phe        | Ile<br>380 | Leu        | Ser        | Gln        | Gly        |
| Asn<br>385 | Leu        | Ile        | Ala        | Asn        | 390<br>CAa | Ala        | Ser        | Ile        | Leu        | Cys<br>395 | ГÀа        | CAa        | Tyr        | Thr        | Thr<br>400 |
| Gly        | Thr        | Ile        | Ile        | Asn<br>405 | Gln        | Asp        | Pro        | Asp        | Lys<br>410 | Ile        | Leu        | Thr        | Tyr        | Ile<br>415 | Ala        |
| Ala        | Asp        | Arg        | Cys<br>420 | Pro        | Val        | Val        | Glu        | Val<br>425 | Asn        | Gly        | Val        | Thr        | Ile<br>430 | Gln        | Val        |
| Gly        | Ser        | Arg<br>435 | Arg        | Tyr        | Pro        | Asp        | Ala<br>440 | Val        | Tyr        | Leu        | His        | Arg<br>445 | Ile        | Asp        | Leu        |
| Gly        | Pro<br>450 | Pro        | Ile        | Ser        | Leu        | Glu<br>455 | Arg        | Leu        | Asp        | Val        | Gly<br>460 | Thr        | Asn        | Leu        | Gly        |
| Asn<br>465 | Ala        | Ile        | Ala        | Lys        | Leu<br>470 | Glu        | Asp        | Ala        | Lys        | Glu<br>475 | Leu        | Leu        | Glu        | Ser        | Ser<br>480 |
| Asp        | Gln        | Ile        | Leu        | Arg<br>485 | Ser        | Met        | Lys        | Gly        | Leu<br>490 | Ser        | Ser        | Thr        | Ser        | Ile<br>495 | Val        |
| Tyr        | Ile        | Leu        | Ile<br>500 | Ala        | Val        | СЛа        | Leu        | Gly<br>505 | Gly        | Leu        | Ile        | Gly        | Ile<br>510 | Pro        | Thr        |
| Leu        | Ile        | Сув<br>515 | Cys        | Сла        | Arg        | Gly        | Arg<br>520 | Cys        | Asn        | Lys        | Lys        | Gly<br>525 | Glu        | Gln        | Val        |

Gly Met Ser Arg Pro Gly Leu Lys Pro Asp Leu Thr Gly Thr Ser Lys 535 Ser Tyr Val Arg Ser Leu <210> SEQ ID NO 48 <211> LENGTH: 550 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 48 Met Gly Leu Lys Val Asn Val Ser Val Ile Phe Met Ala Val Leu Leu Thr Leu Gln Thr Pro Thr Gly Gln Ile His Trp Gly Asn Leu Ser Lys Ile Gly Val Val Gly Val Gly Ser Ala Ser Tyr Lys Val Met Thr Arg 40 Ser Ser His Gln Ser Leu Val Ile Lys Leu Met Pro Asn Ile Thr Leu 55 Leu Asn Asn Cys Thr Arg Val Gly Ile Ala Glu Tyr Arg Arg Leu Leu Arg Thr Val Leu Glu Pro Ile Arg Asp Ala Leu Asn Ala Met Thr Gln Asn Ile Arg Pro Val Gln Ser Val Ala Ser Ser Arg Arg His Lys Arg 105 Phe Ala Gly Val Val Leu Ala Gly Ala Ala Leu Gly Val Ala Thr Ala 120 Ala Gln Ile Thr Ala Gly Ile Ala Leu His Gln Ser Met Leu Asn Ser Gln Ala Ile Asp Asn Leu Arg Ala Ser Leu Glu Thr Thr Asn Gln Ala 155 Ile Glu Ala Ile Arg Gln Ala Gly Gln Glu Met Ile Leu Ala Val Gln Gly Val Gln Asp Tyr Ile Asn Asn Glu Leu Ile Pro Ser Met Asn Gln Leu Ser Cys Asp Leu Ile Gly Gln Lys Leu Gly Leu Lys Leu Leu Arg Tyr Tyr Thr Glu Ile Leu Ser Leu Phe Gly Pro Ser Leu Arg Asp Pro Ile Ser Ala Glu Ile Ser Ile Gln Ala Leu Ser Tyr Ala Leu Gly Gly Asp Ile Asn Lys Val Leu Glu Lys Leu Gly Tyr Ser Gly Gly Asp Leu Leu Gly Ile Leu Glu Ser Arg Gly Ile Lys Ala Arg Ile Thr His Val 265 Asp Thr Glu Ser Tyr Phe Ile Val Leu Ser Ile Ala Tyr Pro Thr Leu Ser Glu Ile Lys Gly Val Ile Val His Arg Leu Glu Gly Val Ser Tyr 295 Asn Ile Gly Ser Gln Glu Trp Tyr Thr Thr Val Pro Lys Tyr Val Ala 310 Thr Gln Gly Tyr Leu Ile Ser Asn Phe Asp Glu Ser Ser Cys Thr Phe 330

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Met Pro Glu Gly Thr Val Cys Ser Gln Asn Ala Leu Tyr Pro Met Ser Pro Leu Leu Gln Glu Cys Leu Arg Gly Ser Thr Lys Ser Cys Ala Arg Thr Leu Val Ser Gly Ser Phe Gly Asn Arg Phe Ile Leu Ser Gln Gly Asn Leu Ile Ala Asn Cys Ala Ser Ile Leu Cys Lys Cys Tyr Thr Thr Gly Thr Ile Ile Asn Gln Asp Pro Asp Lys Ile Leu Thr Tyr Ile Ala Ala Asp His Cys Pro Val Val Glu Val Asn Gly Val Thr Ile Gln Val Gly Ser Arg Arg Tyr Pro Asp Ala Val Tyr Leu His Arg Ile Asp Leu Gly Pro Pro Ile Ser Leu Glu Arg Leu Asp Val Gly Thr Asn Leu Gly 455 Asn Ala Ile Ala Lys Leu Glu Asp Ala Lys Glu Leu Leu Glu Ser Ser Asp Gln Ile Leu Arg Ser Met Lys Gly Leu Ser Ser Thr Ser Ile Val Tyr Ile Leu Ile Ala Val Cys Leu Gly Gly Leu Ile Gly Ile Pro Ala 505 Leu Ile Cys Cys Cys Arg Gly Arg Cys Asn Lys Lys Gly Glu Gln Val 520 Gly Met Ser Arg Pro Gly Leu Lys Pro Asp Leu Thr Gly Thr Ser Lys 535 Ser Tyr Val Arg Ser Leu <210> SEQ ID NO 49 <211> LENGTH: 617 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 49 Met Ser Pro Gln Arg Asp Arg Ile Asn Ala Phe Tyr Lys Asp Asn Pro Tyr Pro Lys Gly Ser Arg Ile Val Ile Asn Arg Glu His Leu Met Ile Asp Arg Pro Tyr Val Leu Leu Ala Val Leu Phe Val Met Phe Leu Ser Leu Ile Gly Leu Leu Ala Ile Ala Gly Ile Arg Leu His Arg Ala Ala 50 55 60 Ile Tyr Thr Ala Glu Ile His Lys Ser Leu Ser Thr Asn Leu Asp Val Thr Asn Ser Ile Glu His Gln Val Lys Asp Val Leu Thr Pro Leu Phe Lys Ile Ile Gly Asp Glu Val Gly Leu Arg Thr Pro Gln Arg Phe Thr 105 Asp Leu Val Lys Phe Ile Ser Asp Lys Ile Lys Phe Leu Asn Pro Asp 120 Arg Glu Tyr Asp Phe Arg Asp Leu Thr Trp Cys Ile Asn Pro Pro Glu 135

| Arg        | Ile        | Lys        | Leu        | Asp        | Tyr<br>150 | Asp        | Gln        | Tyr        | Cys        | Ala<br>155 | Asp        | Val        | Ala        | Ala        | Glu<br>160 |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Glu        | Leu        | Met        | Asn        | Ala<br>165 | Leu        | Val        | Asn        | Ser        | Thr        | Leu        | Leu        | Glu        | Thr        | Arg<br>175 | Thr        |
| Thr        | Thr        | Gln        | Phe<br>180 | Leu        | Ala        | Val        | Ser        | Lys<br>185 | Gly        | Asn        | Сув        | Ser        | Gly<br>190 | Pro        | Thr        |
| Thr        | Ile        | Arg<br>195 | Gly        | Gln        | Phe        | Ser        | Asn<br>200 | Met        | Ser        | Leu        | Ser        | Leu<br>205 | Leu        | Asp        | Leu        |
| Tyr        | Leu<br>210 | Gly        | Arg        | Gly        | Tyr        | Asn<br>215 | Val        | Ser        | Ser        | Ile        | Val<br>220 | Thr        | Met        | Thr        | Ser        |
| Gln<br>225 | Gly        | Met        | Tyr        | Gly        | Gly<br>230 | Thr        | Tyr        | Leu        | Val        | Glu<br>235 | Lys        | Pro        | Asn        | Leu        | Asn<br>240 |
| Ser        | ГЛа        | Gly        | Ser        | Glu<br>245 | Leu        | Ser        | Gln        | Leu        | Ser<br>250 | Met        | Tyr        | Arg        | Val        | Phe<br>255 | Glu        |
| Val        | Gly        | Val        | Ile<br>260 | Arg        | Asn        | Pro        | Gly        | Leu<br>265 | Gly        | Ala        | Pro        | Val        | Phe<br>270 | His        | Met        |
| Thr        | Asn        | Tyr<br>275 | Phe        | Glu        | Gln        | Pro        | Val<br>280 | Ser        | Asn        | Gly        | Leu        | Gly<br>285 | Asn        | CÀa        | Met        |
| Val        | Ala<br>290 | Leu        | Gly        | Glu        | Leu        | Lys<br>295 | Leu        | Ala        | Ala        | Leu        | 300<br>Cys | His        | Gly        | Asp        | Asp        |
| Ser<br>305 | Ile        | Ile        | Ile        | Pro        | Tyr<br>310 | Gln        | Gly        | Ser        | Gly        | Lys<br>315 | Gly        | Val        | Ser        | Phe        | Gln<br>320 |
| Leu        | Val        | Lys        | Leu        | Gly<br>325 | Val        | Trp        | Lys        | Ser        | Pro<br>330 | Thr        | Asp        | Met        | Gln        | Ser<br>335 | Trp        |
| Val        | Pro        | Leu        | Ser<br>340 | Thr        | Asp        | Asp        | Pro        | Val<br>345 | Val        | Asp        | Arg        | Leu        | Tyr<br>350 | Leu        | Ser        |
| Ser        | His        | Arg<br>355 | Gly        | Val        | Ile        | Ala        | Asp<br>360 | Asn        | Gln        | Ala        | Lys        | Trp<br>365 | Ala        | Val        | Pro        |
| Thr        | Thr<br>370 | Arg        | Thr        | Asp        | Asp        | Lys<br>375 | Leu        | Arg        | Met        | Glu        | Thr<br>380 | CAa        | Phe        | Gln        | Gln        |
| Ala<br>385 | Сув        | Lys        | Gly        | Lys        | Ile<br>390 | Gln        | Ala        | Leu        | CAa        | Glu<br>395 | Asn        | Pro        | Glu        | Trp        | Val<br>400 |
| Pro        | Leu        | ГЛа        | Asp        | Asn<br>405 | Arg        | Ile        | Pro        | Ser        | Tyr<br>410 | Gly        | Val        | Leu        | Ser        | Val<br>415 | Asp        |
| Leu        | Ser        | Leu        | Thr<br>420 | Val        | Glu        | Leu        | Lys        | Ile<br>425 | ГЛа        | Ile        | Ala        | Ser        | Gly<br>430 | Phe        | Gly        |
| Pro        |            | Ile<br>435 |            | His        | Gly        | Ser        | Gly<br>440 |            | Asp        | Leu        |            | Lys<br>445 | Ser        | Asn        | Cys        |
| Asn        | Asn<br>450 | Val        | Tyr        | Trp        | Leu        | Thr<br>455 | Ile        | Pro        | Pro        | Met        | Arg<br>460 | Asn        | Leu        | Ala        | Leu        |
| Gly<br>465 | Val        | Ile        | Asn        | Thr        | Leu<br>470 | Glu        | Trp        | Ile        | Pro        | Arg<br>475 | Phe        | ГÀа        | Val        | Ser        | Pro<br>480 |
| Asn        | Leu        | Phe        | Thr        | Val<br>485 | Pro        | Ile        | Lys        | Glu        | Ala<br>490 | Gly        | Glu        | Asp        | Cha        | His<br>495 | Ala        |
| Pro        | Thr        | Tyr        | Leu<br>500 | Pro        | Ala        | Glu        | Val        | Asp<br>505 | Gly        | Asp        | Val        | Lys        | Leu<br>510 | Ser        | Ser        |
| Asn        | Leu        | Val<br>515 | Ile        | Leu        | Pro        | Gly        | Gln<br>520 | Asp        | Leu        | Gln        | Tyr        | Val<br>525 | Leu        | Ala        | Thr        |
| Tyr        | Asp<br>530 | Thr        | Ser        | Arg        | Val        | Glu<br>535 | His        | Ala        | Val        | Val        | Tyr<br>540 | Tyr        | Val        | Tyr        | Ser        |
| Pro<br>545 | Ser        | Arg        | Ser        | Phe        | Ser<br>550 | Tyr        | Phe        | Tyr        | Pro        | Phe<br>555 | Arg        | Leu        | Pro        | Ile        | Lys<br>560 |
| Gly        | Val        | Pro        | Ile        | Glu        | Leu        | Gln        | Val        | Glu        | Сув        | Phe        | Thr        | Trp        | Asp        | Gln        | Lys        |

| -continued |
|------------|
|            |

Leu Trp Cys Arg His Phe Cys Val Leu Ala Asp Ser Glu Ser Gly Gly 585 Leu Ile Thr His Ser Gly Met Val Gly Met Gly Val Ser Cys Thr Ala 600 Thr Arg Glu Asp Gly Thr Asn Arg Arg <210> SEQ ID NO 50 <211> LENGTH: 617 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 50 Met Ser Pro Gln Arg Asp Arg Ile Asn Ala Phe Tyr Lys Asp Asn Pro His Pro Lys Gly Ser Arg Ile Val Ile Asn Arg Glu His Leu Met Ile Asp Arg Pro Tyr Val Leu Leu Ala Val Leu Phe Val Met Phe Leu Ser Leu Ile Gly Leu Leu Ala Ile Ala Gly Ile Arg Leu His Arg Ala Ala 55 Ile Tyr Thr Ala Glu Ile His Lys Ser Leu Ser Thr Asn Leu Asp Val 70 Thr Asn Ser Ile Glu His Gln Val Lys Asp Val Leu Thr Pro Leu Phe Lys Ile Ile Gly Asp Glu Val Gly Leu Arg Thr Pro Gln Arg Phe Thr 105 Asp Leu Val Lys Phe Ile Ser Asp Lys Ile Lys Phe Leu Asn Pro Asp  $\hbox{Arg Glu Tyr Asp Phe Arg Asp Leu Thr Trp Cys Ile Asn Pro Pro Glu } \\$ Arg Ile Lys Leu Asp Tyr Asp Gln Tyr Cys Ala Asp Val Ala Ala Glu Glu Leu Met Asn Ala Leu Val Asn Ser Thr Leu Leu Glu Thr Arg Ala 170 Thr Asn Gln Phe Leu Ala Val Ser Lys Gly Asn Cys Ser Gly Pro Thr Thr Ile Arg Gly Gln Phe Ser Asn Met Ser Leu Ser Leu Leu Asp Leu Tyr Leu Ser Arg Gly Tyr Asn Val Ser Ser Ile Val Thr Met Thr Ser Gln Gly Met Tyr Gly Gly Thr Tyr Leu Val Glu Lys Pro Asn Leu Ser Ser Lys Gly Ser Glu Leu Ser Gln Leu Ser Met His Arg Val Phe Glu Val Gly Val Ile Arg Asn Pro Gly Leu Gly Ala Pro Val Phe His Met 265 Thr Asn Tyr Leu Glu Gln Pro Val Ser Asn Asp Phe Ser Asn Cys Met Val Ala Leu Gly Glu Leu Lys Phe Ala Ala Leu Cys His Arg Glu Asp Ser Ile Thr Ile Pro Tyr Gln Gly Ser Gly Lys Gly Val Ser Phe Gln

| 305 310 315  | 320                          |
|--|------------------------------|
| Leu Val Lys Leu Gly Val Trp Lys Ser Pro Thi<br>325 330   | r Asp Met Gln Ser Trp<br>335 |
| Val Pro Leu Ser Thr Asp Asp Pro Val Ile Asp 340 345  | Arg Leu Tyr Leu Ser<br>350   |
| Ser His Arg Gly Val Ile Ala Asp Asn Gln Ala<br>355 360   | a Lys Trp Ala Val Pro<br>365 |
| Thr Thr Arg Thr Asp Asp Lys Leu Arg Met Glu<br>370 375   | ı Thr Cys Phe Gln Gln<br>380 |
| Ala Cys Lys Gly Lys Ile Gln Ala Leu Cys Glu<br>385 390 395   |                              |
| Pro Leu Lys Asp Asn Arg Ile Pro Ser Tyr Gly<br>405 410   | y Val Leu Ser Val Asp<br>415 |
| Leu Ser Leu Thr Val Glu Leu Lys Ile Lys Ile<br>420 425   | e Val Ser Gly Phe Gly<br>430 |
| Pro Leu Ile Thr His Gly Ser Gly Met Asp Leu<br>435 440   | ı Tyr Lys Ser Asn His<br>445 |
| Asn Asn Met Tyr Trp Leu Thr Ile Pro Pro Met 450 455  | Lys Asn Leu Ala Leu<br>460   |
| Gly Val Ile Asn Thr Leu Glu Trp Ile Pro Arc<br>465 470 475   |                              |
| Asn Leu Phe Thr Val Pro Ile Lys Glu Ala Gly<br>485 490   | y Glu Asp Cys His Ala<br>495 |
| Pro Thr Tyr Leu Pro Ala Glu Val Asp Gly Asp<br>500 505   | o Val Lys Leu Ser Ser<br>510 |
| Asn Leu Val Ile Leu Pro Gly Gln Asp Leu Glr<br>515 520   | n Tyr Val Leu Ala Thr<br>525 |
| Tyr Asp Thr Ser Arg Val Glu His Ala Val Val  | l Tyr Tyr Val Tyr Ser<br>540 |
| Pro Ser Arg Ser Phe Ser Tyr Phe Tyr Pro Phe 545 550 555  |                              |
| Gly Val Pro Ile Glu Leu Gln Val Glu Cys Phe<br>565 570   | e Thr Trp Asp Gln Lys<br>575 |
| Leu Trp Cys Arg His Phe Cys Val Leu Ala Asp<br>580 585   | Ser Glu Ser Gly Gly<br>590   |
| His Ile Thr His Ser Gly Met Val Gly Met Gly 595 600  | y Val Ser Cys Thr Ala<br>605 |
| Thr Arg Glu Asp Gly Thr Ser Arg Arg<br>610 615   |                              |
| <210> SEQ ID NO 51 <211> LENGTH: 1729 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucl | Leotide                      |
| tcaagctttt ggaccctcgt acagaagcta atacgactca  | a ctatagggaa ataagagaga 60   |
| aaagaagagt aagaagaaat ataagagcca ccatggcaca  | a agtcattaat acaaacagcc 120  |
| tgtcgctgtt gacccagaat aacctgaaca aatcccagtc  | c cgcactgggc actgctatcg 180  |
| agegtttgte tteeggtetg egtateaaca gegegaaaga  | a cgatgcggca ggacaggcga 240  |
| ttgctaaccg ttttaccgcg aacatcaaag gtctgactca  | a ggcttcccgt aacgctaacg 300  |

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| -concinued   |      |
|--|------|
| acggtatete cattgegeag accaetgaag gegegetgaa egaaateaac aacaaeetge  | 360  |
| agegtgtgeg tgaactggeg gttcagtetg egaatggtac taacteecag tetgaceteg  | 420  |
| actocatoca ggotgaaato accoagogoo tgaacgaaat cgaccgtgta tooggocaga  | 480  |
| ctcagttcaa cggcgtgaaa gtcctggcgc aggacaacac cctgaccatc caggttggtg  | 540  |
| ccaacgacgg tgaaactatc gatattgatt taaaagaaat cagctctaaa acactgggac  | 600  |
| ttgataagct taatgtccaa gatgcctaca ccccgaaaga aactgctgta accgttgata  | 660  |
| aaactaccta taaaaatggt acagatccta ttacagccca gagcaatact gatatccaaa  | 720  |
| ctgcaattgg cggtggtgca acgggggtta ctggggctga tatcaaattt aaagatggtc  | 780  |
| aatactattt agatgttaaa ggcggtgctt ctgctggtgt ttataaagcc acttatgatg  | 840  |
| aaactacaaa gaaagttaat attgatacga ctgataaaac teegttggca actgeggaag  | 900  |
| ctacagctat tcggggaacg gccactataa cccacaacca aattgctgaa gtaacaaaag  | 960  |
| agggtgttga tacgaccaca gttgcggctc aacttgctgc agcaggggtt actggcgccg  | 1020 |
| ataaggacaa tactagcctt gtaaaactat cgtttgagga taaaaacggt aaggttattg  | 1080 |
| atggtggcta tgcagtgaaa atgggcgacg atttctatgc cgctacatat gatgagaaaa  | 1140 |
| caggtgcaat tactgctaaa accactactt atacagatgg tactggcgtt gctcaaactg  | 1200 |
| gagctgtgaa atttggtggc gcaaatggta aatctgaagt tgttactgct accgatggta  | 1260 |
| agacttactt agcaagcgac cttgacaaac ataacttcag aacaggcggt gagcttaaag  | 1320 |
| aggttaatac agataagact gaaaacccac tgcagaaaat tgatgctgcc ttggcacagg  | 1380 |
| ttgatacact tcgttctgac ctgggtgcgg ttcagaaccg tttcaactcc gctatcacca  | 1440 |
| acctgggcaa taccgtaaat aacctgtctt ctgcccgtag ccgtatcgaa gattccgact  | 1500 |
| acgcaaccga agtctccaac atgtctcgcg cgcagattct gcagcaggcc ggtacctccg  | 1560 |
| ttctggcgca ggcgaaccag gttccgcaaa acgtcctctc tttactgcgt tgataatagg  | 1620 |
| ctggagcete ggtggceatg ettettgeee ettgggeete eecceageee etceteeeet  | 1680 |
| teetgeacee gtaceeeegt ggtetttgaa taaagtetga gtgggegge  | 1729 |
| <210> SEQ ID NO 52 <211> LENGTH: 1518 <212> TYPE: DNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |      |
| <400> SEQUENCE: 52   |      |
| atggcacaag tcattaatac aaacagcctg tcgctgttga cccagaataa cctgaacaaa  | 60   |
| teccagteeg caetgggeae tgetategag egtttgtett eeggtetgeg tateaacage  | 120  |
| gcgaaagacg atgcggcagg acaggcgatt gctaaccgtt ttaccgcgaa catcaaaggt  | 180  |
| ctgactcagg cttcccgtaa cgctaacgac ggtatctcca ttgcgcagac cactgaaggc  | 240  |
| gcgctgaacg aaatcaacaa caacctgcag cgtgtgcgtg aactggcggt tcagtctgcg  | 300  |
| aatggtacta acteecagte tgacetegae tecatecagg etgaaateae eeagegeetg  | 360  |
| aacgaaatcg accgtgtatc cggccagact cagttcaacg gcgtgaaagt cctggcgcag  | 420  |
| gacaacaccc tgaccatcca ggttggtgcc aacgacggtg aaactatcga tattgattta  | 480  |
| aaagaaatca gctctaaaac actgggactt gataagctta atgtccaaga tgcctacacc  | 540  |
| ccgaaagaaa ctgctgtaac cgttgataaa actacctata aaaatggtac agatcctatt  | 600  |

660

acageceaga geaataetga tatecaaaet geaattggeg gtggtgeaae gggggttaet

| ggggctgata tcaaatttaa agatggtcaa tactatttag atgttaaagg cggtgcttct  | 720  |
|--|--|
| gctggtgttt ataaagccac ttatgatgaa actacaaaga aagttaatat tgatacgact  | 780  |
| gataaaacte egttggcaac tgeggaaget acagetatte ggggaacgge cactataace  | 840  |
| cacaaccaaa ttgctgaagt aacaaaagag ggtgttgata cgaccacagt tgcggctcaa  | 900  |
| cttgctgcag caggggttac tggcgccgat aaggacaata ctagccttgt aaaactatcg  | 960  |
| tttgaggata aaaacggtaa ggttattgat ggtggctatg cagtgaaaat gggcgacgat  | 1020   |
| ttctatgccg ctacatatga tgagaaaaca ggtgcaatta ctgctaaaac cactacttat  | 1080   |
| acagatggta ctggcgttgc tcaaactgga gctgtgaaat ttggtggcgc aaatggtaaa  | 1140   |
| totgaagttg ttactgotac ogatggtaag acttacttag caagogacot tgacaaacat  | 1200   |
| aacttcagaa caggcggtga gcttaaagag gttaatacag ataagactga aaacccactg  | 1260   |
| cagaaaattg atgctgcctt ggcacaggtt gatacacttc gttctgacct gggtgcggtt  | 1320   |
| cagaaccgtt tcaactccgc tatcaccaac ctgggcaata ccgtaaataa cctgtcttct  | 1380   |
| gecegtagee gtategaaga tteegaetae geaaeegaag teteeaaeat gtetegegeg  | 1440   |
| cagattetge ageaggeegg taceteegtt etggegeagg egaaceaggt teegeaaaac  | 1500   |
| gtcctctctt tactgcgt  | 1518   |
| <210> SEQ ID NO 53 <211> LENGTH: 1790 <212> TYPE: RNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide <400> SEQUENCE: 53  |  |
| 1100 DEGUERCE. 33  |  |
| dddaaallaa dadadaaad aadadlaada adaallallaa dadccaccall ddcacaadlic  | 60   |
| ggggaaauaa gagagaaaag aagaguaaga agaaauauaa gagccaccau ggcacaaguc  | 60<br>120  |
| auuaauacaa acagecugue geuguugaee eagaauaaee ugaacaaaue eeagueegea  | 120  |
| auuaauacaa acagecugue geuguugace cagaauaace ugaacaaaue ecagueegea cugggeacug cuauegageg uuugueuuce ggueugegua ucaacagege gaaagaegau  | 120<br>180   |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca<br>cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau<br>gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu  | 120<br>180<br>240  |
| auuaauacaa acagecugue geuguugace cagaauaace ugaacaaaue ecagueegea cugggeacug cuaucgageg uuugueuuce ggucugegua ucaacagege gaaagacgau gegggeaggac aggeegauuge uaaceguuuu acegegaaca ucaaaggueu gacucaggeu ucceguaacg cuaacgacgg uaucuecauu gegeagacea cugaaggege geugaacgaa  | 120<br>180<br>240<br>300   |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac  | 120<br>180<br>240<br>300<br>360  |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgggaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac  | 120<br>180<br>240<br>300<br>360<br>420   |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu uccccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug   | 120<br>180<br>240<br>300<br>360<br>420   |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccgguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc   | 120<br>180<br>240<br>300<br>360<br>420<br>480  |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540   |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccgguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugcaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac ugggacuuga uaagcuuaau guccaagaug ccuacaccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600  |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660                                     |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccgguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugcaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac ugggacuuga uaagcuuaau guccaagaug ccuacaccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600  |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaacaa ucaaaggucu gacucaggcu uccccguaacg cuaacgacgg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaaacac ugggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc aauacugaua uccaaacug gggguuacugg ggugaaacgg ggguuacugg ggcugauauc   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660                                     |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugcaaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc aauacugaua uccaaacugc aauuggcggu ggugcaacgg ggguuacugg ggcugauauc aaauuuaaag auggucaaua cuauuuagau guuaaaggcg gugcuucugc ugguguuuau   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660<br>720                              |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc aauacugaua uccaaacugc aauuggcggu ggugcaacgg ggguuacugg ggcugauauc aaauuuaaag auggucaaua cuauuuagau guuaaaggcg gugcuucugc ugguguuuau aaagccacuu augaugaaac uacaaagaaa guuaauauug auacgacuga uaaaacuccg   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>660<br>720<br>780<br>840                       |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaacaa ucaaaggucu gacucaggcu uccccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac cguguauccg gccagacuca guucaacggc gugaaagucc agcgcugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac ugggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag ggguuacugg ggcugauauc aaauuuaaaa auggucaaua cuauuuagau guuaaaggcg gugcuucugc ugguguuuau aaagccacuu augaugaaac uacaaagaaa guuaauauug auacgacuga uaaaacuccg uuggcaacug cggaagcuac agcuauucgg ggaacggcca cuauaacca caaccaaauu  | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>660<br>720<br>780<br>840<br>900                |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaaca ucaaaggucu gacucaggcu ucccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac ucccagucug accucgacuc cauccaggcu gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugcaaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag auccuauuac agcccagagc aauacugaua uccaaacugc aauuggcggu ggugcaacgg ggguuacugg ggcugauauc aaauuuaaag auggucaaua cuauuuagau guuaaaggcg gugcuucugc ugguguuuau aaagccacuu augaugaaac uacaaagaaa guuaauauug auacgacuga uaaaacuccg uuggcaacug cggaagcuac agcuauucgg ggaacggcca cuauaaccca caaccaaauu gcugaaguaa caaaagaggg uguugauacg accacaguug cggcucaacu ugcugcagca   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>660<br>720<br>780<br>840<br>900                |
| auuaauacaa acagccuguc gcuguugacc cagaauaacc ugaacaaauc ccaguccgca cugggcacug cuaucgagcg uuugucuucc ggucugcgua ucaacagcgc gaaagacgau gcggcaggac aggcgauugc uaaccguuuu accgcgaacaa ucaacagcgc gaaagacgau uccccguaacg cuaacgacgg uaucuccauu gcgcagacca cugaaggcgc gcugaacgaa aucaacaaca accugcagcg ugugcgugaa cuggcgguuc agucugcgaa ugguacuaac cguguauccg gccagacuca guucaacggc gaaaucaccc agcgccugaa cgaaaucgac cguguauccg gccagacuca guucaacggc gugaaagucc uggcgcagga caacacccug accauccagg uuggugccaa cgacggugaa acuaucgaua uugauuuaaa agaaaucagc ucuaaaacac uggggacuuga uaagcuuaau guccaagaug ccuacacccc gaaagaaacu gcuguaaccg uugauaaaac uaccuauaaa aaugguacag ggguuacugg ggcugauauc aaauuuaaaa auggucaaua cuauuuagau guuaaaggcg gugcuucugc ugguguuuau aaagccacuu augaugaaac uacaaagaaa guuaauauug auacgacuga uaaaacuccg uuggcaacuu cgaaagaaacu gcugaagcaacuu augaugaaac agcuauucgg ggaacggcca cuauaacca caaccaaauu gcugaaaguaa caaaagaggg uguugauacg accacaguug cggcucaacu ugcugcagca ggggguuacug gcgcaacuu ggcgcaacaa agcuauucgg accacaguug cggcucaacu ugcugcagca ggggguuacug gggguuacug gggguuacug gggguuacug aacacacaaauu gcugaaguaa caaaagaggg uguugauacg accacaguug cggcucaacu ugcugcagca ggggguuacug gggguuacug gggguuacug ggggaaagaca aacacaaauu gcugaaguaa caaaagaaggg agacaauacu agccuuguaa aacuaccgu ugcugcagca ggggguuacug gggguuacug gggguuacug ggggguuacug aacacacuu ugcugcagca ggggguuacug gggguuacug ggggguuacug aacacacuu ugcugcagca gggggggggg | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>660<br>720<br>780<br>840<br>900<br>960<br>1020 |

| ggcg                                 | guugo                                     | cuc a                                    | aaacı               | ıggaç               | ge u       | gugaa      | aauuu      | ı ggı      | ıggcç         | gcaa       | aug        | guaaa      | auc 1      | ugaaç      | guuguu     | 1200 |
|--------------------------------------|---|--|---------------------|---------------------|------------|------------|------------|------------|---------------|------------|------------|------------|------------|------------|------------|------|
| acuç                                 | gcuad                                     | ccg a                                    | auggı               | ıaaga               | ac ui      | ıacuı      | ıagca      | a ago      | cgaco         | cuug       | acaa       | acau       | ıaa (      | cuuca      | agaaca     | 1260 |
| ggcg                                 | gguga                                     | agc 1                                    | uaaa                | agago               | gu ua      | aauao      | cagau      | ı aaç      | gacuç         | gaaa       | acco       | cacuç      | gca 🤅      | gaaaa      | auugau     | 1320 |
| gcu                                  | gccui                                     | ıgg (                                    | cacaç               | gguug               | ga ua      | acacı      | ıucgu      | ı ucı      | ıgaco         | cugg       | gug        | eggui      | ıca ç      | gaaco      | eguuuc     | 1380 |
| aacı                                 | ıccgo                                     | cua 1                                    | ıcaco               | caaco               | cu gọ      | ggcaa      | auaco      | gua        | aaaua         | aacc       | ugu        | cuucı      | ıgc (      | ccgua      | agccgu     | 1440 |
| auco                                 | gaaga                                     | auu (                                    | ccga                | cuacç               | gc aa      | accga      | aaguo      | uco        | caaca         | augu       | cuc        | gegeç      | gca q      | gauud      | cugcag     | 1500 |
| cago                                 | geegg                                     | gua (                                    | ccuc                | guu                 | cu g       | gegea      | aggcg      | g aad      | ccago         | guuc       | cgca       | aaaa       | cgu (      | ccucı      | ıcuuua     | 1560 |
| cugo                                 | eguug                                     | gau a                                    | aauaç               | ggcu                | gg ag      | gccu       | eggue      | g gco      | caugo         | cuuc       | uug        | cccı       | ug 9       | ggccı      | ıccccc     | 1620 |
| cago                                 | cccu                                      | icc i                                    | iccc                | cuuc                | cu go      | cacco      | cguac      | 000        | ccgu          | gguc       | uuuç       | gaaua      | aaa 🤉      | gucuç      | gagugg     | 1680 |
| gegg                                 | gcaaa                                     | aaa a                                    | aaaa                | aaaa                | aa aa      | aaaa       | aaaa       | a aaa      | aaaa          | aaaa       | aaaa       | aaaa       | aaa a      | aaaaa      | aaaaaa     | 1740 |
| aaaa                                 | aaaa                                      | aaa a                                    | aaaa                | aaaa                | aa aa      | aaaaa      | aaaa       | a aaa      | aaaa          | aaaa       | aaaa       | aaucı      | ıag        |            |            | 1790 |
| <211<br><212<br><213<br><220<br><223 | L> LE<br>2> TY<br>3> OF<br>0> FE<br>3> OT | ENGTI<br>YPE :<br>RGAN:<br>EATUI<br>THER | ISM:<br>RE:<br>INFO | D6<br>Art:<br>DRMA: |            |            | -          |            | ?oly <u>r</u> | oept:      | ide        |            |            |            |            |      |
|                                      |   | _  | VCE:                |                     |            |            |            |            |               |            |            |            |            |            |            |      |
| Met<br>1                             | Ala                                       | Gln                                      | Val                 | Ile<br>5            | Asn        | Thr        | Asn        | Ser        | Leu<br>10     | Ser        | Leu        | Leu        | Thr        | Gln<br>15  | Asn        |      |
| Asn                                  | Leu                                       | Asn                                      | Lys<br>20           | Ser                 | Gln        | Ser        | Ala        | Leu<br>25  | Gly           | Thr        | Ala        | Ile        | Glu<br>30  | Arg        | Leu        |      |
| Ser                                  | Ser                                       | Gly<br>35                                | Leu                 | Arg                 | Ile        | Asn        | Ser<br>40  | Ala        | Lys           | Asp        | Asp        | Ala<br>45  | Ala        | Gly        | Gln        |      |
| Ala                                  | Ile<br>50                                 | Ala                                      | Asn                 | Arg                 | Phe        | Thr<br>55  | Ala        | Asn        | Ile           | Lys        | Gly<br>60  | Leu        | Thr        | Gln        | Ala        |      |
| Ser<br>65                            | Arg                                       | Asn                                      | Ala                 | Asn                 | Asp<br>70  | Gly        | Ile        | Ser        | Ile           | Ala<br>75  | Gln        | Thr        | Thr        | Glu        | Gly<br>80  |      |
| Ala                                  | Leu                                       | Asn                                      | Glu                 | Ile<br>85           | Asn        | Asn        | Asn        | Leu        | Gln<br>90     | Arg        | Val        | Arg        | Glu        | Leu<br>95  | Ala        |      |
| Val                                  | Gln                                       | Ser                                      | Ala<br>100          | Asn                 | Gly        | Thr        | Asn        | Ser<br>105 | Gln           | Ser        | Asp        | Leu        | Asp<br>110 | Ser        | Ile        |      |
| Gln                                  | Ala                                       | Glu<br>115                               | Ile                 | Thr                 | Gln        | Arg        | Leu<br>120 | Asn        | Glu           | Ile        | Asp        | Arg<br>125 | Val        | Ser        | Gly        |      |
| Gln                                  | Thr<br>130                                | Gln                                      | Phe                 | Asn                 | Gly        | Val<br>135 | Lys        | Val        | Leu           | Ala        | Gln<br>140 | Asp        | Asn        | Thr        | Leu        |      |
| Thr<br>145                           | Ile                                       | Gln                                      | Val                 | Gly                 | Ala<br>150 | Asn        | Asp        | Gly        | Glu           | Thr<br>155 | Ile        | Asp        | Ile        | Asp        | Leu<br>160 |      |
| Lys                                  | Glu                                       | Ile                                      | Ser                 | Ser<br>165          | Lys        | Thr        | Leu        | Gly        | Leu<br>170    | Asp        | Lys        | Leu        | Asn        | Val<br>175 | Gln        |      |
| Asp                                  | Ala                                       | Tyr                                      | Thr<br>180          | Pro                 | Lys        | Glu        | Thr        | Ala<br>185 | Val           | Thr        | Val        | Asp        | Lys<br>190 | Thr        | Thr        |      |
| Tyr                                  | Lys                                       | Asn<br>195                               | Gly                 | Thr                 | Asp        | Pro        | Ile<br>200 | Thr        | Ala           | Gln        | Ser        | Asn<br>205 | Thr        | Asp        | Ile        |      |
| Gln                                  | Thr<br>210                                | Ala                                      | Ile                 | Gly                 | Gly        | Gly<br>215 | Ala        | Thr        | Gly           | Val        | Thr<br>220 | Gly        | Ala        | Asp        | Ile        |      |
| Lys<br>225                           | Phe                                       | Lys                                      | Asp                 | Gly                 | Gln<br>230 | Tyr        | Tyr        | Leu        | Asp           | Val<br>235 | ГЛа        | Gly        | Gly        | Ala        | Ser<br>240 |      |
| Ala                                  | Gly                                       | Val                                      | Tyr                 | Lys<br>245          | Ala        | Thr        | Tyr        | Asp        | Glu<br>250    | Thr        | Thr        | Lys        | Lys        | Val<br>255 | Asn        |      |
|                                      |   |  |                     |                     |            |            |            |            |               |            |            |            |            |            |            |      |

| Ile As                              | p Thr                             | Thr<br>260                  | Asp        | Lys        | Thr        | Pro        | Leu<br>265 | Ala           | Thr        | Ala        | Glu        | Ala<br>270 | Thr        | Ala        |
|-------------------------------------|-----------------------------------|-----------------------------|------------|------------|------------|------------|------------|---------------|------------|------------|------------|------------|------------|------------|
| Ile Ar                              | g Gly<br>275                      | Thr                         | Ala        | Thr        | Ile        | Thr<br>280 | His        | Asn           | Gln        | Ile        | Ala<br>285 | Glu        | Val        | Thr        |
| Lys Gl<br>29                        |                                   | Val                         | Asp        | Thr        | Thr<br>295 | Thr        | Val        | Ala           | Ala        | Gln<br>300 | Leu        | Ala        | Ala        | Ala        |
| Gly Va<br>305                       | l Thr                             | Gly                         | Ala        | Asp<br>310 | Lys        | Asp        | Asn        | Thr           | Ser<br>315 | Leu        | Val        | Lys        | Leu        | Ser<br>320 |
| Phe Gl                              | u Asp                             | Lys                         | Asn<br>325 | Gly        | Lys        | Val        | Ile        | Asp<br>330    | Gly        | Gly        | Tyr        | Ala        | Val<br>335 | ГÀа        |
| Met Gl                              | y Asp                             | Asp<br>340                  | Phe        | Tyr        | Ala        | Ala        | Thr<br>345 | Tyr           | Asp        | Glu        | Lys        | Thr<br>350 | Gly        | Ala        |
| Ile Th                              | r Ala<br>355                      | Lys                         | Thr        | Thr        | Thr        | Tyr<br>360 | Thr        | Asp           | Gly        | Thr        | Gly<br>365 | Val        | Ala        | Gln        |
| Thr Gl                              | _                                 | Val                         | ГÀа        | Phe        | Gly<br>375 | Gly        | Ala        | Asn           | Gly        | 380        | Ser        | Glu        | Val        | Val        |
| Thr Al<br>385                       | a Thr                             | Asp                         | Gly        | 390<br>TÀa | Thr        | Tyr        | Leu        | Ala           | Ser<br>395 | Asp        | Leu        | Asp        | Lys        | His<br>400 |
| Asn Ph                              | e Arg                             | Thr                         | Gly<br>405 | Gly        | Glu        | Leu        | Lys        | Glu<br>410    | Val        | Asn        | Thr        | Asp        | Lys<br>415 | Thr        |
| Glu As                              | n Pro                             | Leu<br>420                  | Gln        | ГÀа        | Ile        | Asp        | Ala<br>425 | Ala           | Leu        | Ala        | Gln        | Val<br>430 | Asp        | Thr        |
| Leu Ar                              | g Ser<br>435                      | Asp                         | Leu        | Gly        | Ala        | Val<br>440 | Gln        | Asn           | Arg        | Phe        | Asn<br>445 | Ser        | Ala        | Ile        |
| Thr As                              |                                   | Gly                         | Asn        | Thr        | Val<br>455 | Asn        | Asn        | Leu           | Ser        | Ser<br>460 | Ala        | Arg        | Ser        | Arg        |
| Ile Gl<br>465                       | u Asp                             | Ser                         | Asp        | Tyr<br>470 | Ala        | Thr        | Glu        | Val           | Ser<br>475 | Asn        | Met        | Ser        | Arg        | Ala<br>480 |
| Gln Il                              | e Leu                             | Gln                         | Gln<br>485 | Ala        | Gly        | Thr        | Ser        | Val<br>490    | Leu        | Ala        | Gln        | Ala        | Asn<br>495 | Gln        |
| Val Pr                              | o Gln                             | Asn<br>500                  | Val        | Leu        | Ser        | Leu        | Leu<br>505 | Arg           |            |            |            |            |            |            |
| <210><211><211><212><213><220><223> | LENGT<br>TYPE :<br>ORGAN<br>FEATU | H: 6:<br>PRT<br>ISM:<br>RE: | 98<br>Art: |            |            | _          |            | Poly <u>r</u> | pept:      | ide        |            |            |            |            |
| <400>                               | SEQUE                             | NCE :                       | 55         |            |            |            |            |               |            |            |            |            |            |            |
| Met Al<br>1                         | a Gln                             | Val                         | Ile<br>5   | Asn        | Thr        | Asn        | Ser        | Leu<br>10     | Ser        | Leu        | Leu        | Thr        | Gln<br>15  | Asn        |
| Asn Le                              | u Asn                             | Lys<br>20                   | Ser        | Gln        | Ser        | Ala        | Leu<br>25  | Gly           | Thr        | Ala        | Ile        | Glu<br>30  | Arg        | Leu        |
| Ser Se                              | r Gly<br>35                       | Leu                         | Arg        | Ile        | Asn        | Ser<br>40  | Ala        | Lys           | Asp        | Asp        | Ala<br>45  | Ala        | Gly        | Gln        |
| Ala Il<br>50                        | e Ala                             | Asn                         | Arg        | Phe        | Thr<br>55  | Ala        | Asn        | Ile           | Lys        | Gly<br>60  | Leu        | Thr        | Gln        | Ala        |
| Ser Ar<br>65                        | g Asn                             | Ala                         | Asn        | Asp<br>70  | Gly        | Ile        | Ser        | Ile           | Ala<br>75  | Gln        | Thr        | Thr        | Glu        | Gly<br>80  |
| Ala Le                              | u Asn                             | Glu                         | Ile<br>85  | Asn        | Asn        | Asn        | Leu        | Gln<br>90     | Arg        | Val        | Arg        | Glu        | Leu<br>95  | Ala        |
| Val Gl                              | n Ser                             | Ala<br>100                  | Asn        | Ser        | Thr        | Asn        | Ser<br>105 | Gln           | Ser        | Asp        | Leu        | Asp<br>110 | Ser        | Ile        |

| Gln        | Ala        | Glu<br>115 | Ile        | Thr        | Gln        | Arg        | Leu<br>120 | Asn        | Glu        | Ile        | Asp        | Arg<br>125 | Val        | Ser        | Gly        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Gln        | Thr        | Gln        | Phe        | Asn        | Gly        | Val<br>135 | ГЛа        | Val        | Leu        | Ala        | Gln<br>140 | Asp        | Asn        | Thr        | Leu        |
| Thr<br>145 | Ile        | Gln        | Val        | Gly        | Ala<br>150 | Asn        | Asp        | Gly        | Glu        | Thr<br>155 | Ile        | Asp        | Ile        | Asp        | Leu<br>160 |
| Lys        | Gln        | Ile        | Asn        | Ser<br>165 | Gln        | Thr        | Leu        | Gly        | Leu<br>170 | Asp        | Thr        | Leu        | Asn        | Val<br>175 | Gln        |
| Gln        | Lys        | Tyr        | Lys<br>180 | Val        | Ser        | Asp        | Thr        | Ala<br>185 | Ala        | Thr        | Val        | Thr        | Gly<br>190 | Tyr        | Ala        |
| Asp        | Thr        | Thr<br>195 | Ile        | Ala        | Leu        | Asp        | Asn<br>200 | Ser        | Thr        | Phe        | Lys        | Ala<br>205 | Ser        | Ala        | Thr        |
| Gly        | Leu<br>210 | Gly        | Gly        | Thr        | Asp        | Gln<br>215 | Lys        | Ile        | Asp        | Gly        | Asp<br>220 | Leu        | Lys        | Phe        | Asp        |
| Asp<br>225 | Thr        | Thr        | Gly        | ГÀа        | Tyr<br>230 | Tyr        | Ala        | Lys        | Val        | Thr<br>235 | Val        | Thr        | Gly        | Gly        | Thr<br>240 |
| Gly        | Lys        | Asp        | Gly        | Tyr<br>245 | Tyr        | Glu        | Val        | Ser        | Val<br>250 | Asp        | Lys        | Thr        | Asn        | Gly<br>255 | Glu        |
| Val        | Thr        | Leu        | Ala<br>260 | Gly        | Gly        | Ala        | Thr        | Ser<br>265 | Pro        | Leu        | Thr        | Gly        | Gly<br>270 | Leu        | Pro        |
| Ala        | Thr        | Ala<br>275 | Thr        | Glu        | Asp        | Val        | Lys<br>280 | Asn        | Val        | Gln        | Val        | Ala<br>285 | Asn        | Ala        | Asp        |
| Leu        | Thr<br>290 | Glu        | Ala        | ГÀа        | Ala        | Ala<br>295 | Leu        | Thr        | Ala        | Ala        | Gly<br>300 | Val        | Thr        | Gly        | Thr        |
| Ala<br>305 | Ser        | Val        | Val        | rys        | Met<br>310 | Ser        | Tyr        | Thr        | Asp        | Asn<br>315 | Asn        | Gly        | Lys        | Thr        | Ile<br>320 |
| Asp        | Gly        | Gly        | Leu        | Ala<br>325 | Val        | Lys        | Val        | Gly        | Asp<br>330 | Asp        | Tyr        | Tyr        | Ser        | Ala<br>335 | Thr        |
| Gln        | Asn        | Lys        | Asp<br>340 | Gly        | Ser        | Ile        | Ser        | Ile<br>345 | Asn        | Thr        | Thr        | Lys        | Tyr<br>350 | Thr        | Ala        |
| Asp        | Asp        | Gly<br>355 | Thr        | Ser        | ГÀз        | Thr        | Ala<br>360 | Leu        | Asn        | Lys        | Leu        | Gly<br>365 | Gly        | Ala        | Asp        |
| Gly        | Lys<br>370 | Thr        | Glu        | Val        | Val        | Ser<br>375 | Ile        | Gly        | Gly        | Lys        | Thr<br>380 | Tyr        | Ala        | Ala        | Ser        |
| 385<br>Tàs | Ala        | Glu        | Gly        | His        | Asn<br>390 | Phe        | Lys        | Ala        | Gln        | Pro<br>395 | Asp        | Leu        | Ala        | Glu        | Ala<br>400 |
| Ala        | Ala        | Thr        | Thr        | Thr<br>405 | Glu        | Asn        | Pro        | Leu        | Gln<br>410 | Lys        | Ile        | Asp        | Ala        | Ala<br>415 | Leu        |
| Ala        | Gln        | Val        | Asp<br>420 | Thr        | Leu        | Arg        | Ser        | Asp<br>425 | Leu        | Gly        | Ala        | Val        | Gln<br>430 | Asn        | Arg        |
| Phe        | Asn        | Ser<br>435 | Ala        | Ile        | Thr        | Asn        | Leu<br>440 | Gly        | Asn        | Thr        | Val        | Asn<br>445 | Asn        | Leu        | Thr        |
| Ser        | Ala<br>450 | Arg        | Ser        | Arg        | Ile        | Glu<br>455 | Asp        | Ser        | Asp        | Tyr        | Ala<br>460 | Thr        | Glu        | Val        | Ser        |
| Asn<br>465 | Met        | Ser        | Arg        | Ala        | Gln<br>470 | Ile        | Leu        | Gln        | Gln        | Ala<br>475 | Gly        | Thr        | Ser        | Val        | Leu<br>480 |
| Ala        | Gln        | Ala        | Asn        | Gln<br>485 | Val        | Pro        | Gln        | Asn        | Val<br>490 | Leu        | Ser        | Leu        | Leu        | Arg<br>495 | Gly        |
| Gly        | Gly        | Gly        | Ser<br>500 | Gly        | Gly        | Gly        | Gly        | Ser<br>505 | Met        | Met        | Ala        | Pro        | Asp<br>510 | Pro        | Asn        |
| Ala        | Asn        | Pro<br>515 | Asn        | Ala        | Asn        | Pro        | Asn<br>520 | Ala        | Asn        | Pro        | Asn        | Ala<br>525 | Asn        | Pro        | Asn        |

| Ala Asn<br>530   |                                 |  |  |   |  |   |  |                                       |  |  |   |  |                                     |   |
|--|---------------------------------|--|--|---|--|---|--|---------------------------------------|--|--|---|--|-------------------------------------|---|
|  | Pro                             | Asn  | Ala  | Asn                                     | Pro<br>535   | Asn   | Ala  | Asn                                   | Pro  | Asn<br>540   | Ala                                     | Asn  | Pro                                 | Asn   |
| Ala Asn<br>545   | Pro                             | Asn  | Ala  | Asn<br>550                              | Pro  | Asn   | Ala  | Asn                                   | Pro<br>555                                   | Asn  | Ala                                     | Asn  | Pro                                 | Asn<br>560                                  |
| Ala Asn  | Pro                             | Asn  | Ala<br>565   | Asn                                     | Pro  | Asn   | Ala  | Asn<br>570                            | Pro  | Asn  | Ala                                     | Asn  | Pro<br>575                          | Asn   |
| Ala Asn  | Pro                             | Asn<br>580                                     | Ala  | Asn                                     | Pro  | Asn   | Ala<br>585                                 | Asn                                   | Pro  | Asn  | Lys                                     | Asn<br>590                                   | Asn                                 | Gln   |
| Gly Asn  | Gly<br>595                      | Gln  | Gly  | His                                     | Asn  | Met<br>600                                      | Pro  | Asn                                   | Asp  | Pro  | Asn<br>605                              | Arg  | Asn                                 | Val   |
| Asp Glu<br>610   | Asn                             | Ala  | Asn  | Ala                                     | Asn<br>615   | Asn   | Ala  | Val                                   | Lys  | Asn<br>620   | Asn                                     | Asn  | Asn                                 | Glu   |
| Glu Pro<br>625   | Ser                             | Asp  | ГЛа  | His<br>630                              | Ile  | Glu   | Gln  | Tyr                                   | Leu<br>635                                   | ГЛа  | ГЛа                                     | Ile  | Lys                                 | Asn<br>640                                  |
| Ser Ile  | Ser                             | Thr  | Glu<br>645   | Trp                                     | Ser  | Pro   | Cys  | Ser<br>650                            | Val  | Thr  | Cys                                     | Gly  | Asn<br>655                          | Gly   |
| Ile Gln  | Val                             | Arg<br>660                                     | Ile  | Lys                                     | Pro  | Gly   | Ser<br>665                                 | Ala                                   | Asn  | Lys  | Pro                                     | Lys<br>670                                   | Asp                                 | Glu   |
| Leu Asp  | Tyr<br>675                      | Glu  | Asn  | Asp                                     | Ile  | Glu<br>680                                      | Lys  | ГЛа                                   | Ile  | CÀa  | Lys<br>685                              | Met  | Glu                                 | Lys   |
| Cys Ser<br>690   | Ser                             | Val  | Phe  | Asn                                     | Val<br>695   | Val   | Asn  | Ser                                   |  |  |   |  |                                     |   |
| <220> F<br><223> O   |                                 |  | DRMA:  | CION                                    | : Svi  | nthet   | ia I                                       | 20111                                 |  |  |   |  |                                     |   |
| <223> 0  | IHER                            | TMFC   | JKMA.  | I TON                                   |  |   |  |                                       |  |  |   |  |                                     |   |
| <400> S  | EQUE                            | ICE :  |  |   | 2 -  |   | .10 1                                      | POTAF                                 | рерс.  | ide  |   |  |                                     |   |
| <400> S<br>Met Met<br>1  | -                               |  | 56   |   |  |   |  |                                       | _  |  | Asn                                     | Pro  | Asn<br>15                           | Ala   |
| Met Met  | Ala                             | Pro  | 56<br>Asp<br>5                                       | Pro                                     | Asn  | Ala   | Asn  | Pro<br>10                             | Asn  | Ala  |   |  | 15                                  |   |
| Met Met<br>1   | Ala<br>Asn                      | Pro<br>Ala<br>20                               | 56<br>Asp<br>5<br>Asn                                | Pro<br>Pro                              | Asn<br>Asn   | Ala<br>Ala                                      | Asn<br>Asn<br>25                           | Pro<br>10                             | Asn<br>Asn                                   | Ala<br>Ala   | Asn                                     | Pro<br>30                                    | 15<br>Asn                           | Ala   |
| Met Met<br>1<br>Asn Pro  | Ala<br>Asn<br>Asn<br>35         | Pro<br>Ala<br>20<br>Ala                        | 56<br>Asp<br>5<br>Asn<br>Asn                         | Pro<br>Pro                              | Asn<br>Asn<br>Asn                                  | Ala<br>Ala<br>Ala<br>40                         | Asn<br>Asn<br>25<br>Asn                    | Pro<br>10<br>Pro                      | Asn<br>Asn<br>Asn                            | Ala<br>Ala<br>Ala                                  | Asn<br>Asn<br>45                        | Pro<br>30<br>Pro                             | 15<br>Asn<br>Asn                    | Ala<br>Ala                                  |
| Met Met 1 Asn Pro Asn Pro  | Ala<br>Asn<br>Asn<br>35<br>Asn  | Pro Ala 20 Ala Ala                             | Asp<br>5<br>Asn<br>Asn<br>Asn                        | Pro<br>Pro<br>Pro                       | Asn<br>Asn<br>Asn<br>Asn<br>55                     | Ala<br>Ala<br>Ala<br>40<br>Ala                  | Asn<br>Asn<br>25<br>Asn<br>Asn             | Pro<br>10<br>Pro<br>Pro               | Asn<br>Asn<br>Asn                            | Ala<br>Ala<br>Ala<br>Ala                           | Asn<br>Asn<br>45<br>Asn                 | Pro<br>30<br>Pro                             | 15<br>Asn<br>Asn<br>Asn             | Ala<br>Ala<br>Ala                           |
| Met Met 1 Asn Pro Asn Pro Asn Pro So Asn Pro   | Ala<br>Asn<br>Asn<br>35<br>Asn  | Pro Ala 20 Ala Ala Ala                         | Asp<br>5<br>Asn<br>Asn<br>Asn                        | Pro Pro Pro Pro 70                      | Asn<br>Asn<br>Asn<br>55<br>Asn                     | Ala<br>Ala<br>Ala<br>40<br>Ala                  | Asn<br>25<br>Asn<br>Asn                    | Pro<br>10<br>Pro<br>Pro               | Asn<br>Asn<br>Asn<br>Asn<br>75               | Ala<br>Ala<br>Ala<br>Ala<br>60                     | Asn<br>45<br>Asn<br>Asn                 | Pro<br>30<br>Pro<br>Pro                      | Asn<br>Asn<br>Asn<br>Asn            | Ala<br>Ala<br>Ala<br>Ala                    |
| Met 1 Asn Pro Asn Pro 50 Asn Pro 65  | Ala Asn Asn S Asn Asn Asn       | Pro Ala 20 Ala Ala Ala Lys                     | Asn Asn Asn Asn Asn                                  | Pro Pro Pro Asn                         | Asn<br>Asn<br>Asn<br>Asn<br>Gln                    | Ala<br>Ala<br>Ala<br>Ala<br>Ala                 | Asn<br>25<br>Asn<br>Asn<br>Asn             | Pro 10 Pro Pro Gly 90                 | Asn<br>Asn<br>Asn<br>Asn<br>Gln              | Ala<br>Ala<br>Ala<br>60<br>Ala                     | Asn<br>45<br>Asn<br>Asn                 | Pro<br>30<br>Pro<br>Pro<br>Pro               | Asn Asn Asn Met                     | Ala<br>Ala<br>Ala<br>Ala<br>80<br>Pro       |
| Met Met 1 Asn Pro Asn Pro 50 Asn Pro 65 Asn Pro                                      | Ala Asn Asn Asn Asn Pro         | Pro Ala 20 Ala Ala Lys Asn 100                 | Asp<br>Asn<br>Asn<br>Asn<br>Asn<br>Asn               | Pro Pro Pro Pro Asn                     | Asn<br>Asn<br>Asn<br>Asn<br>Gln<br>Val             | Ala Ala Ala Ala Ala Ala Ala Ala                 | Asn 25 Asn Asn Asn Glu 105                 | Pro 10 Pro Pro Pro Asn                | Asn<br>Asn<br>Asn<br>Asn<br>Gln              | Ala Ala Ala Ala Gly Asn                            | Asn<br>Asn<br>Asn<br>Asn<br>Ala         | Pro<br>30<br>Pro<br>Pro<br>Asn<br>Asn        | Asn Asn Asn Met 95 Asn              | Ala Ala Ala 80 Pro                          |
| Met 1 Asn Pro Asn Pro 50 Asn Pro 65 Asn Pro Asn Asn                                  | Ala Asn Asn 35 Asn Asn Asn Asn  | Pro Ala 20 Ala Ala Ala Ala Lys Asn 100 Asn     | Asp<br>5<br>Asn<br>Asn<br>Asn<br>Asn<br>Asn          | Pro Pro Pro Pro Asn Asn                 | Asn<br>Asn<br>Asn<br>S5<br>Asn<br>Gln<br>Val       | Ala Ala Ala Ala Ala Ala Ala Gly Asp Glu 120     | Asn 25 Asn Asn Asn Glu 105 Pro             | Pro 10 Pro Pro Pro Asn Ser            | Asn<br>Asn<br>Asn<br>Asn<br>75<br>Gln<br>Ala | Ala Ala Ala Ala 60 Ala Gly Asn                     | Asn Asn Asn Asn Asn His Ala             | Pro Pro Pro Asn Asn 110                      | Asn Asn Asn Asn Glu                 | Ala Ala Ala Ala Pro Ala Gln                 |
| Met 1 Asn Pro Asn Pro 50 Asn Pro 65 Asn Pro Asn Asp Val Lys Tyr Leu                  | Ala Asn Asn Asn Asn Asn Asn Lys | Pro Ala 20 Ala Ala Ala Lys Asn 100 Asn         | Asp<br>Asn<br>Asn<br>Asn<br>Asn<br>Asn<br>Asn<br>Ile | Pro Pro Pro Pro Asn Asn Lys             | Asn Asn Asn Asn Cln Cln Clu Asn Clu Asn            | Ala Ala Ala Ala Ala Ala Gly Asp Glu 120 Ser     | Asn 25 Asn Asn Asn Fro                     | Pro 10 Pro Pro Pro Asn Ser            | Asn Asn Asn Asn Asn Thr                      | Ala Ala Ala Ala 60 Ala Gly Asn Lys Glu 140         | Asn Asn 45 Asn Asn Ala His 125          | Pro<br>30<br>Pro<br>Pro<br>Asn<br>Asn<br>110 | Asn Asn Asn Asn Asn Glu Pro         | Ala Ala Ala Ala Pro Ala Gln Cys             |
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| Met 1 Asn Pro Asn Pro 50 Asn Pro 65 Asn Pro Asn Asp Val Lys Tyr Leu 130 Ser Val      | Ala Asn Asn Asn Asn Asn Lys Lys | Pro Ala 20 Ala Ala Ala Ala Lys Asn 100 Asn Lys | Asn Asn Asn Asn Asn Ile Gly Lys 165                  | Pro Pro Pro Pro Asn Asn Asn Asn Asn Asn | Asn Asn Asn Asn S5 Asn Gln Val Glu Asn 135 Gly Glu | Ala Ala Ala Ala Ala Ala Gly Asp Glu 120 Ser Ile | Asn 25 Asn Asn Asn Glu 105 Pro Ile Gln Asp | Pro 10 Pro Pro Pro Asn Ser Val        | Asn Asn Asn Asn Asn Thr Arg 155 Glu          | Ala Ala Ala Ala 60 Ala Gly Asn Lys Glu 140 Ile Asn | Asn Asn Asn Asn Asn His Ala His Lys Asp | Pro 30 Pro Pro Asn Asn 110 Ile Fro Ile       | Asn Asn Asn Asn Glu Pro Gly Glu 175 | Ala Ala Ala Ala Pro Ala Gln Cys Ser 160 Lys |

| Ser        | Arg        | Pro<br>195 | Val        | Thr        | Met        | Ala        | Gln<br>200 | Val        | Ile        | Asn        | Thr        | Asn<br>205 | Ser        | Leu        | Ser        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Leu        | Leu<br>210 | Thr        | Gln        | Asn        | Asn        | Leu<br>215 | Asn        | Lys        | Ser        | Gln        | Ser<br>220 | Ala        | Leu        | Gly        | Thr        |
| Ala<br>225 | Ile        | Glu        | Arg        | Leu        | Ser<br>230 | Ser        | Gly        | Leu        | Arg        | Ile<br>235 | Asn        | Ser        | Ala        | Lys        | Asp<br>240 |
| Asp        | Ala        | Ala        | Gly        | Gln<br>245 | Ala        | Ile        | Ala        | Asn        | Arg<br>250 | Phe        | Thr        | Ala        | Asn        | Ile<br>255 | Lys        |
| Gly        | Leu        | Thr        | Gln<br>260 | Ala        | Ser        | Arg        | Asn        | Ala<br>265 | Asn        | Asp        | Gly        | Ile        | Ser<br>270 | Ile        | Ala        |
| Gln        | Thr        | Thr<br>275 | Glu        | Gly        | Ala        | Leu        | Asn<br>280 | Glu        | Ile        | Asn        | Asn        | Asn<br>285 | Leu        | Gln        | Arg        |
| Val        | Arg<br>290 | Glu        | Leu        | Ala        | Val        | Gln<br>295 | Ser        | Ala        | Asn        | Ser        | Thr<br>300 | Asn        | Ser        | Gln        | Ser        |
| Asp<br>305 | Leu        | Asp        | Ser        | Ile        | Gln<br>310 | Ala        | Glu        | Ile        | Thr        | Gln<br>315 | Arg        | Leu        | Asn        | Glu        | Ile<br>320 |
| Asp        | Arg        | Val        | Ser        | Gly<br>325 | Gln        | Thr        | Gln        | Phe        | Asn<br>330 | Gly        | Val        | ГÀа        | Val        | Leu<br>335 | Ala        |
| Gln        | Asp        | Asn        | Thr<br>340 | Leu        | Thr        | Ile        | Gln        | Val<br>345 | Gly        | Ala        | Asn        | Asp        | Gly<br>350 | Glu        | Thr        |
| Ile        | Asp        | Ile<br>355 | Asp        | Leu        | ГÀа        | Gln        | Ile<br>360 | Asn        | Ser        | Gln        | Thr        | Leu<br>365 | Gly        | Leu        | Asp        |
| Thr        | Leu<br>370 | Asn        | Val        | Gln        | Gln        | Lys<br>375 | Tyr        | Lys        | Val        | Ser        | Asp<br>380 | Thr        | Ala        | Ala        | Thr        |
| Val<br>385 | Thr        | Gly        | Tyr        | Ala        | Asp<br>390 | Thr        | Thr        | Ile        | Ala        | Leu<br>395 | Asp        | Asn        | Ser        | Thr        | Phe<br>400 |
| ГÀв        | Ala        | Ser        | Ala        | Thr<br>405 | Gly        | Leu        | Gly        | Gly        | Thr<br>410 | Asp        | Gln        | Lys        | Ile        | Asp<br>415 | Gly        |
| Asp        | Leu        | Lys        | Phe<br>420 | Asp        | Asp        | Thr        | Thr        | Gly<br>425 | Lys        | Tyr        | Tyr        | Ala        | Lys<br>430 | Val        | Thr        |
| Val        | Thr        | Gly<br>435 | Gly        | Thr        | Gly        | Lys        | Asp<br>440 | Gly        | Tyr        | Tyr        | Glu        | Val<br>445 | Ser        | Val        | Asp        |
| rya        | Thr<br>450 | Asn        | Gly        | Glu        | Val        | Thr<br>455 | Leu        | Ala        | Gly        | Gly        | Ala<br>460 | Thr        | Ser        | Pro        | Leu        |
| Thr<br>465 | Gly        | Gly        | Leu        | Pro        | Ala<br>470 | Thr        | Ala        | Thr        | Glu        | Asp<br>475 | Val        | Lys        | Asn        | Val        | Gln<br>480 |
| Val        | Ala        | Asn        | Ala        | Asp<br>485 | Leu        | Thr        | Glu        | Ala        | Lys<br>490 | Ala        | Ala        | Leu        | Thr        | Ala<br>495 | Ala        |
| Gly        | Val        | Thr        | Gly<br>500 | Thr        | Ala        | Ser        | Val        | Val<br>505 | Lys        | Met        | Ser        | Tyr        | Thr<br>510 | Asp        | Asn        |
| Asn        | Gly        | Lys<br>515 | Thr        | Ile        | Asp        | Gly        | Gly<br>520 | Leu        | Ala        | Val        | Lys        | Val<br>525 | Gly        | Asp        | Asp        |
| Tyr        | Tyr<br>530 | Ser        | Ala        | Thr        | Gln        | Asn<br>535 | Lys        | Asp        | Gly        | Ser        | Ile<br>540 | Ser        | Ile        | Asn        | Thr        |
| Thr<br>545 | Lys        | Tyr        | Thr        | Ala        | Asp<br>550 | Asp        | Gly        | Thr        | Ser        | Lys<br>555 | Thr        | Ala        | Leu        | Asn        | Lys<br>560 |
| Leu        | Gly        | Gly        | Ala        | Asp<br>565 | Gly        | Lys        | Thr        | Glu        | Val<br>570 | Val        | Ser        | Ile        | Gly        | Gly<br>575 | ГЛа        |
| Thr        | Tyr        | Ala        | Ala<br>580 | Ser        | Lys        | Ala        | Glu        | Gly<br>585 | His        | Asn        | Phe        | Lys        | Ala<br>590 | Gln        | Pro        |
| Asp        | Leu        | Ala<br>595 | Glu        | Ala        | Ala        | Ala        | Thr        | Thr        | Thr        | Glu        | Asn        | Pro<br>605 | Leu        | Gln        | Lys        |
| Ile        | Asp        | Ala        | Ala        | Leu        | Ala        | Gln        | Val        | Asp        | Thr        | Leu        | Arg        | Ser        | Asp        | Leu        | Gly        |

120

180 240

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360

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1500

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559 560

-continued 610 615 Ala Val Gln Asn Arg Phe Asn Ser Ala Ile Thr Asn Leu Gly Asn Thr 625 630 635 Val Asn Asn Leu Thr Ser Ala Arg Ser Arg Ile Glu Asp Ser Asp Tyr 650 Ala Thr Glu Val Ser Asn Met Ser Arg Ala Gln Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro Gln Asn Val Leu Ser Leu Leu Arg <210> SEQ ID NO 57 <211> LENGTH: 1620 <212> TYPE: RNA <213 > ORGANISM: Unknown <220> FEATURE: <223 > OTHER INFORMATION: Human metapneumovirus <400> SEQUENCE: 57 augagcugga agguggugau uaucuucagc cugcugauua caccucaaca cggccugaag qaqaqcuacc uqqaaqaqaq cuqcuccacc aucaccqaqq qcuaccuqaq cquqcuqcqq accggcuggu acaccaacgu guucacccug qaggugggcg acguggagaa ccugaccugc agegacggee cuagecugau caagacegag cuggaccuga ccaagagege ucugagagag cugaagaceg ugucegeega ceageuggee agagaggaac agauegagaa cecueggeag agcagauucg ugcugggege caucgeucug ggaguegeeg cugeegeuge agugacageu ggaguggcca uugcuaagac caucagacug gaaagcgagg ugacagccau caacaaugcc cugaagaaga ccaacgaggc cgugagcacc cugggcaaug gagugagagu gcuggccaca gccgugcggg agcugaagga cuucgugagc aagaaccuga ccagagccau caacaagaac aagugcgaca ucgaugaccu gaagauggcc gugagcuucu cccaguucaa cagacgguuc cugaacgugg ugagacaguu cuccgacaac gcuggaauca caccugccau uagccuggac cugaugaccg acgccgagcu ggcuagagcc gugcccaaca ugcccaccag cgcuggccag aucaagcuga ugcuggagaa cagagccaug gugcggagaa agggcuucgg cauccugauu gggguguaug gaagcuccgu gaucuacaug gugcagcugc ccaucuucgg cgugaucgac acacccugcu ggaucgugaa ggccgcuccu agcugcuccg agaagaaagg aaacuaugcc ugucugcuga gagaggacca gggcugguac ugccagaacg ccggaagcac aguguacuau cccaacgaga aggacugcga gaccagaggc gaccacgugu ucugcgacac cgcugccgga 1020 1080 aucaacquqq ccqaqcaqaq caaqqaquqc aacaucaaca ucaqcacaac caacuacccc ugcaagguga gcaccggacg gcaccccauc agcauggugg cucugagccc ucugggcgcu 1140 1200 cugguggccu gcuauaaggg cguguccugu agcaucggca gcaaucgggu gggcaucauc aagcagcuga acaagggaug cuccuacauc accaaccagg acgccgacac cgugaccauc 1260

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cuguccageg cugagaaggg caacacegge uucaucauug ugaucauucu gaucgeegug

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| acagguuggu   | acaccaaugu | cuuuacauua  | gaaguuggug  | auguugaaaa | ucuuacaugu | 180  |
|--|------------|-------------|-------------|------------|------------|------|
| acugauggac   | cuagcuuaau | caaaacagaa  | cuugaccuaa  | ccaaaagugc | uuuaagagaa | 240  |
| cucaaaacag   | uuucugcuga | ucaguuagcg  | agagaagaac  | aaauugaaaa | ucccagacaa | 300  |
| ucaagguuug   | uccuaggugc | aauagcucuu  | ggaguugcca  | cagcagcagc | agucacagca | 360  |
| ggcauugcaa   | uagccaaaac | uauaaggcuu  | gagagugaag  | ugaaugcaau | caaaggugcu | 420  |
| cucaaaacaa   | ccaaugaggc | aguaucaaca  | cuaggaaaug  | gagugcgggu | ccuagccacu | 480  |
| gcaguaagag   | agcugaaaga | auuugugagc  | aaaaaccuga  | cuagugcgau | caacaagaac | 540  |
| aagugugaca   | uugcugauuu | gaagauggcu  | gucagcuuca  | gucaguucaa | cagaagauuc | 600  |
| cuaaauguug   | ugcggcaguu | uucagacaau  | gcagggauaa  | caccagcaau | aucauuggac | 660  |
| cugaugaaug   | augcugagcu | ggccagagcu  | guaucauaca  | ugccaacauc | ugcaggacag | 720  |
| auaaaacuaa   | uguuagagaa | ccgugcaaug  | gugaggagaa  | aaggauuugg | aaucuugaua | 780  |
| ggggucuacg   | gaagcucugu | gauuuacaug  | guccagcugc  | cgaucuuugg | ugucauaaau | 840  |
| acaccuuguu   | ggauaaucaa | ggcagcuccc  | ucuuguucag  | aaaaagaugg | aaauuaugcu | 900  |
| ugccuccuaa   | gagaggauca | agggugguau  | uguaaaaaug  | caggauccac | uguuuacuac | 960  |
| ccaaaugaaa   | aagacugcga | aacaagaggu  | gaucauguuu  | uuugugacac | agcagcaggg | 1020 |
| aucaauguug   | cugagcaauc | aagagaaugc  | aacaucaaca  | uaucuaccac | caacuaccca | 1080 |
| ugcaaaguca   | gcacaggaag | acacccuauc  | agcaugguug  | cacuaucacc | ucucggugcu | 1140 |
| uugguagcuu   | gcuacaaagg | gguuagcugc  | ucgacuggca  | guaaucaggu | uggaauaauc | 1200 |
| aaacaacuac   | cuaaaggcug | cucauacaua  | acuaaccagg  | acgcagacac | uguaacaauu | 1260 |
| gacaacacug   | uguaucaacu | aagcaaaguu  | gagggugaac  | agcauguaau | aaaagggaga | 1320 |
| ccaguuucaa   | gcaguuuuga | uccaaucagg  | uuuccugagg  | aucaguucaa | uguugcgcuu | 1380 |
| gaucaagucu   | uugaaagcau | ugaaaacagu  | caagcacuag  | uggaccaguc | aaacaaaauu | 1440 |
| cugaacagug   | cagaaaaagg | aaacacuggu  | uucauuauug  | uaauaauuuu | gauugcuguu | 1500 |
| cuuggguuaa   | ccaugauuuc | agugagcauc  | aucaucauaa  | ucaaaaaaac | aaggaagccc | 1560 |
| acaggggcac   | cuccggagcu | gaaugguguu  | accaacggcg  | guuucauacc | gcauaguuag | 1620 |
| <210> SEQ I<br><211> LENG<br><212> TYPE<br><213> ORGAI | ΓH: 1725   | respiratory | y syncytial | virus      |            |      |
| <400> SEQUI  | ENCE: 60   |             |             |            |            |      |
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| uguuucgcuu   | ccagucaaaa | caucacugaa  | gaauuuuauc  | aaucaacaug | cagugcaguu | 120  |
| agcaaaggcu   | aucuuagugc | ucuaagaacu  | gguugguaua  | cuaguguuau | aacuauagaa | 180  |
| uuaaguaaua   | ucaaggaaaa | uaaguguaau  | ggaacagaug  | cuaagguaaa | auugauaaaa | 240  |
| caagaauuag   | auaaauauaa | aaaugcugua  | acagaauugc  | aguugcucau | gcaaagcaca | 300  |
| ccagcagcca   | acaaucgagc | cagaagagaa  | cuaccaaggu  | uuaugaauua | uacacucaau | 360  |
| aauaccaaaa   | auaccaaugu | aacauuaagc  | aagaaaagga  | aaagaagauu | ucuuggcuuu | 420  |
| uuguuaggug   | uuggaucugc | aaucgccagu  | ggcauugcug  | uaucuaaggu | ccugcaccua | 480  |
| gaagggaag  | ugaacaaaau | caaaagugcu  | cuacuaucca  | caaacaaggc | uguagucagc | 540  |

uuaucaaaug gaguuagugu cuuaaccagc aaaguguuag accucaaaaa cuauauagau

| aaacaguugu uaccuauug  | ı gaacaagcaa   | agcugcagca  | uaucaaacau   | ugaaacugug  | 660   |  |
|---|--|---|--|---|---|--|
| auagaguucc aacaaaaga  | a caacagacua   | cuagagauua  | ccagggaauu   | uaguguuaau  | 720   |  |
| gcagguguaa cuacaccug  | ı aagcacuuau   | auguuaacua  | auagugaauu   | auuaucauua  | 780   |  |
| aucaaugaua ugccuauaa  | c aaaugaucag   | aaaaaguuaa  | uguccaacaa   | uguucaaaua  | 840   |  |
| guuagacagc aaaguuacu  | c uaucaugucc   | auaauaaagg  | aggaagucuu   | agcauaugua  | 900   |  |
| guacaauuac cacuauaug  | g uguaauagau   | acacccuguu  | ggaaacugca   | cacauccccu  | 960   |  |
| cuauguacaa ccaacacaa  | a ggaagggucc   | aacaucugcu  | uaacaagaac   | cgacagagga  | 1020  |  |
| ugguauugug acaaugcag  | g aucaguaucu   | uucuucccac  | aagcugaaac   | auguaaaguu  | 1080  |  |
| caaucgaauc ggguauuuu  | g ugacacaaug   | aacaguuuaa  | cauuaccaag   | ugaaguaaau  | 1140  |  |
| cucugcaaca uugacauau  | ı caaccccaaa   | uaugauugca  | aaauuaugac   | uucaaaaaca  | 1200  |  |
| gauguaagca gcuccguua  | ı cacaucucua   | ggagccauug  | ugucaugcua   | uggcaaaacu  | 1260  |  |
| aaauguacag cauccaaua  | a aaaucguggg   | aucauaaaga  | cauuuucuaa   | cgggugugau  | 1320  |  |
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| aagcaagaag gcaaaaguc  | ı cuauguaaaa   | ggugaaccaa  | uaauaaauuu   | cuaugaccca  | 1440  |  |
| uuaguguucc ccucugaug  | a auuugaugca   | ucaauaucuc  | aagucaauga   | gaagauuaac  | 1500  |  |
| cagagecuag cauuuauuc  | g uaaauccgau   | gaauuauuac  | auaauguaaa   | ugcugguaaa  | 1560  |  |
| uccaccacaa auaucauga  | ı aacuacuaua   | auuauaguga  | uuauaguaau   | auuguuauca  | 1620  |  |
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| <pre>&lt;211&gt; LENGTH: 1617 &lt;212&gt; TYPE: RNA &lt;213&gt; ORGANISM: Huma: &lt;400&gt; SEQUENCE: 61 augccaauuu caauacugu: gacaucacaa aacuacagc: ucacaaaacu ucgaaacaa: aacucuugug gugaccaac: ccuuuauaug auggacuaac: gaaaacacug aucccagaa: ggaguagcaa ccucagcac: agaucagaca uugaaaaaac: guucagagcu cuguaggaa: aaagaaaucg ugccaucga: auugcauuaa cacagcauu: uuacaagaaa aaggaauaa</pre>   | a aauuauuaca a uguaggugua g auaucuaauc a gaucaagcaa g auuacagaag c agaacgauuc a aauuacagca u caaggaagca a uuugauagua u ugcgagacua a cucagaauua a auuacaaggu              | accaugauca uuggucaaca cugagucuca uacaagaggu gaugugauag uuuggagggg gcaguugcuc aucagggaca gcaauuaaau gguugugaag acaaauauau auagcaucau uaugauauuu            | gucccaaagg uaccaaaaau uauuggauag ugacuaauca uaauuggaac ugguugaagc caaauaaagc caguccagga cagcaggacu uuggugauaa uauaccguac augaucuauu            | gaugaagaua agaagaucauu agaauccaau uauugcucua caagcaggca agugcaguca uuaugucaac ucaguuaggg cauaggaucg aaauaucaca                      | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660        |  |
| <pre>&lt;211&gt; LENGTH: 1617 &lt;212&gt; TYPE: RNA &lt;213&gt; ORGANISM: Huma: &lt;400&gt; SEQUENCE: 61 augccaauuu caauacugu gacaucacaa aacuacagc ucacaaaacu ucgaaacaac aacucuugug gugaccaac ccuuuauaug auggacuaac gaaaacacug aucccagaac ggaguagcaa ccucagcac agaucagaca uugaaaaaac guucagagcu cuguaggaa aaagaaaucg ugccaucga auugcauuaa cacagcauu uuacaagaaa aaggaauaa gaaauauuca caacaucaac gaaauauca caacaucaac gaaauaacacacacacacacacacacacacacacacacac</pre> | a aauuauuaca a uguaggugua g auaucuaauc a gaucaagcaa g auuacagaag c agaacgauuc a aauuacagca a uuugauagua u ugcgagacua a cucagaauua a auuacaaggu c aguugacaaa u agauguugau | accaugauca uuggucaaca cugagucuca uacaagaggu gaugugauag uuuggagggg gcaguugcuc aucagggaca gcaauuaaau gguugugaag acaaauauau auagcaucau uaugauauuu uugaaugauu | gucccaaagg uaccaaaaau uauuggauag ugacuaauca uaauuggaac ugguugaagc caaauaaagc caguccagga cagcaggacu uuggugauaa uauaccguac augaucuauu acucaauaac | gaugaagaua agaagauucu acugaucaau uauugcucua caagcaggca agugcaguca uuaugucaac ucaguuaggg cauaggaucg aaauaucaca auuuacagaa ccuccaaguc | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>720<br>780 |  |

gcauuucuag guggagcaga ugucaaagaa ugcauagaag cauucagcag uuauauaugc 1020

| ccuucugauc  | caggauuugu                       | acuaaaccau | gaaauggaga  | gcugucuauc | aggaaacaua | 1080 |
|---|----------------------------------|------------|-------------|------------|------------|------|
| ucccaauguc  | caagaaccac                       | agucacauca | gacauaguuc  | cuagguaugc | auuugucaau | 1140 |
| ggaggagugg  | uugcgaauug                       | uauaacaacu | acauguacau  | gcaaugguau | cgguaauaga | 1200 |
| aucaaccaac  | caccugauca                       | aggagucaaa | auuauaacac  | auaaagaaug | uaauacaaua | 1260 |
| gguaucaacg  | gaaugcuauu                       | caacacaaac | aaagaaggaa  | cucuugcauu | cuacacacca | 1320 |
| gacgacauaa  | cauuaaacaa                       | uucuguugca | cuugauccga  | uugacauauc | aaucgagcuc | 1380 |
| aacaaggcca  | aaucagaucu                       | ugaggaauca | aaagaaugga  | uaagaagguc | aaaucaaaag | 1440 |
| cuagauucua  | uuggaaguug                       | gcaucaaucu | agcacuacaa  | ucauaguuau | uuugauaaug | 1500 |
| augauuauau  | uguuuauaau                       | uaauauaaca | auaauuacaa  | uugcaauuaa | guauuacaga | 1560 |
| auucaaaaga  | gaaaucgagu                       | ggaucaaaau | gauaagccgu  | auguauuaac | aaacaag    | 1617 |
| <210 > SEQ : <211 > LENG' <212 > TYPE <213 > ORGAI <400 > SEQUI | IH: 1716<br>: RNA<br>NISM: Human | parainflue | nza virus 3 |            |            |      |
| auggaauacu  | ggaagcacac                       | caaccacgga | aaggaugcug  | guaaugagcu | ggagacaucc | 60   |
| acagccacuc  | auggcaacaa                       | gcucaccaac | aagauaacau  | auauauugug | gacgauaacc | 120  |
| cugguguuau  | uaucaauagu                       | cuucaucaua | gugcuaacua  | auuccaucaa | aagugaaaag | 180  |
| gcccgcgaau  | cauugcuaca                       | agacauaaau | aaugaguuua  | uggaaguuac | agaaaagauc | 240  |
| caaguggcau  | cggauaauac                       | uaaugaucua | auacagucag  | gagugaauac | aaggcuucuu | 300  |
| acaauucaga  | gucaugucca                       | gaauuauaua | ccaauaucau  | ugacacaaca | aauaucggau | 360  |
| cuuaggaaau  | ucauuaguga                       | aauuacaauu | agaaaugaua  | aucaagaagu | gccaccacaa | 420  |
| agaauaacac  | augauguggg                       | uauaaaaccu | uuaaauccag  | augauuucug | gagaugcacg | 480  |
| ucuggucuuc  | caucuuugau                       | gaaaacucca | aaaauaagau  | uaaugccggg | accaggauua | 540  |
| uuagcuaugc  | caacgacugu                       | ugauggcugu | gucagaaccc  | cguccuuagu | gauaaaugau | 600  |
| cugauuuaug  | cuuacaccuc                       | aaaucuaauu | acucgagguu  | gccaggauau | agggaaauca | 660  |
| uaucaaguau  | uacagauagg                       | gauaauaacu | guaaacucag  | acuugguacc | ugacuuaaau | 720  |
| ccuaggaucu  | cucauaccuu                       | caacauaaau | gacaauagaa  | agucauguuc | ucuagcacuc | 780  |
| cuaaauacag  | auguauauca                       | acuguguuca | accccaaaag  | uugaugaaag | aucagauuau | 840  |
| gcaucaucag  | gcauagaaga                       | uauuguacuu | gauauuguca  | auuaugaugg | cucaaucucg | 900  |
| acaacaagau  | uuaagaauaa                       | uaauauaagu | uuugaucaac  | cauaugegge | auuauaccca | 960  |
| ucuguuggac  | cagggauaua                       | cuacaaaggc | aaaauaauau  | uucucgggua | uggaggucuu | 1020 |
| gaacauccaa  | uaaaugagaa                       | ugcaaucugc | aacacaacug  | gguguccugg | gaaaacacag | 1080 |
| agagacugua  | aucaagcauc                       | ucauagucca | ugguuuucag  | auagaaggau | ggucaacucu | 1140 |
| auaauuguug  | uugacaaggg                       | cuugaacuca | guuccaaaau  | ugaagguaug | gacgauaucu | 1200 |
| augagacaaa  | auuacugggg                       | gucagaagga | agauuacuuc  | uacuagguaa | caagaucuac | 1260 |
| auauacacaa  | gaucuacaag                       | uuggcacagc | aaguuacaau  | uaggaauaau | ugacauuacu | 1320 |
| gacuacagug  | auauaaggau                       | aaaauggaca | uggcauaaug  | ugcuaucaag | accaggaaac | 1380 |
| aaugaauguc  | cauggggaca                       | uucauguccg | gauggaugua  | uaacgggagu | auauaccgau | 1440 |
|   |                                  |            |             |            |            |      |

gcauauccac ucaaucccac aggaagcauu guaucaucug ucauauugga cucacaaaaa 1500

-continued

|             |   |                              |            |            | 1404       |      |  |
|-------------|---|------------------------------|------------|------------|------------|------|--|
| ucgagaguca  | acccagucau                                | aacuuacuca                   | acagcaaccg | aaaggguaaa | cgagcuggcu | 1560 |  |
| auccgaaaca  | aaacacucuc                                | agcuggguac                   | acaacaacaa | gcugcauuac | acacuauaac | 1620 |  |
| aaaggguauu  | guuuucauau                                | aguagaaaua                   | aaucauaaaa | gcuuaaacac | auuucaaccc | 1680 |  |
| auguuguuca  | aaacagagau                                | uccaaaaagc                   | ugcagu     |            |            | 1716 |  |
| <220> FEATU | TH: 1716<br>: RNA<br>NISM: Artif:<br>JRE: | icial Sequer<br>DN: Syntheti |            | eotide     |            |      |  |
| <400> SEQUE | ENCE: 63                                  |                              |            |            |            |      |  |
| auggaauacu  | ggaagcacac                                | caaccacggc                   | aaggacgccg | gcaacgagcu | ggaaaccagc | 60   |  |
| acagccacac  | acggcaacaa                                | gcugaccaac                   | aagaucaccu | acauccugug | gaccaucacc | 120  |  |
| cuggugcugc  | ugagcaucgu                                | guucaucauc                   | gugcugacca | auagcaucaa | gagcgagaag | 180  |  |
| gccagagaga  | gccugcugca                                | ggacaucaac                   | aacgaguuca | uggaagugac | cgagaagauc | 240  |  |
| cagguggcca  | gcgacaacac                                | caacgaccug                   | auccagagcg | gcgugaacac | ccggcugcug | 300  |  |
| accauccaga  | gccacgugca                                | gaacuacauc                   | cccaucagcc | ugacccagca | gaucagcgac | 360  |  |
| cugcggaagu  | ucaucagcga                                | gaucaccauc                   | cggaacgaca | accaggaagu | gccccccag  | 420  |  |
| agaaucaccc  | acgacguggg                                | caucaagccc                   | cugaaccccg | acgauuucug | gcgguguaca | 480  |  |
| ageggeeuge  | ccagccugau                                | gaagaccccc                   | aagauccggc | ugaugccugg | cccuggacug | 540  |  |
| cuggccaugc  | cuaccacagu                                | ggauggcugu                   | gugeggaeee | ccagccucgu | gaucaacgau | 600  |  |
| cugaucuacg  | ccuacaccag                                | caaccugauc                   | acccggggcu | gccaggauau | cggcaagagc | 660  |  |
| uaccaggugc  | ugcagaucgg                                | caucaucacc                   | gugaacuccg | accuggugcc | cgaccugaac | 720  |  |
| ccucggauca  | gccacaccuu                                | caacaucaac                   | gacaacagaa | agagcugcag | ccuggcucug | 780  |  |
| cugaacaccg  | acguguacca                                | gcugugcagc                   | acccccaagg | uggacgagag | aagcgacuac | 840  |  |
| gccagcagcg  | gcaucgagga                                | uaucgugcug                   | gacaucguga | acuacgacgg | cagcaucagc | 900  |  |
| accacccggu  | ucaagaacaa                                | caacaucagc                   | uucgaccagc | ccuacgccgc | ccuguacccu | 960  |  |
| ucugugggcc  | cuggcaucua                                | cuacaagggc                   | aagaucaucu | uccugggcua | cggcggccug | 1020 |  |
| gaacacccca  | ucaacgagaa                                | cgccaucugc                   | aacaccaccg | gcugcccugg | caagacccag | 1080 |  |
| agagacugca  | aucaggccag                                | ccacagcccc                   | ugguucagcg | accgcagaau | ggucaacucu | 1140 |  |
| aucaucgugg  | uggacaaggg                                | ccugaacagc                   | gugcccaagc | ugaaagugug | gacaaucagc | 1200 |  |
| augegeeaga  | acuacugggg                                | cagcgagggc                   | agacuucugc | ugcugggaaa | caagaucuac | 1260 |  |
| aucuacaccc  | gguccaccag                                | cuggcacagc                   | aaacugcagc | ugggaaucau | cgacaucacc | 1320 |  |
| gacuacagcg  | acauccggau                                | caaguggacc                   | uggcacaacg | ugcugagcag | acccggcaac | 1380 |  |
| aaugagugcc  | cuuggggcca                                | cagcugcccc                   | gauggaugua | ucaccggcgu | guacaccgac | 1440 |  |
| gccuaccccc  | ugaauccuac                                | cggcuccauc                   | guguccagcg | ugauccugga | cagccagaaa | 1500 |  |
| agcagaguga  | accccgugau                                | cacauacage                   | accgccaccg | agagagugaa | cgaacuggcc | 1560 |  |
| aucagaaaca  | agacccugag                                | cgccggcuac                   | accaccacaa | gcugcaucac | acacuacaac | 1620 |  |
| aagggcuacu  | gcuuccacau                                | cguggaaauc                   | aaccacaagu | cccugaacac | cuuccagccc | 1680 |  |
| augcuguuca  | agaccgagau                                | ccccaagagc                   | ugcucc     |            |            | 1716 |  |
|             |   |                              |            |            |            |      |  |

<210> SEQ ID NO 64 <211> LENGTH: 1617

571 572

<212> TYPE: RNA <213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEOUENCE: 64

augeccauca geauceugeu gaucaucace acaaugauca uggecageea eugecagaue 60 gacaucacca agcugcagca cgugggcgug cucgugaaca gccccaaggg caugaagauc 120 agccagaacu ucgagacacg cuaccugauc cugagccuga uccccaagau cgaggacagc 180 aacagcugcg gcgaccagca gaucaagcag uacaagcggc ugcuggacag acugaucauc 240 ccccuguacg acggccugcg gcugcagaaa gacgugaucg ugaccaacca ggaaagcaac gagaacaccg acccccggac cgagagauuc uucggcggcg ugaucggcac aaucgcccug 360 420 qqaquqqcca caaqcqccca qauuacaqcc qcuquqqccc uqquqqaaqc caaqcaqqcc agaagegaca ucgagaageu gaaagaggee aucegggaca ecaacaagge egugeagage 480 gugcagucca gcgugggcaa ucugaucgug gccaucaagu ccgugcagga cuacgugaac 540 600 aaaqaaaucq uqcccucuau cqcccqqcuq qqcuquqaaq cuqccqqacu qcaqcuqqqc auugeeeuga cacageaeua cagegageug accaaeaueu ueggegaeaa caueggeage 660 cuqcaqqaaa aqqqcauuaa qcuqcaqqqa aucqccaqcc uquaccqcac caacaucacc 720 gagaucuuca ccaccagcac cguggauaag uacgacaucu acgaccugcu guucaccgag 780 840 aqcaucaaaq uqcqcquqau cqacquqqac cuqaacqacu acaqcaucac ccuqcaaquq eggeugeece ugeugaceag acugeugaac acceagaueu acaaggugga eageaueuee 900 960 uacaacauce agaacegega gugguacauc ceucugeeca gecacauuau gaccaaggge gccuuucugg gcggagccga cgugaaagag ugcaucgagg ccuucagcag cuacaucugc 1020 cccagcgacc cuggcuucgu gcugaaccac gagauggaaa gcugccugag cggcaacauc 1080 agccagugec ccagaaccac egugaccuce gacaueguge ecagauaege euuegugaau 1140 ggcggcgugg uggccaacug caucaccacc accuguaccu gcaacggcau cggcaaccgg 1200 aucaaccage cuccegauca gggegugaag auuaucaeee acaaagagug uaacaecaue 1260 ggcaucaacg gcaugcuguu caauaccaac aaagagggca cccuggccuu cuacaccccc 1320 gacgauauca cccugaacaa cuccguggcu cuggacccca ucgacaucuc caucgagcug 1380 aacaaggcca agagcgaccu ggaagagucc aaagagugga uccggcggag caaccagaag 1440 cuggacucua ucggcagcug gcaccagagc agcaccacca ucaucgugau ccugauuaug 1500 augauuaucc uguucaucau caacauuacc aucaucacua ucgccauuaa guacuaccgg 1560 auccagaaac ggaaccgggu ggaccagaau gacaagcccu acgugcugac aaacaag 1617

<400> SEQUENCE: 65

| augauacacu | caguguuucu | acugauguuc | uuguuaacac | cuacagaaag | uuacguugau | 60  |
|------------|------------|------------|------------|------------|------------|-----|
| guagggccag | auucuguuaa | gucugcuugu | auugagguug | auauacaaca | gaccuucuuu | 120 |
| gauaaaacuu | ggccuaggcc | aauugauguu | ucuaaggcug | acgguauuau | auacccucaa | 180 |
| ggccguacau | auucuaacau | aacuaucacu | uaucaagguc | uuuuucccua | ucagggagac | 240 |

<sup>&</sup>lt;210> SEQ ID NO 65

<sup>&</sup>lt;211> LENGTH: 4062

<sup>&</sup>lt;212> TYPE: RNA

<sup>&</sup>lt;213 > ORGANISM: Unknown

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Middle East respiratory syndrome coronavirus

| cauggugaua | uguauguuua | cucugcagga | caugcuacag | gcacaacucc | acaaaaguug | 300  |
|------------|------------|------------|------------|------------|------------|------|
| uuuguagcua | acuauucuca | ggacgucaaa | caguuugcua | auggguuugu | cguccguaua | 360  |
| ggagcagcug | ccaauuccac | uggcacuguu | auuauuagcc | caucuaccag | cgcuacuaua | 420  |
| cgaaaaauuu | acccugcuuu | uaugcugggu | ucuucaguug | guaauuucuc | agaugguaaa | 480  |
| augggccgcu | ucuucaauca | uacucuaguu | cuuuugcccg | auggaugugg | cacuuuacuu | 540  |
| agagcuuuuu | auuguauucu | agagccucgc | ucuggaaauc | auuguccugc | uggcaauucc | 600  |
| uauacuucuu | uugccacuua | ucacacuccu | gcaacagauu | guucugaugg | caauuacaau | 660  |
| cguaaugcca | gucugaacuc | uuuuaaggag | uauuuuaauu | uacguaacug | caccuuuaug | 720  |
| uacacuuaua | acauuaccga | agaugagauu | uuagaguggu | uuggcauuac | acaaacugcu | 780  |
| caagguguuc | accucuucuc | aucucgguau | guugauuugu | acggcggcaa | uauguuucaa | 840  |
| uuugccaccu | ugccuguuua | ugauacuauu | aaguauuauu | cuaucauucc | ucacaguauu | 900  |
| cguucuaucc | aaagugauag | aaaagcuugg | gcugccuucu | acguauauaa | acuucaaccg | 960  |
| uuaacuuucc | uguuggauuu | uucuguugau | gguuauauac | gcagagcuau | agacuguggu | 1020 |
| uuuaaugauu | ugucacaacu | ccacugcuca | uaugaauccu | ucgauguuga | aucuggaguu | 1080 |
| uauucaguuu | cgucuuucga | agcaaaaccu | ucuggcucag | uuguggaaca | ggcugaaggu | 1140 |
| guugaaugug | auuuuucacc | ucuucugucu | ggcacaccuc | cucagguuua | uaauuucaag | 1200 |
| cguuugguuu | uuaccaauug | caauuauaau | cuuaccaaau | ugcuuucacu | uuuuucugug | 1260 |
| aaugauuuua | cuuguaguca | aauaucucca | gcagcaauug | cuagcaacug | uuauucuuca | 1320 |
| cugauuuugg | auuauuuuc  | auacccacuu | aguaugaaau | ccgaucucag | uguuaguucu | 1380 |
| gcugguccaa | uaucccaguu | uaauuauaaa | caguccuuuu | cuaaucccac | auguuugauc | 1440 |
| uuagcgacug | uuccucauaa | ccuuacuacu | auuacuaagc | cucuuaagua | cagcuauauu | 1500 |
| aacaagugcu | cucgucuucu | uucugaugau | cguacugaag | uaccucaguu | agugaacgcu | 1560 |
| aaucaauacu | cacccugugu | auccauuguc | ccauccacug | ugugggaaga | cggugauuau | 1620 |
| uauaggaaac | aacuaucucc | acuugaaggu | gguggcuggc | uuguugcuag | uggcucaacu | 1680 |
| guugccauga | cugagcaauu | acagaugggc | uuugguauua | caguucaaua | ugguacagac | 1740 |
| accaauagug | uuugccccaa | gcuugaauuu | gcuaaugaca | caaaaauugc | cucucaauua | 1800 |
| ggcaauugcg | uggaauauuc | ccucuauggu | guuucgggcc | gugguguuuu | ucagaauugc | 1860 |
| acagcuguag | guguucgaca | gcagcgcuuu | guuuaugaug | cguaccagaa | uuuaguuggc | 1920 |
| uauuauucug | augauggcaa | cuacuacugu | cugcgugcuu | guguuagugu | uccuguuucu | 1980 |
| gucaucuaug | auaaagaaac | uaaaacccac | gcuacucuau | uugguagugu | ugcaugugaa | 2040 |
| cacauuucuu | cuaccauguc | ucaauacucc | cguucuacgc | gaucaaugcu | uaaacggcga | 2100 |
| gauucuacau | auggcccccu | ucagacaccu | guugguugug | uccuaggacu | uguuaauucc | 2160 |
| ucuuuguucg | uagaggacug | caaguugccu | cucggucaau | cucucugugc | ucuuccugac | 2220 |
| acaccuagua | cucucacacc | ucgcagugug | cgcucugugc | caggugaaau | gcgcuuggca | 2280 |
| uccauugcuu | uuaaucaucc | cauucagguu | gaucaacuua | auaguaguua | uuuuaaauua | 2340 |
| aguauaccca | cuaauuuuuc | cuuuggugug | acucaggagu | acauucagac | aaccauucag | 2400 |
| aaaguuacug | uugauuguaa | acaguacguu | ugcaaugguu | uccagaagug | ugagcaauua | 2460 |
| cugcgcgagu | auggccaguu | uuguuccaaa | auaaaccagg | cucuccaugg | ugccaauuua | 2520 |
| cgccaggaug | auucuguacg | uaauuuguuu | gcgagcguga | aaagcucuca | aucaucuccu | 2580 |
| aucauaccag | guuuuggagg | ugacuuuaau | uugacacuuc | uagaaccugu | uucuauaucu | 2640 |
|            |            |            |            |            |            |      |

| acuggcaguc   | guagugcacg   | uagugcuauu | gaggauuugc | uauuugacaa | agucacuaua | 2700 |  |  |
|--|--|------------|------------|------------|------------|------|--|--|
| gcugauccug   | guuauaugca   | agguuacgau | gauuguaugc | agcaaggucc | agcaucagcu | 2760 |  |  |
| cgugaucuua   | uuugugcuca   | auauguggcu | gguuauaaag | uauuaccucc | ucuuauggau | 2820 |  |  |
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| ccucuucucg   | gcaauuccac   | cgggauugac | uuccaagaug | aguuggauga | guuuuucaaa | 3720 |  |  |
| aauguuagca   | ccaguauacc   | uaauuuuggu | ucucuaacac | agauuaauac | uacauuacuc | 3780 |  |  |
| gaucuuaccu   | acgagauguu   | gucucuucaa | caaguuguua | aagcccuuaa | ugagucuuac | 3840 |  |  |
| auagaccuua   | aagagcuugg   | caauuauacu | uauuacaaca | aauggccgug | guacauuugg | 3900 |  |  |
| cuugguuuca   | uugcugggcu   | uguugccuua | gcucuaugcg | ucuucuucau | acugugcugc | 3960 |  |  |
| acugguugug   | gcacaaacug   | uaugggaaaa | cuuaagugua | aucguuguug | ugauagauac | 4020 |  |  |
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| gauaaaacuu   | ggccuaggcc   | aauugauguu | ucuaaggcug | acgguauuau | auacccucaa | 180  |  |  |
| ggccguacau   | auucuaacau   | aacuaucacu | uaucaagguc | uuuuucccua | ucagggagac | 240  |  |  |
| cauggugaua   | uguauguuua   | cucugcagga | caugcuacag | gcacaacucc | acaaaaguug | 300  |  |  |
| uuuguagcua   | acuauucuca   | ggacgucaaa | caguuugcua | auggguuugu | cguccguaua | 360  |  |  |
| ggagcagcug   | ccaauuccac   | uggcacuguu | auuauuagcc | caucuaccag | cgcuacuaua | 420  |  |  |
| cgaaaaauuu   | acccugcuuu   | uaugcugggu | ucuucaguug | guaauuucuc | agaugguaaa | 480  |  |  |
| augggccgcu   | ucuucaauca   | uacucuaguu | cuuuugcccg | auggaugugg | cacuuuacuu | 540  |  |  |

agagcuuuuu auuguauucu ggagccucgc ucuggaaauc auuguccugc uggcaauucc

| uauacuucuu | uugccacuua | ucacacuccu | gcaacagauu | guucugaugg | caauuacaau | 660  |
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| cguaaugcca | gucugaacuc | uuuuaaggag | uauuuuaauu | uacguaacug | caccuuuaug | 720  |
| uacacuuaua | acauuaccga | agaugagauu | uuagaguggu | uuggcauuac | acaaacugcu | 780  |
| caagguguuc | accucuucuc | aucucgguau | guugauuugu | acggcggcaa | uauguuucaa | 840  |
| uuugccaccu | ugccuguuua | ugauacuauu | aaguauuauu | cuaucauucc | ucacaguauu | 900  |
| cguucuaucc | aaagugauag | aaaagcuugg | gcugccuucu | acguauauaa | acuucaaccg | 960  |
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| acagcuguag | guguucgaca | gcagcgcuuu | guuuaugaug | cguaccagaa | uuuaguuggc | 1920 |
| uauuauucug | augauggcaa | cuacuacugu | uugcgugcuu | guguuagugu | uccuguuucu | 1980 |
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| aguauaccca | cuaauuuuuc | cuuuggugug | acucaggagu | acauucagac | aaccauucag | 2400 |
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| aucauaccag | guuuuggagg | ugacuuuaau | uugacacuuc | uggaaccugu | uucuauaucu | 2640 |
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|            | gcauuacuca |            |            |            |            | 3000 |
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|                            |   |                              | cagcugcuga          |            |            | 360  |
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| guggccgggu                 | acaaagugcu                                | gccuccucug                   | auggauguga          | acauggaggc | cgcuuauacu | 720  |
| ucgucccugc                 | ucggcucuau                                | cgccggcgug                   | ggguggaccg          | ccggccuguc | cuccuucgcc | 780  |
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| agcuaugugg acgugggccc cgauagcgug aaguccgccu guaucgaagu ggacauccag cagaccuuuu ucgacaagac cuggccaga cccaucgacg uguccaaggc cgacggcauc aucuauccac aaggccggac cuacagcaac aucaccauua ccuaccaggg ccuguuccca uaucaaggcg accacggcga uauguacgug uacucugccg gccacgccac   | 120 180 240 300 360 420 480 540 600 660 720 780 840 900 960               |

1200

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| ugcuacagca | gccugauccu | ggacuacuuc | agcuaccccc | ugagcaugaa | guccgaucug | 1380 |
| agcguguccu | ccgccggacc | caucagccag | uucaacuaca | agcagagcuu | cagcaacccu | 1440 |
| accugccuga | uucuggccac | cgugccccac | aaucugacca | ccaucaccaa | gccccugaag | 1500 |
| uacagcuaca | ucaacaagug | cagcagacug | cuguccgacg | accggaccga | agugccccag | 1560 |
| cucgugaacg | ccaaccagua | cagccccugc | guguccaucg | ugcccagcac | cgugugggag | 1620 |
| gacggcgacu | acuacagaaa | gcagcugagc | ccccuggaag | geggeggaug | gcugguggcu | 1680 |
| ucuggaagca | caguggccau | gaccgagcag | cugcagaugg | gcuuuggcau | caccgugcag | 1740 |
| uacggcaccg | acaccaacag | cgugugcccc | aagcuggaau | ucgccaauga | caccaagauc | 1800 |
| gccagccagc | ugggaaacug | cguggaauac | ucccuguaug | gcguguccgg | acggggcgug | 1860 |
| uuccagaauu | gcacagcagu | gggagugcgg | cagcagagau | ucguguacga | ugccuaccag | 1920 |
| aaccucgugg | gcuacuacag | cgacgacggc | aauuacuacu | gccugcgggc | cugugugucc | 1980 |
| gugcccgugu | ccgugaucua | cgacaaagag | acaaagaccc | acgccacacu | guucggcucc | 2040 |
| guggccugcg | agcacaucag | cuccaccaug | agccaguacu | cccgcuccac | ccgguccaug | 2100 |
| cugaagcgga | gagauagcac | cuacggcccc | cugcagacac | cugugggaug | ugugcugggc | 2160 |
| cucgugaaca | gcucccuguu | uguggaagau | ugcaagcugc | cccugggcca | gagccugugu | 2220 |
| gcccugccag | auaccccuag | cacccugacc | ccuagaagcg | ugcgcucugu | gcccggcgaa | 2280 |
| augcggcugg | ccucuaucgc | cuucaaucac | cccauccagg | uggaccagcu | gaacuccagc | 2340 |
| uacuucaagc | ugagcauccc | caccaacuuc | agcuucggcg | ugacccagga | guacauccag | 2400 |
| accacaaucc | agaaagugac | cguggacugc | aagcaguacg | ugugcaacgg | cuuucagaag | 2460 |
| ugcgaacagc | ugcugcgcga | guacggccag | uucugcagca | agaucaacca | ggcccugcac | 2520 |
| ggcgccaacc | ugagacagga | ugacagcgug | cggaaccugu | ucgccagcgu | gaaaagcagc | 2580 |
| caguccagcc | ccaucauccc | uggcuucggc | ggcgacuuua | accugacccu | gcuggaaccu | 2640 |
| guguccauca | gcaccggcuc | cagaagcgcc | agauccgcca | ucgaggaccu | gcuguucgac | 2700 |
| aaagugacca | uugccgaccc | cggcuacaug | cagggcuacg | acgauugcau | gcagcagggc | 2760 |
| ccagccagcg | ccagggaucu | gaucugugcc | caguaugugg | ccggcuacaa | ggugcugccc | 2820 |
| ccccugaugg | acgugaacau | ggaagccgcc | uacaccucca | gccugcuggg | cucuauugcu | 2880 |
| ggcgugggau | ggacagccgg | ccugucuagc | uuugccgcca | ucccuuucgc | ccagagcauc | 2940 |
| uucuaccggc | ugaacggcgu | gggcaucaca | caacaggugc | ugagcgagaa | ccagaagcug | 3000 |
| aucgccaaca | aguuuaacca | ggcacugggc | gccaugcaga | ccggcuucac | caccaccaac | 3060 |
| gaggccuuca | gaaaggugca | ggacgccgug | aacaacaacg | cccaggcucu | gagcaagcug | 3120 |
| gccuccgagc | ugagcaauac | cuucggcgcc | aucagegeeu | ccaucggcga | caucauccag | 3180 |
| cggcuggacg | ugcuggaaca | ggacgcccag | aucgaccggc | ugaucaacgg | cagacugacc | 3240 |
| acccugaacg | ccuucguggc | acagcagcuc | gugcggagcg | aaucugccgc | ucugucugcu | 3300 |
| cagcuggcca | aggacaaagu | gaacgagugc | gugaaggccc | aguccaagcg | gagcggcuuu | 3360 |
| uguggccagg | gcacccacau | cguguccuuc | gucgugaaug | ccccaacgg  | ccuguacuuu | 3420 |
| augcacgugg | gcuauuaccc | cagcaaccac | aucgaggugg | uguccgccua | uggccugugc | 3480 |
| gacgccgcca | auccuaccaa | cuguaucgcc | cccgugaacg | gcuacuucau | caagaccaac | 3540 |
|            |            |            |            |            |            |      |

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|--|------|
| aacaccegga ueguggaega gugguecuae acaggeagea geuucuaege eecegageee  | 3600 |
| aucaccucce ugaacaccaa auacguggee eeccaaguga cauaccagaa caucuccaec  | 3660 |
| aaccugcccc cuccacugcu gggaaauucc accggcaucg acuuccagga cgagcuggac  | 3720 |
| gaguucuuca agaacguguc caccuccauc cccaacuucg gcagccugac ccagaucaac  | 3780 |
| accacucuge uggaccugae cuacgagaug cugucccuge aacaggucgu gaaagcccug  | 3840 |
| aacgagagcu acaucgaccu gaaagagcug gggaacuaca ccuacuacaa caaguggccu  | 3900 |
| ugguacauuu ggcugggcuu uaucgccggc cugguggccc uggcccugug cguguucuuc  | 3960 |
| auccugugcu gcaccggcug cggcaccaau ugcaugggca agcugaaaug caaccggugc  | 4020 |
| ugcgacagau acgaggaaua cgaccuggaa ccucacaaag ugcaugugca c   | 4071 |
| <210> SEQ ID NO 69 <211> LENGTH: 1864 <212> TYPE: RNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |      |
| <400> SEQUENCE: 69   |      |
| ucaagcuuuu ggacccucgu acagaagcua auacgacuca cuauagggaa auaagagaga aaaqaaqaqu aaqaagaaau auaaqagcca ccaugggucu caaggugaac gucucugccg                        | 120  |
| uauucauggo aguacuguua acucuccaaa cacccgccgg ucaaauucau uggggcaauc  | 180  |
| ucucuaagau agggguagua ggaauaggaa gugcaagcua caaaguuaug acucguucca  | 240  |
| gccaucaauc auuagucaua aaauuaaugc ccaauauaac ucuccucaau aacugcacga  | 300  |
| ggguagagau ugcagaauac aggagacuac uaagaacagu uuuggaacca auuagggaug  | 360  |
| cacuuaaugc aaugacccag aacauaaggc cgguucagag cguagcuuca aguaggagac  | 420  |
| acaagagauu ugcgggagua guccuggcag gugcggcccu agguguugcc acagcugcuc  | 480  |
| agauaacage eggeauugea euucaeeggu eeaugeugaa eucucaggee auegaeaaue  | 540  |
| ugagagcgag ccuggaaacu acuaaucagg caauugaggc aaucagacaa gcagggcagg  | 600  |
| agaugauauu ggcuguucag gguguccaag acuacaucaa uaaugagcug auaccgucua  | 660  |
| ugaaccagcu aucuugugau cuaaucgguc agaagcucgg gcucaaauug cuuagauacu  | 720  |
| auacagaaau ccugucauua uuuggcccca gccuacggga ccccauaucu gcggagauau  | 780  |
| cuauccaggc uuugaguuau gcacuuggag gagauaucaa uaagguguua gaaaagcucg  | 840  |
| gauacagugg aggcgauuua cuaggcaucu uagagagcag aggaauaaag gcucggauaa  | 900  |
| cucacguega cacagaguee uacuucauag uecucaguau agecuaueeg aegeugueeg  | 960  |
| agauuaaggg ggugauuguc caccggcuag agggggucuc guacaacaua ggcucucaag  | 1020 |
| agugguauac cacugugccc aaguauguug caacccaagg guaccuuauc ucgaauuuug  | 1080 |
| augagucauc auguacuuuc augccagagg ggacugugug cagccaaaau gccuuguacc  | 1140 |
| cgaugaguce ucugcuccaa gaaugccuce ggggguccae caaguccugu gcucguacae  | 1200 |
| ucguauccgg gucuuuuggg aaccgguuca uuuuaucaca agggaaccua auagccaauu  | 1260 |
| gugcaucaau ucuuuguaag uguuacacaa cagguacgau uauuaaucaa gacccugaca  | 1320 |
| agauccuaac auacauugcu gccgaucgcu gcccgguagu cgaggugaac ggcgugacca  | 1380 |
| uccaagucgg gagcaggagg uauccagacg cuguguacuu gcacagaauu gaccucgguc  | 1440 |
| cucccauauc auuggagagg uuggacguag ggacaaaucu ggggaaugca auugccaaau  | 1500 |
|  |      |

uggaggaugc caaggaauug uuggaaucau cggaccagau auugagaagu augaaagguu 1560

587 588

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| uaucgagcac   | uagcauaguc   | uacauccuga | uugcagugug | ucuuggaggg | uugauaggga | 1620 |  |
|--|--|------------|------------|------------|------------|------|--|
| uccccacuuu   | aauauguugc   | ugcagggggc | guuguaacaa | aaagggagaa | caaguuggua | 1680 |  |
| ugucaagacc   | aggccuaaag   | ccugaccuua | caggaacauc | aaaauccuau | guaagaucgc | 1740 |  |
| uuugaugaua   | auaggcugga   | gccucggugg | ccaagcuucu | ugccccuugg | gccuccccc  | 1800 |  |
| agccccuccu   | ccccuuccug   | cacccguacc | cccguggucu | uugaauaaag | ucugaguggg | 1860 |  |
| cggc   |  |            |            |            |            | 1864 |  |
| <211> LENG'<br><212> TYPE<br><213> ORGAL<br><220> FEAT | <210> SEQ ID NO 70 <211> LENGTH: 1653 <212> TYPE: RNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |            |            |            |            |      |  |
| <400> SEQUI  | ENCE: 70   |            |            |            |            |      |  |
| augggucuca   | aggugaacgu   | cucugeegua | uucauggcag | uacuguuaac | ucuccaaaca | 60   |  |
| cccgccgguc   | aaauucauug   | gggcaaucuc | ucuaagauag | ggguaguagg | aauaggaagu | 120  |  |
| gcaagcuaca   | aaguuaugac   | ucguuccagc | caucaaucau | uagucauaaa | auuaaugccc | 180  |  |
| aauauaacuc   | uccucaauaa   | cugcacgagg | guagagauug | cagaauacag | gagacuacua | 240  |  |
| agaacaguuu   | uggaaccaau   | uagggaugca | cuuaaugcaa | ugacccagaa | cauaaggccg | 300  |  |
| guucagagcg   | uagcuucaag   | uaggagacac | aagagauuug | cgggaguagu | ccuggcaggu | 360  |  |
| gcggcccuag   | guguugccac   | agcugcucag | auaacagccg | gcauugcacu | ucaccggucc | 420  |  |
| augcugaacu   | cucaggccau   | cgacaaucug | agagcgagcc | uggaaacuac | uaaucaggca | 480  |  |
| auugaggcaa   | ucagacaagc   | agggcaggag | augauauugg | cuguucaggg | uguccaagac | 540  |  |
| uacaucaaua   | augagcugau   | accgucuaug | aaccagcuau | cuugugaucu | aaucggucag | 600  |  |
| aagcucgggc   | ucaaauugcu   | uagauacuau | acagaaaucc | ugucauuauu | uggccccagc | 660  |  |
| cuacgggacc   | ccauaucugc   | ggagauaucu | auccaggcuu | ugaguuaugc | acuuggagga | 720  |  |
| gauaucaaua   | agguguuaga   | aaagcucgga | uacaguggag | gcgauuuacu | aggcaucuua | 780  |  |
| gagagcagag   | gaauaaaggc   | ucggauaacu | cacgucgaca | cagaguccua | cuucauaguc | 840  |  |
| cucaguauag   | ccuauccgac   | gcuguccgag | auuaaggggg | ugauugucca | ccggcuagag | 900  |  |
| ggggucucgu   | acaacauagg   | cucucaagag | ugguauacca | cugugcccaa | guauguugca | 960  |  |
| acccaagggu   | accuuaucuc   | gaauuuugau | gagucaucau | guacuuucau | gccagagggg | 1020 |  |
| acugugugca   | gccaaaaugc   | cuuguacccg | augaguccuc | ugcuccaaga | augccuccgg | 1080 |  |
| ggguccacca   | aguccugugc   | ucguacacuc | guauccgggu | cuuuugggaa | ccgguucauu | 1140 |  |
| uuaucacaag   | ggaaccuaau   | agccaauugu | gcaucaauuc | uuuguaagug | uuacacaaca | 1200 |  |
| gguacgauua   | uuaaucaaga   | cccugacaag | auccuaacau | acauugcugc | cgaucgcugc | 1260 |  |
| ccgguagucg   | aggugaacgg   | cgugaccauc | caagucggga | gcaggaggua | uccagacgcu | 1320 |  |
| guguacuugc   | acagaauuga   | ccucgguccu | cccauaucau | uggagagguu | ggacguaggg | 1380 |  |
| acaaaucugg   | ggaaugcaau   | ugccaaauug | gaggaugcca | aggaauuguu | ggaaucaucg | 1440 |  |
|  | ugagaaguau   |            |            |            |            | 1500 |  |
|  | uuggaggguu   |            |            |            |            | 1560 |  |
|  |  |            |            |            |            | 1620 |  |
| uyuaacaada   | agggagaaca   | ayuuyguaug | ucaayaccag | geedaaagee | ugaccuuaca | 1020 |  |

ggaacaucaa aauccuaugu aagaucgcuu uga

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<210> SEQ ID NO 71

<211> LENGTH: 1925

<212> TYPE: RNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEQUENCE: 71

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<212> TYPE: RNA

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEQUENCE: 72

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<sup>&</sup>lt;210> SEQ ID NO 73

<sup>&</sup>lt;211> LENGTH: 1653

<sup>&</sup>lt;212> TYPE: RNA

<sup>&</sup>lt;213> ORGANISM: Artificial Sequence

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223 > OTHER INFORMATION: Synthetic Polynucleotide

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| <400> SEQUENCE: 73  |      |
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| augggucuca aggugaacgu cucugucaua uucauggcag uacuguuaac ucuucaaaca   | 60   |
| cccaccgguc aaauccauug gggcaaucuc ucuaagauag gggugguagg gguaggaagu   | 120  |
| gcaagcuaca aaguuaugac ucguuccagc caucaaucau uagucauaaa guuaaugccc   | 180  |
| aauauaacuc uccucaacaa uugcacgagg guagggauug cagaauacag gagacuacug   | 240  |
| agaacaguuc uggaaccaau uagagaugca cuuaaugcaa ugacccagaa uauaagaccg   | 300  |
| guucagagug uagcuucaag uaggagacac aagagauuug cgggaguugu ccuggcaggu   | 360  |
| geggeecuag geguugeeac ageugeucaa auaacageeg guauugeacu ucaecaguee   | 420  |
| augcugaacu cucaagccau cgacaaucug agagcgagcc uagaaacuac uaaucaggca   | 480  |
| auugaggcaa ucagacaagc agggcaggag augauauugg cuguucaggg uguccaagac   | 540  |
| uacaucaaua augagcugau accgucuaug aaucaacuau cuugugauuu aaucggccag   | 600  |
| aagcuagggc ucaaauugcu cagauacuau acagaaaucc ugucauuauu uggccccagc   | 660  |
| uuacgggacc ccauaucugc ggagauaucu auccaggcuu ugagcuaugc gcuuggagga   | 720  |
| gauaucaaua agguguugga aaagcucgga uacaguggag gugaucuacu gggcaucuua   | 780  |
| gagagcagag gaauaaaggc ccggauaacu cacgucgaca cagaguccua cuucauugua   | 840  |
| cucaguauag ccuauccgac gcuauccgag auuaaggggg ugauugucca ccggcuagag   | 900  |
| ggggucucgu acaacauagg cucucaagag ugguauacca cugugcccaa guauguugca   | 960  |
| acccaagggu accuuaucuc gaauuuugau gagucaucau gcacuuucau gccagagggg   | 1020 |
| acugugugca gccagaaugc cuuguacceg augaguccuc ugcuccaaga augccuccgg   | 1080 |
| ggguccacua aguccugugc ucguacacuc guauccgggu cuuucgggaa ccgguucauu   | 1140 |
| uuaucacagg ggaaccuaau agccaauugu gcaucaaucc uuugcaagug uuacacaaca   | 1200 |
| ggaacaauca uuaaucaaga cccugacaag auccuaacau acauugcugc cgaucacugc   | 1260 |
| ccgguggucg aggugaaugg cgugaccauc caagucggga gcaggaggua uccggacgcu   | 1320 |
| guguacuugc acaggauuga ccucgguccu cccauaucuu uggagagguu ggacguaggg   | 1380 |
| acaaaucugg ggaaugcaau ugcuaaguug gaggaugcca aggaauuguu ggagucaucg   | 1440 |
| gaccagauau ugaggaguau gaaagguuua ucgagcacua guauaguuua cauccugauu   | 1500 |
| gcaguguguc uuggaggauu gauagggauc cccgcuuuaa uauguugcug cagggggcgu   | 1560 |
| uguaacaaga agggagaaca aguugguaug ucaagaccag gccuaaagcc ugaucuuaca   | 1620 |
| ggaacaucaa aauccuaugu aaggucacuc uga  | 1653 |
| <210> SEQ ID NO 74 <211> LENGTH: 1925 <212> TYPE: RNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide <400> SEQUENCE: 74 |      |
| ggggaaauaa gagagaaaag aagaguaaga agaaauauaa gagccaccau gggucucaag   | 60   |
| gugaacgucu cugucauauu cauggcagua cuguuaacuc uucaaacacc caccggucaa   | 120  |
| auccauuggg gcaaucucuc uaagauaggg gugguagggg uaggaagugc aagcuacaaa   | 180  |
| guuaugacuc guuccagoca ucaaucauua gucauaaagu uaaugoccaa uauaacucuc   | 240  |
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| cucaacaauu gcacgagggu agggauugca gaauacagga gacuacugag aacaguucug   | 300  |

gaaccaauua gagaugcacu uaaugcaaug acccagaaua uaagaccggu ucagagugua

| gcuucaagua                 | ggagacacaa  | gagauuugcg | ggaguugucc | uggcaggugc | ggcccuaggc | 420  |
|----------------------------|---|------------|------------|------------|------------|------|
| guugccacag                 | cugcucaaau  | aacagccggu | auugcacuuc | accaguccau | gcugaacucu | 480  |
| caagccaucg                 | acaaucugag  | agegageeua | gaaacuacua | aucaggcaau | ugaggcaauc | 540  |
| agacaagcag                 | ggcaggagau  | gauauuggcu | guucagggug | uccaagacua | caucaauaau | 600  |
| gagcugauac                 | cgucuaugaa  | ucaacuaucu | ugugauuuaa | ucggccagaa | gcuagggcuc | 660  |
| aaauugcuca                 | gauacuauac  | agaaauccug | ucauuauuug | gccccagcuu | acgggacccc | 720  |
| auaucugcgg                 | agauaucuau  | ccaggcuuug | agcuaugcgc | uuggaggaga | uaucaauaag | 780  |
| guguuggaaa                 | agcucggaua  | caguggaggu | gaucuacugg | gcaucuuaga | gagcagagga | 840  |
| auaaaggccc                 | ggauaacuca  | cgucgacaca | gaguccuacu | ucauuguacu | caguauagcc | 900  |
| uauccgacgc                 | uauccgagau  | uaagggggug | auuguccacc | ggcuagaggg | ggucucguac | 960  |
| aacauaggcu                 | cucaagagug  | guauaccacu | gugcccaagu | auguugcaac | ccaaggguac | 1020 |
| cuuaucucga                 | auuuugauga  | gucaucaugc | acuuucaugc | cagaggggac | ugugugcagc | 1080 |
| cagaaugccu                 | uguacccgau  | gaguccucug | cuccaagaau | gccuccgggg | guccacuaag | 1140 |
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| aaccuaauag                 | ccaauugugc  | aucaauccuu | ugcaaguguu | acacaacagg | aacaaucauu | 1260 |
| aaucaagacc                 | cugacaagau  | ccuaacauac | auugcugccg | aucacugccc | gguggucgag | 1320 |
| gugaauggcg                 | ugaccaucca  | agucgggagc | aggagguauc | cggacgcugu | guacuugcac | 1380 |
| aggauugacc                 | ucgguccucc  | cauaucuuug | gagagguugg | acguagggac | aaaucugggg | 1440 |
| aaugcaauug                 | cuaaguugga  | ggaugccaag | gaauuguugg | agucaucgga | ccagauauug | 1500 |
| aggaguauga                 | aagguuuauc  | gagcacuagu | auaguuuaca | uccugauugc | agugugucuu | 1560 |
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| ggagaacaag                 | uugguauguc  | aagaccaggc | cuaaagccug | aucuuacagg | aacaucaaaa | 1680 |
| uccuauguaa                 | ggucacucug  | augauaauag | gcuggagccu | cgguggccaa | gcuucuugcc | 1740 |
| ccuugggccu                 | cccccagcc   | ccuccucccc | uuccugcacc | cguacccccg | uggucuuuga | 1800 |
| auaaagucug                 | agugggcggc  | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | 1860 |
| aaaaaaaaa                  | aaaaaaaaa   | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | 1920 |
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| aaagaagagu                 | aagaagaaau  | auaagagcca | ccaugucacc | gcaacgagac | cggauaaaug | 120  |
| ccuucuacaa                 | agauaacccu  | uaucccaagg | gaaguaggau | aguuauuaac | agagaacauc | 180  |
| uuaugauuga                 | cagacccuau  | guucugcugg | cuguucuguu | cgucauguuu | cugagcuuga | 240  |
| ucggauugcu                 | ggcaauugca  | ggcauuagac | uucaucgggc | agccaucuac | accgcggaga | 300  |
| uccauaaaag                 | ccucaguacc  | aaucuggaug | ugacuaacuc | caucgagcau | caggucaagg | 360  |

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| gauucacuga ccuagugaaa uucaucucgg acaagauuaa auuccuuaau ccggauaggg  | 480  |
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| augaucaaua cugugcagau guggcugcug aagagcucau gaaugcauug gugaacucaa  | 600  |
| cucuacugga gaccagaaca accacucagu uccuagcugu cucaaaggga aacugcucag  | 660  |
| ggcccacuac aaucagaggu caauucucaa acaugucgcu guccuuguug gacuuguacu  | 720  |
| uaggucgagg uuacaaugug ucaucuauag ucacuaugac aucccaggga auguaugggg  | 780  |
| gaaccuaccu aguugaaaag ccuaaucuga acagcaaagg gucagaguug ucacaacuga  | 840  |
| gcauguaccg aguguuugaa guagguguga ucagaaaccc ggguuugggg gcuccggugu  | 900  |
| uccauaugac aaacuauuuu gagcaaccag ucaguaaugg ucucggcaac uguauggugg  | 960  |
| cuuuggggga gcucaaacuc gcagcccuuu gucacgggga cgauucuauc auaauucccu  | 1020 |
| aucagggauc agggaaaggu gucagcuucc agcucgucaa gcuggguguc uggaaauccc  | 1080 |
| caaccgacau gcaauccugg guccccuuau caacggauga uccaguggua gacaggcuuu  | 1140 |
| accucucauc ucacagaggu gucaucgcug acaaucaagc aaaaugggcu gucccgacaa  | 1200 |
| cacgaacaga ugacaaguug cgaauggaga caugcuucca gcaggcgugu aaagguaaaa  | 1260 |
| uccaagcacu cugcgagaau cccgaguggg uaccauugaa ggauaacagg auuccuucau  | 1320 |
| acgggguccu gucuguugau cugagucuga cgguugagcu uaaaaucaaa auugcuucgg  | 1380 |
| gauucgggcc auugaucaca cacggcucag ggauggaccu auacaaaucc aacugcaaca  | 1440 |
| auguguauug gcugacuauu ccgccaauga gaaaucuagc cuuaggcgua aucaacacau  | 1500 |
| uggaguggau accgagauuc aagguuaguc ccaaccucuu cacuguccca auuaaggaag  | 1560 |
| caggcgaaga cugccaugcc ccaacauacc uaccugcgga gguggacggu gaugucaaac  | 1620 |
| ucaguuccaa ccuggugauu cuaccugguc aagaucucca auauguuuug gcaaccuacg  | 1680 |
| auaccuccag gguugagcau gcugugguuu auuacguuua cagcccaagc cgcucauuuu  | 1740 |
| cuuacuuuua uccuuuuagg uugccuauaa aggggguccc aaucgaacua caaguggaau  | 1800 |
| gcuucacaug ggaucaaaaa cucuggugcc gucacuucug ugugcuugcg gacucagaau  | 1860 |
| ccgguggacu uaucacucac ucugggaugg ugggcauggg agucagcugc acagcuaccc  | 1920 |
| gggaagaugg aaccaaucgc agauaaugau aauaggcugg agccucggug gccaagcuuc  | 1980 |
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| guucuguucg ucauguuucu gagcuugauc ggauugcugg caauugcagg cauuagacuu  | 180  |
| caucgggcag ccaucuacac cgcggagauc cauaaaagcc ucaguaccaa ucuggaugug  | 240  |
| acuaacucca ucgagcauca ggucaaggac gugcugacac cacucuuuaa aaucaucggg  | 300  |
| gaugaagugg gecugagaac accucagaga uucacugace uagugaaauu caucucggac  | 360  |
| 5 5 5 55 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5   |      |

420

aagauuaaau uccuuaaucc ggauagggag uacgacuuca gagaucucac uuggugcauc

| aacccgccag                 | agaggaucaa  | acuagauuau | gaucaauacu          | gugcagaugu | ggcugcugaa | 480  |
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| gagcucauga                 | augcauuggu  | gaacucaacu | cuacuggaga          | ccagaacaac | cacucaguuc | 540  |
| cuagcugucu                 | caaagggaaa  | cugcucaggg | cccacuacaa          | ucagagguca | auucucaaac | 600  |
| augucgcugu                 | ccuuguugga  | cuuguacuua | ggucgagguu          | acaauguguc | aucuauaguc | 660  |
| acuaugacau                 | cccagggaau  | guauggggga | accuaccuag          | uugaaaagcc | uaaucugaac | 720  |
| agcaaagggu                 | cagaguuguc  | acaacugagc | auguaccgag          | uguuugaagu | aggugugauc | 780  |
| agaaacccgg                 | guuugggggc  | uccgguguuc | cauaugacaa          | acuauuuuga | gcaaccaguc | 840  |
| aguaaugguc                 | ucggcaacug  | uaugguggcu | uugggggagc          | ucaaacucgc | agcccuuugu | 900  |
| cacggggacg                 | auucuaucau  | aauucccuau | cagggaucag          | ggaaaggugu | cagcuuccag | 960  |
| cucgucaagc                 | ugggugucug  | gaaaucccca | accgacaugc          | aauccugggu | ccccuuauca | 1020 |
| acggaugauc                 | cagugguaga  | caggcuuuac | cucucaucuc          | acagaggugu | caucgcugac | 1080 |
| aaucaagcaa                 | aaugggcugu  | cccgacaaca | cgaacagaug          | acaaguugcg | aauggagaca | 1140 |
| ugcuuccagc                 | aggcguguaa  | agguaaaauc | caagcacucu          | gcgagaaucc | cgagugggua | 1200 |
| ccauugaagg                 | auaacaggau  | uccuucauac | gggguccugu          | cuguugaucu | gagucugacg | 1260 |
| guugagcuua                 | aaaucaaaau  | ugcuucggga | uucgggccau          | ugaucacaca | cggcucaggg | 1320 |
| auggaccuau                 | acaaauccaa  | cugcaacaau | guguauuggc          | ugacuauucc | gccaaugaga | 1380 |
| aaucuagccu                 | uaggcguaau  | caacacauug | gaguggauac          | cgagauucaa | gguuaguccc | 1440 |
| aaccucuuca                 | cugucccaau  | uaaggaagca | ggcgaagacu          | gccaugcccc | aacauaccua | 1500 |
| ccugcggagg                 | uggacgguga  | ugucaaacuc | aguuccaacc          | uggugauucu | accuggucaa | 1560 |
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| uacguuuaca                 | gcccaagccg  | cucauuuucu | uacuuuuauc          | cuuuuagguu | gccuauaaag | 1680 |
| ggggucccaa                 | ucgaacuaca  | aguggaaugc | uucacauggg          | aucaaaaacu | cuggugccgu | 1740 |
| cacuucugug                 | ugcuugcgga  | cucagaaucc | gguggacuua          | ucacucacuc | ugggauggug | 1800 |
| ggcaugggag                 | ucagcugcac  | agcuacccgg | gaagauggaa          | ccaaucgcag | auaa       | 1854 |
| <220> FEATU<br><223> OTHER | TH: 2126<br>: RNA<br>NISM: Artif:<br>JRE:<br>R INFORMATIO | _          | nce<br>ic Polynucle | eotide     |            |      |
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| auuaacagag                 | aacaucuuau  | gauugacaga | cccuauguuc          | ugcuggcugu | ucuguucguc | 180  |
| auguuucuga                 | gcuugaucgg  | auugcuggca | auugcaggca          | uuagacuuca | ucgggcagcc | 240  |
| aucuacaccg                 | cggagaucca  | uaaaagccuc | aguaccaauc          | uggaugugac | uaacuccauc | 300  |
| gagcaucagg                 | ucaaggacgu  | gcugacacca | cucuuuaaaa          | ucaucgggga | ugaagugggc | 360  |
| cugagaacac                 | cucagagauu  | cacugaccua | gugaaauuca          | ucucggacaa | gauuaaauuc | 420  |
| cuuaauccgg                 | auagggagua  | cgacuucaga | gaucucacuu          | ggugcaucaa | cccgccagag | 480  |
| aggaucaaac                 | uagauuauga  | ucaauacugu | gcagaugugg          | cugcugaaga | gcucaugaau | 540  |
|                            |   |            |                     |            |            |      |

gcauugguga acucaacucu acuggagacc agaacaacca cucaguuccu agcugucuca

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| uuguuggacu  | uguacuuagg                        | ucgagguuac | aaugugucau | cuauagucac | uaugacaucc | 720  |  |
| cagggaaugu  | augggggaac                        | cuaccuaguu | gaaaagccua | aucugaacag | caaaggguca | 780  |  |
| gaguugucac  | aacugagcau                        | guaccgagug | uuugaaguag | gugugaucag | aaacccgggu | 840  |  |
| uugggggcuc  | cgguguucca                        | uaugacaaac | uauuuugagc | aaccagucag | uaauggucuc | 900  |  |
| ggcaacugua  | ugguggcuuu                        | gggggagcuc | aaacucgcag | cccuuuguca | cggggacgau | 960  |  |
| ucuaucauaa  | uucccuauca                        | gggaucaggg | aaagguguca | gcuuccagcu | cgucaagcug | 1020 |  |
| ggugucugga  | aauccccaac                        | cgacaugcaa | uccugggucc | ccuuaucaac | ggaugaucca | 1080 |  |
| gugguagaca  | ggcuuuaccu                        | cucaucucac | agagguguca | ucgcugacaa | ucaagcaaaa | 1140 |  |
| ugggcugucc  | cgacaacacg                        | aacagaugac | aaguugcgaa | uggagacaug | cuuccagcag | 1200 |  |
| gcguguaaag  | guaaaaucca                        | agcacucugc | gagaaucccg | aguggguacc | auugaaggau | 1260 |  |
| aacaggauuc  | cuucauacgg                        | gguccugucu | guugaucuga | gucugacggu | ugagcuuaaa | 1320 |  |
| aucaaaauug  | cuucgggauu                        | cgggccauug | aucacacacg | gcucagggau | ggaccuauac | 1380 |  |
| aaauccaacu  | gcaacaaugu                        | guauuggcug | acuauuccgc | caaugagaaa | ucuagccuua | 1440 |  |
| ggcguaauca  | acacauugga                        | guggauaccg | agauucaagg | uuagucccaa | ccucuucacu | 1500 |  |
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| guuuuggcaa  | ccuacgauac                        | cuccaggguu | gagcaugcug | ugguuuauua | cguuuacagc | 1680 |  |
| ccaagccgcu  | cauuuucuua                        | cuuuuauccu | uuuagguugc | cuauaaaggg | ggucccaauc | 1740 |  |
| gaacuacaag  | uggaaugcuu                        | cacaugggau | caaaaacucu | ggugccguca | cuucugugug | 1800 |  |
| cuugcggacu  | cagaauccgg                        | uggacuuauc | acucacucug | ggaugguggg | caugggaguc | 1860 |  |
| agcugcacag  | cuacccggga                        | agauggaacc | aaucgcagau | aaugauaaua | ggcuggagcc | 1920 |  |
| ucgguggcca  | agcuucuugc                        | cccuugggcc | uccccccagc | cccuccuccc | cuuccugcac | 1980 |  |
| ccguaccccc  | guggucuuug                        | aauaaagucu | gagugggcgg | caaaaaaaaa | aaaaaaaaa  | 2040 |  |
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| aaagaagagu  | aagaagaaau                        | auaagagcca | ccaugucacc | acaacgagac | cggauaaaug | 120  |  |
| ccuucuacaa  | agacaacccc                        | cauccuaagg | gaaguaggau | aguuauuaac | agagaacauc | 180  |  |
| uuaugauuga  | uagaccuuau                        | guuuugcugg | cuguucuauu | cgucauguuu | cugagcuuga | 240  |  |
| ucggguugcu  | agccauugca                        | ggcauuagac | uucaucgggc | agccaucuac | accgcagaga | 300  |  |
| uccauaaaag  | ccucagcacc                        | aaucuggaug | uaacuaacuc | aaucgagcau | cagguuaagg | 360  |  |
| acgugcugac  | accacucuuc                        | aagaucaucg | gugaugaagu | gggcuugagg | acaccucaga | 420  |  |
| gauucacuga  | ccuagugaag                        | uucaucucug | acaagauuaa | auuccuuaau | ccggacaggg | 480  |  |
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540

aauacgacuu cagagaucuc acuuggugua ucaacccgcc agagagaauc aaauuggauu

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| cucuacugga                 | gaccagggca  | accaaucagu                   | uccuagcugu | cucaaaggga | aacugcucag | 660  |
| ggcccacuac                 | aaucagaggc  | caauucucaa                   | acaugucgcu | gucccuguug | gacuuguauu | 720  |
| uaagucgagg                 | uuacaaugug  | ucaucuauag                   | ucacuaugac | aucccaggga | auguacgggg | 780  |
| gaacuuaccu                 | aguggaaaag  | ccuaaucuga                   | gcagcaaagg | gucagaguug | ucacaacuga | 840  |
| gcaugcaccg                 | aguguuugaa  | guagguguua                   | ucagaaaucc | ggguuugggg | gcuccgguau | 900  |
| uccauaugac                 | aaacuaucuu  | gagcaaccag                   | ucaguaauga | uuucagcaac | ugcauggugg | 960  |
| cuuuggggga                 | gcucaaguuc  | gcagcccucu                   | gucacaggga | agauucuauc | acaauucccu | 1020 |
| aucagggauc                 | agggaaaggu  | gucagcuucc                   | agcuugucaa | gcuagguguc | uggaaauccc | 1080 |
| caaccgacau                 | gcaauccugg  | gucccccuau                   | caacggauga | uccagugaua | gacaggcuuu | 1140 |
| accucucauc                 | ucacagaggc  | guuaucgcug                   | acaaucaagc | aaaaugggcu | gucccgacaa | 1200 |
| cacggacaga                 | ugacaaguug  | cgaauggaga                   | caugcuucca | gcaggcgugu | aaggguaaaa | 1260 |
| uccaagcacu                 | uugcgagaau  | cccgagugga                   | caccauugaa | ggauaacagg | auuccuucau | 1320 |
| acggggucuu                 | gucuguugau  | cugagucuga                   | caguugagcu | uaaaaucaaa | auuguuucag | 1380 |
| gauucgggcc                 | auugaucaca  | cacgguucag                   | ggauggaccu | auacaaaucc | aaccacaaca | 1440 |
| auauguauug                 | gcugacuauc  | ccgccaauga                   | agaaccuggc | cuuaggugua | aucaacacau | 1500 |
| uggaguggau                 | accgagauuc  | aagguuaguc                   | ccaaccucuu | cacuguucca | auuaaggaag | 1560 |
| caggcgagga                 | cugccaugcc  | ccaacauacc                   | uaccugcgga | gguggauggu | gaugucaaac | 1620 |
| ucaguuccaa                 | ucuggugauu  | cuaccugguc                   | aagaucucca | auauguucug | gcaaccuacg | 1680 |
| auacuuccag                 | aguugaacau  | gcuguaguuu                   | auuacguuua | cagcccaagc | cgcucauuuu | 1740 |
| cuuacuuuua                 | uccuuuuagg  | uugccuguaa                   | gggggguccc | cauugaauua | caaguggaau | 1800 |
| gcuucacaug                 | ggaccaaaaa  | cucuggugcc                   | gucacuucug | ugugcuugcg | gacucagaau | 1860 |
| cugguggaca                 | uaucacucac  | ucugggaugg                   | ugggcauggg | agucagcugc | acagccacuc | 1920 |
| gggaagaugg                 | aaccagccgc  | agauagugau                   | aauaggcugg | agccucggug | gccaagcuuc | 1980 |
| uugccccuug                 | ggccuccccc  | cageceeuce                   | uccccuuccu | gcacccguac | ccccgugguc | 2040 |
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| <220> FEATU<br><223> OTHER | TH: 1854<br>: RNA<br>NISM: Artif:<br>URE:<br>R INFORMATIO | icial Sequer<br>DN: Synthet: |            | eotide     |            |      |
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| aguaggauag                 | uuauuaacag  | agaacaucuu                   | augauugaua | gaccuuaugu | uuugcuggcu | 120  |
| guucuauucg                 | ucauguuucu  | gagcuugauc                   | ggguugcuag | ccauugcagg | cauuagacuu | 180  |
| caucgggcag                 | ccaucuacac  | cgcagagauc                   | cauaaaagcc | ucagcaccaa | ucuggaugua | 240  |
| acuaacucaa                 | ucgagcauca  | gguuaaggac                   | gugcugacac | cacucuucaa | gaucaucggu | 300  |
| gaugaagugg                 | gcuugaggac  | accucagaga                   | uucacugacc | uagugaaguu | caucucugac | 360  |
| aagauuaaau                 | uccuuaaucc  | ggacagggaa                   | uacgacuuca | gagaucucac | uugguguauc | 420  |

aacccgccag agagaaucaa auuggauuau gaucaauacu gugcagaugu ggcugcugaa

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|             |                                   |            |            | COIICII    | iaca       |      |  |
|-------------|-----------------------------------|------------|------------|------------|------------|------|--|
| gaacucauga  | augcauuggu                        | gaacucaacu | cuacuggaga | ccagggcaac | caaucaguuc | 540  |  |
| cuagcugucu  | caaagggaaa                        | cugcucaggg | cccacuacaa | ucagaggcca | auucucaaac | 600  |  |
| augucgcugu  | cccuguugga                        | cuuguauuua | agucgagguu | acaauguguc | aucuauaguc | 660  |  |
| acuaugacau  | cccagggaau                        | guacggggga | acuuaccuag | uggaaaagcc | uaaucugagc | 720  |  |
| agcaaagggu  | cagaguuguc                        | acaacugagc | augcaccgag | uguuugaagu | agguguuauc | 780  |  |
| agaaauccgg  | guuugggggc                        | uccgguauuc | cauaugacaa | acuaucuuga | gcaaccaguc | 840  |  |
| aguaaugauu  | ucagcaacug                        | caugguggcu | uugggggagc | ucaaguucgc | ageceueugu | 900  |  |
| cacagggaag  | auucuaucac                        | aauucccuau | cagggaucag | ggaaaggugu | cagcuuccag | 960  |  |
| cuugucaagc  | uaggugucug                        | gaaaucccca | accgacaugc | aauccugggu | ccccuauca  | 1020 |  |
| acggaugauc  | cagugauaga                        | caggcuuuac | cucucaucuc | acagaggcgu | uaucgcugac | 1080 |  |
| aaucaagcaa  | aaugggcugu                        | cccgacaaca | cggacagaug | acaaguugcg | aauggagaca | 1140 |  |
| ugcuuccagc  | aggcguguaa                        | ggguaaaauc | caagcacuuu | gcgagaaucc | cgaguggaca | 1200 |  |
| ccauugaagg  | auaacaggau                        | uccuucauac | ggggucuugu | cuguugaucu | gagucugaca | 1260 |  |
| guugagcuua  | aaaucaaaau                        | uguuucagga | uucgggccau | ugaucacaca | cgguucaggg | 1320 |  |
| auggaccuau  | acaaauccaa                        | ccacaacaau | auguauuggc | ugacuauccc | gccaaugaag | 1380 |  |
| aaccuggccu  | uagguguaau                        | caacacauug | gaguggauac | cgagauucaa | gguuaguccc | 1440 |  |
| aaccucuuca  | cuguuccaau                        | uaaggaagca | ggcgaggacu | gccaugcccc | aacauaccua | 1500 |  |
| ccugcggagg  | uggaugguga                        | ugucaaacuc | aguuccaauc | uggugauucu | accuggucaa | 1560 |  |
| gaucuccaau  | auguucuggc                        | aaccuacgau | acuuccagag | uugaacaugc | uguaguuuau | 1620 |  |
| uacguuuaca  | gcccaagccg                        | cucauuuucu | uacuuuuauc | cuuuuagguu | gccuguaagg | 1680 |  |
| ggggucccca  | uugaauuaca                        | aguggaaugc | uucacauggg | accaaaaacu | cuggugccgu | 1740 |  |
| cacuucugug  | ugcuugcgga                        | cucagaaucu | gguggacaua | ucacucacuc | ugggauggug | 1800 |  |
| ggcaugggag  | ucagcugcac                        | agccacucgg | gaagauggaa | ccagccgcag | auag       | 1854 |  |
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| <400> SEQUI | ENCE: 80                          |            |            |            |            |      |  |
| ggggaaauaa  | gagagaaaag                        | aagaguaaga | agaaauauaa | gagccaccau | gucaccacaa | 60   |  |
| cgagaccgga  | uaaaugccuu                        | cuacaaagac | aacccccauc | cuaagggaag | uaggauaguu | 120  |  |
| auuaacagag  | aacaucuuau                        | gauugauaga | ccuuauguuu | ugcuggcugu | ucuauucguc | 180  |  |
| auguuucuga  | gcuugaucgg                        | guugcuagcc | auugcaggca | uuagacuuca | ucgggcagcc | 240  |  |
| aucuacaccg  | cagagaucca                        | uaaaagccuc | agcaccaauc | uggauguaac | uaacucaauc | 300  |  |
| gagcaucagg  | uuaaggacgu                        | gcugacacca | cucuucaaga | ucaucgguga | ugaagugggc | 360  |  |
| uugaggacac  | cucagagauu                        | cacugaccua | gugaaguuca | ucucugacaa | gauuaaauuc | 420  |  |
| cuuaauccgg  | acagggaaua                        | cgacuucaga | gaucucacuu | gguguaucaa | cccgccagag | 480  |  |
| agaaucaaau  | uggauuauga                        | ucaauacugu | gcagaugugg | cugcugaaga | acucaugaau | 540  |  |
| gcauugguga  | acucaacucu                        | acuggagacc | agggcaacca | aucaguuccu | agcugucuca | 600  |  |
|             | gcucagggcc                        |            |            |            |            | 660  |  |
| 5555        | 5 5555                            |            | 5 55 2 244 |            | 555        |      |  |

720

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| cagggaaugu  | acgggggaac                        | uuaccuagug | gaaaagccua | aucugagcag | caaaggguca | 780  |
|-------------|-----------------------------------|------------|------------|------------|------------|------|
| gaguugucac  | aacugagcau                        | gcaccgagug | uuugaaguag | guguuaucag | aaauccgggu | 840  |
| uugggggcuc  | cgguauucca                        | uaugacaaac | uaucuugagc | aaccagucag | uaaugauuuc | 900  |
| agcaacugca  | ugguggcuuu                        | gggggagcuc | aaguucgcag | cccucuguca | cagggaagau | 960  |
| ucuaucacaa  | uucccuauca                        | gggaucaggg | aaagguguca | gcuuccagcu | ugucaagcua | 1020 |
| ggugucugga  | aauccccaac                        | cgacaugcaa | uccugggucc | cccuaucaac | ggaugaucca | 1080 |
| gugauagaca  | ggcuuuaccu                        | cucaucucac | agaggcguua | ucgcugacaa | ucaagcaaaa | 1140 |
| ugggcugucc  | cgacaacacg                        | gacagaugac | aaguugcgaa | uggagacaug | cuuccagcag | 1200 |
| gcguguaagg  | guaaaaucca                        | agcacuuugc | gagaaucccg | aguggacacc | auugaaggau | 1260 |
| aacaggauuc  | cuucauacgg                        | ggucuugucu | guugaucuga | gucugacagu | ugagcuuaaa | 1320 |
| aucaaaauug  | uuucaggauu                        | cgggccauug | aucacacacg | guucagggau | ggaccuauac | 1380 |
| aaauccaacc  | acaacaauau                        | guauuggcug | acuaucccgc | caaugaagaa | ccuggccuua | 1440 |
| gguguaauca  | acacauugga                        | guggauaccg | agauucaagg | uuagucccaa | ccucuucacu | 1500 |
| guuccaauua  | aggaagcagg                        | cgaggacugc | caugccccaa | cauaccuacc | ugcggaggug | 1560 |
| gauggugaug  | ucaaacucag                        | uuccaaucug | gugauucuac | cuggucaaga | ucuccaauau | 1620 |
| guucuggcaa  | ccuacgauac                        | uuccagaguu | gaacaugcug | uaguuuauua | cguuuacagc | 1680 |
| ccaagccgcu  | cauuuucuua                        | cuuuuauccu | uuuagguugc | cuguaagggg | gguccccauu | 1740 |
| gaauuacaag  | uggaaugcuu                        | cacaugggac | caaaaacucu | ggugccguca | cuucugugug | 1800 |
| cuugcggacu  | cagaaucugg                        | uggacauauc | acucacucug | ggaugguggg | caugggaguc | 1860 |
| agcugcacag  | ccacucggga                        | agauggaacc | agccgcagau | agugauaaua | ggcuggagcc | 1920 |
| ucgguggcca  | agcuucuugc                        | cccuugggcc | uccccccagc | cccuccuccc | cuuccugcac | 1980 |
| ccguaccccc  | guggucuuug                        | aauaaagucu | gagugggcgg | caaaaaaaaa | aaaaaaaaa  | 2040 |
| aaaaaaaaa   | aaaaaaaaa                         | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | aaaaaaaaa  | 2100 |
| aaaaaaaaa   | aaaaaaaaa                         | aucuag     |            |            |            | 2126 |
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| <400> SEQUI | ENCE: 81                          |            |            |            |            |      |
| ucaagcuuuu  | ggacccucgu                        | acagaagcua | auacgacuca | cuauagggaa | auaagagaga | 60   |
| aaagaagagu  | aagaagaaau                        | auaagagcca | ccauggcaca | agucauuaau | acaaacagcc | 120  |
| ugucgcuguu  | gacccagaau                        | aaccugaaca | aaucccaguc | cgcacugggc | acugcuaucg | 180  |
| agcguuuguc  | uuccggucug                        | cguaucaaca | gcgcgaaaga | cgaugcggca | ggacaggcga | 240  |
| uugcuaaccg  | uuuuaccgcg                        | aacaucaaag | gucugacuca | ggcuucccgu | aacgcuaacg | 300  |
| acgguaucuc  | cauugcgcag                        | accacugaag | gcgcgcugaa | cgaaaucaac | aacaaccugc | 360  |
| agcgugugcg  | ugaacuggcg                        | guucagucug | cgaaugguac | uaacucccag | ucugaccucg | 420  |
| acuccaucca  | ggcugaaauc                        | acccagcgcc | ugaacgaaau | cgaccgugua | uccggccaga | 480  |
| cucaguucaa  | cggcgugaaa                        | guccuggcgc | aggacaacac | ccugaccauc | cagguuggug | 540  |

ccaacgacgg ugaaacuauc gauauugauu uaaaagaaau cagcucuaaa acacugggac

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|---|---|
| uugauaagcu uaauguccaa gaugccuaca ccccgaaaga aacugcugua accguugaua   | 660   |
| aaacuaccua uaaaaauggu acagauccua uuacagccca gagcaauacu gauauccaaa   | 720   |
| cugcaauugg cgguggugca acggggguua cuggggcuga uaucaaauuu aaagaugguc   | 780   |
| aauacuauuu agauguuaaa ggcggugcuu cugcuggugu uuauaaagcc acuuaugaug   | 840   |
| aaacuacaaa gaaaguuaau auugauacga cugauaaaac uccguuggca acugcggaag   | 900   |
| cuacagcuau ucggggaacg gccacuauaa cccacaacca aauugcugaa guaacaaaag   | 960   |
| aggguguuga uacgaccaca guugcggcuc aacuugcugc agcagggguu acuggcgccg   | 1020  |
| auaaggacaa uacuagccuu guaaaacuau cguuugagga uaaaaacggu aagguuauug   | 1080  |
| augguggcua ugcagugaaa augggcgacg auuucuaugc cgcuacauau gaugagaaaa   | 1140  |
| caggugcaau uacugcuaaa accacuacuu auacagaugg uacuggcguu gcucaaacug   | 1200  |
| gagcugugaa auuugguggc gcaaauggua aaucugaagu uguuacugcu accgauggua   | 1260  |
| agacuuacuu agcaagcgac cuugacaaac auaacuucag aacaggcggu gagcuuaaag   | 1320  |
| agguuaauac agauaagacu gaaaacccac ugcagaaaau ugaugcugcc uuggcacagg   | 1380  |
| uugauacacu ucguucugac cugggugcgg uucagaaccg uuucaacucc gcuaucacca   | 1440  |
| accugggcaa uaccguaaau aaccugucuu cugcccguag ccguaucgaa gauuccgacu   | 1500  |
| acgcaaccga agucuccaac augucucgcg cgcagauucu gcagcaggcc gguaccuccg   | 1560  |
| uucuggcgca ggcgaaccag guuccgcaaa acguccucuc uuuacugcgu ugauaauagg   | 1620  |
| cuggagecue gguggecaug cuucuugeee cuugggecue eececageee cuccueeecu   | 1680  |
| uccugcacce guaccecegu ggucuuugaa uaaagucuga gugggegge   | 1729  |
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| <pre>&lt;211&gt; LENGTH: 1518 &lt;212&gt; TYPE: RNA &lt;213&gt; ORGANISM: Artificial Sequence &lt;220&gt; FEATURE: &lt;223&gt; OTHER INFORMATION: Synthetic Polynucleotide &lt;400&gt; SEQUENCE: 82 auggcacaag ucauuaauac aaacagccug ucgcuguuga cccagaauaa ccugaacaaa ucccaguccg cacugggcac ugcuaucgag cguuugucuu ccggucugcg uaucaacagc gcgaaaagacg augcggcagg acaggcgauu gcuaaccguu uuaccgcgaa caucaaaaggu cugacucagg cuucccguaa cgcuaacgac gguaucucca uugcgcagac cacugaaggc gcgcugaacg aaaucaacaa caaccugcag cgugugcgug aacuggcggu ucagucugcg aaugguacua acucccaguc ugaccucgac uccauccagg cugaaaucac ccagcgccug aacgaaaucg accguguauc cggccagacu caguucaacg gcgugaaagu ccuggcgcag gacaacaccc ugaccaucca gguuggugca aacgacggug aaacuaucga uauugauuua aaagaaauca gcucuaaaac acugggacuu gauaagcuua auguccaaga ugccuacacc ccgaaagaaa cugcuguaac cguugauaaa acuaccuaua aaaaugguac agauccuauu acagcccaga gcaauacuga uauccaaacu gcaauuggcg guggugcaac ggggguuacu ggggcugaua ucaaauuuaa agauggucaa uacuauuuag auguuaaagg cggugcuucu</pre>   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660<br>720 |
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cuugcugcag cagggguuac uggcgccgau aaggacaaua cuagccuugu aaaacuaucg

| uuugaggaua | aaaacgguaa  | gguuauugau | gguggcuaug          | cagugaaaau | gggcgacgau | 1020 |
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| acagauggua | cuggcguugc  | ucaaacugga | gcugugaaau          | uugguggcgc | aaaugguaaa | 1140 |
| ucugaaguug | uuacugcuac  | cgaugguaag | acuuacuuag          | caagcgaccu | ugacaaacau | 1200 |
| aacuucagaa | caggcgguga  | gcuuaaagag | guuaauacag          | auaagacuga | aaacccacug | 1260 |
| cagaaaauug | augcugccuu  | ggcacagguu | gauacacuuc          | guucugaccu | gggugcgguu | 1320 |
| cagaaccguu | ucaacuccgc  | uaucaccaac | cugggcaaua          | ccguaaauaa | ccugucuucu | 1380 |
| gcccguagcc | guaucgaaga  | uuccgacuac | gcaaccgaag          | ucuccaacau | gucucgcgcg | 1440 |
| cagauucugc | agcaggccgg  | uaccuccguu | cuggcgcagg          | cgaaccaggu | uccgcaaaac | 1500 |
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| auuaauacaa | acagccuguc  | gcuguugacc | cagaauaacc          | ugaacaaauc | ccaguccgca | 120  |
| cugggcacug | cuaucgagcg  | uuugucuucc | ggucugcgua          | ucaacagcgc | gaaagacgau | 180  |
| geggeaggae | aggcgauugc  | uaaccguuuu | accgcgaaca          | ucaaaggucu | gacucaggcu | 240  |
| ucccguaacg | cuaacgacgg  | uaucuccauu | gcgcagacca          | cugaaggcgc | gcugaacgaa | 300  |
| aucaacaaca | accugcagcg  | ugugcgugaa | cuggcgguuc          | agucugcgaa | ugguacuaac | 360  |
| ucccagucug | accucgacuc  | cauccaggcu | gaaaucaccc          | agcgccugaa | cgaaaucgac | 420  |
| cguguauccg | gccagacuca  | guucaacggc | gugaaagucc          | uggcgcagga | caacacccug | 480  |
| accauccagg | uuggugccaa  | cgacggugaa | acuaucgaua          | uugauuuaaa | agaaaucagc | 540  |
| ucuaaaacac | ugggacuuga  | uaagcuuaau | guccaagaug          | ccuacacccc | gaaagaaacu | 600  |
| gcuguaaccg | uugauaaaac  | uaccuauaaa | aaugguacag          | auccuauuac | agcccagagc | 660  |
| aauacugaua | uccaaacugc  | aauuggcggu | ggugcaacgg          | ggguuacugg | ggcugauauc | 720  |
| aaauuuaaag | auggucaaua  | cuauuuagau | guuaaaggcg          | gugcuucugc | ugguguuuau | 780  |
| aaagccacuu | augaugaaac  | uacaaagaaa | guuaauauug          | auacgacuga | uaaaacuccg | 840  |
| uuggcaacug | cggaagcuac  | agcuauucgg | ggaacggcca          | cuauaaccca | caaccaaauu | 900  |
| gcugaaguaa | caaaagaggg  | uguugauacg | accacaguug          | cggcucaacu | ugcugcagca | 960  |
| gggguuacug | gcgccgauaa  | ggacaauacu | agccuuguaa          | aacuaucguu | ugaggauaaa | 1020 |
| aacgguaagg | uuauugaugg  | uggcuaugca | gugaaaaugg          | gcgacgauuu | cuaugccgcu | 1080 |
| acauaugaug | agaaaacagg  | ugcaauuacu | gcuaaaacca          | cuacuuauac | agaugguacu | 1140 |
| ggcguugcuc | aaacuggagc  | ugugaaauuu | gguggcgcaa          | augguaaauc | ugaaguuguu | 1200 |
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|            |   |            |                     |            |            |      |

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                                                               1560
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Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe
                         40
Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro
Ser Leu Ile Lys Thr Glu Leu Asp Leu Thr Lys Ser Ala Leu Arg Glu
Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu
Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val
                   105
Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Cys Lys Thr Ile
Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr
Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Phe
Ala Val Arg Glu Leu Lys Asp Phe Val Ser Lys Asn Leu Thr Arg Ala
Leu Asn Lys Asn Lys Cys Asp Ile Asp Asp Leu Lys Met Ala Val Ser
                             185
Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser
                         200
Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp
Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln
Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe
```

|                      |   |                                  |                             | 245        |            |            |            |            | 250          |            |            |            |            | 255        |            |
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| Gly                  | Ile   | Leu                              | Cys<br>260                  | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser          | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu                  | Pro   | Ile<br>275                       | Phe                         | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro          | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala                  | Pro<br>290  | Ser                              | CÀa                         | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn          | Tyr        | Ala<br>300 | CAa        | Leu        | Leu        | Arg        |
| Glu<br>305           | Asp   | Gln                              | Gly                         | Trp        | Tyr<br>310 | CÀa        | Gln        | Asn        | Ala          | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro                  | Asn   | Glu                              | ГÀа                         | Asp<br>325 | CÀa        | Glu        | Thr        | Arg        | Gly<br>330   | Asp        | His        | Val        | Phe        | 335        | Asp        |
| Thr                  | Ala   | Ala                              | Gly<br>340                  | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln          | Ser        | Lys        | Glu        | 350<br>250 | Asn        | Ile        |
| Asn                  | Ile   | Ser<br>355                       | Thr                         | Thr        | Asn        | Tyr        | Pro<br>360 | Cys        | Lys          | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro                  | Ile<br>370  | Ser                              | Met                         | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu          | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385           | Lys   | Gly                              | Val                         | Ser        | 390<br>CAa | Ser        | Ile        | Gly        | Ser          | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| ГÀа                  | Gln   | Leu                              | Asn                         | Lys<br>405 | Gly        | CAa        | Ser        | Tyr        | Ile<br>410   | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr                  | Val   | Thr                              | Ile<br>420                  | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln          | Leu        | Ser        | ГÀа        | Val<br>430 | Glu        | Gly        |
| Glu                  | Gln   | His<br>435                       | Val                         | Ile        | ГÀа        | Gly        | Arg<br>440 | Pro        | Val          | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile                  | Lys<br>450  | Phe                              | Pro                         | Glu        | Asp        | Gln<br>455 | Phe        | Asn        | Val          | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465           | Asn   | Ile                              | Glu                         | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val          | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu                  | Ser   | Ser                              | Ala                         | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490   | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu                  | Ile   | Ala                              | Val<br>500                  | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile          | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile                  | Ile   | Lys<br>515                       | Lys                         | Thr        | rys        | Lys        | Pro<br>520 | Thr        | Gly          | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly                  | Val<br>530  | Thr                              | Asn                         | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His          | Asn        |            |            |            |            |            |
| <213<br><213<br><223 | 0 > SI<br>1 > LI<br>2 > TY<br>3 > OI<br>0 > FI<br>3 > O | ENGTI<br>(PE :<br>RGAN)<br>EATUI | H: 53<br>PRT<br>ISM:<br>RE: | 39<br>Art: |            |            | -          |            | oly <u>r</u> | pept:      | ide        |            |            |            |            |
| < 400                | D> SI   | EQUEI                            | ICE :                       | 86         |            |            |            |            |              |            |            |            |            |            |            |
| Met<br>1             | Ser   | Trp                              | Lys                         | Val<br>5   | Val        | Ile        | Ile        | Phe        | Ser<br>10    | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                  | Gly   | Leu                              | Lys<br>20                   | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu          | Ser        | Cys        | Ser        | Thr        | Ile        | Thr        |
| Glu                  | Gly   | Tyr<br>35                        | Leu                         | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly          | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                  | Leu<br>50   | Glu                              | Val                         | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu          | Thr        | Cys        | Ser        | Asp        | Gly        | Pro        |
| Ser                  | Leu   | Ile                              | ГЛа                         | Thr        | Glu        | Leu        | Asp        | Leu        | Thr          | rys        | Ser        | Ala        | Leu        | Arg        | Glu        |

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|------------|---|
|------------|---|

| _          |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| 65         |            |            |            |            | 70         |            |            |            |            | 75         |            |            |            |            | 80         |
| Leu        | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Сув<br>125 | Lys        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | ràa        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | ГЛа        | CÀa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Cys<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | CÀa        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | СЛа        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | Càa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | CÀa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | 335<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | 350<br>Cys | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390<br>CAa | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | ГЛа        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | His        | Gln<br>455 | Trp        | His        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |

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Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser 520 Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 87 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 87 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Leu Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val 105 Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile 120 Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr 155 Ala Val Arg Glu Leu Lys Asp Phe Val Ser Lys Asn Leu Thr Arg Ala Ile Asn Lys Asn Lys Cys Asp Ile Pro Asp Leu Lys Met Ala Val Ser Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln 265 Leu Pro Ile Phe Gly Val Ile Asp Thr Pro Cys Trp Ile Val Lys Ala Ala Pro Ser Cys Ser Glu Lys Lys Gly Asn Tyr Ala Cys Leu Leu Arg 295 300 Glu Asp Gln Gly Trp Tyr Cys Gln Asn Ala Gly Ser Thr Val Tyr Tyr 310 315

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Pro Asn Glu Lys Asp Cys Glu Thr Arg Gly Asp His Val Phe Cys Asp Thr Ala Ala Gly Ile Asn Val Ala Glu Gln Ser Lys Glu Cys Asn Ile Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro 440 Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe 455 Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile 490 Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser 520 Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn 530 <210> SEQ ID NO 88 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 88 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Leu Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile 120 125 Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr 135

| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|            | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 |            | Asn        | Leu        | Thr        | Arg<br>175 |            |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | Cys        | Asp        | Ile        | Pro<br>185 | Asp        | Leu        | Гуз        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | Cys        | Ser        | Glu        | Lув<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CÀa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | ГÀа        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | 335<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | ГÀз        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | СЛа        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390        | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | ГÀа        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | Asn        | Gln<br>455 | Phe        | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu<br>485 | ГÀа        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu        | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile        | Ile        | Lys<br>515 | Lys        | Thr        | ГÀа        | ГЛа        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly        | Val<br>530 | Thr        | Asn        | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |

| <213<br><220 | )> FI            | RGANI<br>EATUF | ISM:<br>RE: |            |            |            | Seque      |            |            |            |            |            |            |            |            |
|--------------|------------------|----------------|-------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|              | 3 > 01<br>0 > SI |                |             |            | rion       | : Syı      | nthet      | ic I       | Polyg      | pept:      | ide        |            |            |            |            |
|              |                  |                |             |            | Val        | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His          | Gly              | Leu            | Lys<br>20   | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | СЛа        | Ser        | Thr        | Ile        | Thr        |
| Glu          | Gly              | Tyr<br>35      | Leu         | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr          | Leu<br>50        | Glu            | Val         | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | 60<br>CÀa  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65    | Leu              | Ile            | Lys         | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Leu        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu          | Lys              | Thr            | Val         | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn          | Pro              | Gly            | Ser<br>100  | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala          | Ala              | Ala<br>115     | Ala         | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГÀа        | Thr        | Ile        |
| Arg          | Leu<br>130       | Glu            | Ser         | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | ГÀа        | Lys        | Thr        |
| Asn<br>145   | Glu              | Ala            | Val         | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala          | Val              | Arg            | Glu         | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Leu<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile          | Asn              | Lys            | Asn<br>180  | ГÀв        | СЛа        | Asp        | Ile        | Pro<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe          | Ser              | Gln<br>195     | Phe         | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp          | Asn<br>210       | Ala            | Gly         | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225   | Glu              | Leu            | Ala         | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile          | Lys              | Leu            | Met         | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly          | Ile              | Leu            | Ile<br>260  | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu          | Pro              | Ile<br>275     | Phe         | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala          | Pro<br>290       | Ser            | Cys         | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Сув        | Leu        | Leu        | Arg        |
| Glu<br>305   | Asp              | Gln            | Gly         | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro          | Asn              | Glu            | ГЛа         | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | 335<br>Cya | Asp        |
| Thr          | Ala              | Ala            | Gly<br>340  | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn          | Ile              | Ser<br>355     | Thr         | Thr        | Asn        | Tyr        | Pro<br>360 | СЛа        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro          | Ile<br>370       | Ser            | Met         | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Сув        |

| Tyr<br>385                   | Lys                              | Gly        | Val                         | Ser        | 390<br>Cys | Ser        | Ile            | Gly        | Ser          | Asn<br>395     | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
|------------------------------|----------------------------------|------------|-----------------------------|------------|------------|------------|----------------|------------|--------------|----------------|------------|------------|------------|------------|------------|
| Lys                          | Gln                              | Leu        | Asn                         | Lys<br>405 | Gly        | Cys        | Ser            | Tyr        | Ile<br>410   | Thr            | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr                          | Val                              | Thr        | Ile<br>420                  | Asp        | Asn        | Thr        | Val            | Tyr<br>425 | Gln          | Leu            | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu                          | Gln                              | His<br>435 | Val                         | Ile        | Lys        | Gly        | Arg<br>440     | Pro        | Val          | Ser            | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile                          | Lys<br>450                       | Phe        | Pro                         | Glu        | Asp        | Gln<br>455 | Phe            | Gln        | Val          | Ala            | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465                   | Asn                              | Ile        | Glu                         | Asn        | Ser<br>470 | Gln        | Ala            | Leu        | Val          | Asp<br>475     | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu                          | Ser                              | Ser        | Ala                         | Glu<br>485 | Lys        | Gly        | Asn            | Thr        | Gly<br>490   | Phe            | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu                          | Ile                              | Ala        | Val<br>500                  | Leu        | Gly        | Ser        | Ser            | Met<br>505 | Ile          | Leu            | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile                          | Ile                              | Lys<br>515 | Lys                         | Thr        | Lys        | Lys        | Pro<br>520     | Thr        | Gly          | Ala            | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly                          | Val<br>530                       | Thr        | Asn                         | Asn        | Gly        | Phe<br>535 | Ile            | Pro        | His          | Asn            |            |            |            |            |            |
| <211<br><212<br><213<br><220 | L> LE<br>2> TY<br>3> OF<br>0> FE | EATUI      | H: 53<br>PRT<br>ISM:<br>RE: | 39<br>Art: |            |            | Seque<br>nthet |            | oly <u>r</u> | p <b>e</b> pt: | ide        |            |            |            |            |
| < 400                        | )> SI                            | EQUEI      | ICE :                       | 90         |            |            |                |            |              |                |            |            |            |            |            |
| Met<br>1                     | Ser                              | Trp        | Lys                         | Val<br>5   | Val        | Ile        | Ile            | Phe        | Ser<br>10    | Leu            | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                          | Gly                              | Leu        | Lys<br>20                   | Glu        | Ser        | Tyr        | Leu            | Glu<br>25  | Glu          | Ser            | Càa        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly                              | Tyr<br>35  | Leu                         | Ser        | Val        | Leu        | Arg<br>40      | Thr        | Gly          | Trp            | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50                        | Glu        | Val                         | Gly        | Asp        | Val<br>55  | Glu            | Asn        | Leu          | Thr            | 60<br>Cys  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu                              | Ile        | Lys                         | Thr        | Glu<br>70  | Leu        | Asp            | Leu        | Leu          | Lys<br>75      | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | Lys                              | Thr        | Val                         | Ser<br>85  | Ala        | Asp        | Gln            | Leu        | Ala<br>90    | Arg            | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn                          | Pro                              | Gly        | Ser<br>100                  | Gly        | Ser        | Phe        | Val            | Leu<br>105 | Gly          | Ala            | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala                          | Ala                              | Ala<br>115 | Ala                         | Ala        | Val        | Thr        | Ala<br>120     | Gly        | Val          | Ala            | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg                          | Leu<br>130                       | Glu        | Ser                         | Glu        | Val        | Thr<br>135 | Ala            | Ile        | Asn          | Asn            | Ala<br>140 | Leu        | ГÀв        | ГÀв        | Thr        |
| Asn<br>145                   | Glu                              | Ala        | Val                         | Ser        | Thr<br>150 | Leu        | Gly            | Asn        | Gly          | Val<br>155     | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala                          | Val                              | Arg        | Glu                         | Leu<br>165 | Lys        | Asp        | Phe            | Val        | Leu<br>170   | Lys            | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile                          |                                  |            | _                           |            |            | 7          | Tle            | Dro        | 7 an         | T.011          | Lve        | Met        | Δla        | 77-7       | Car        |
|                              | Asn                              | Lys        | Asn<br>180                  | Lys        | Cys        | Asp        | 110            | 185        | Asp          | пец            | БуБ        |            | 190        | vai        | Del        |

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Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp 215 Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe 250 Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln Leu Pro Ile Phe Gly Val Ile Asp Thr Pro Cys Trp Ile Val Lys Ala Ala Pro Ser Cys Ser Glu Lys Lys Gly Asn Tyr Ala Cys Leu Leu Arg Glu Asp Gln Gly Trp Tyr Cys Gln Asn Ala Gly Ser Thr Val Tyr Tyr Pro Asn Glu Lys Asp Cys Glu Thr Arg Gly Asp His Val Phe Cys Asp 325 330 335 Thr Ala Ala Gly Ile Asn Val Ala Glu Gln Ser Lys Glu Cys Asn Ile Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys 375 Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile 390 395 Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly 425 Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro 440 Ile Lys Phe Pro Glu Asn Gln Phe Gln Val Ala Leu Asp Gln Val Phe 455 Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile 490 Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 91 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 91 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln 10 His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr 25

| Glu        | Gly        | Tyr<br>35  | Leu        | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Thr        | Leu<br>50  | Pro        | Val        | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | Cys        | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Leu        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu        | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | ГЛа        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | ГÀа        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | ГÀа        | CAa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | ГÀв        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | ГÀа        | Ala        |
| Ala        | Pro<br>290 | Ser        | Càa        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | СЛа        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | СЛа        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        |            | Asp<br>325 | CAa        | Glu        | Thr        |            | Gly<br>330 |            | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | ГÀЗ        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Cys        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | Cys<br>390 | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | Сув        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | ГЛа        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | ГÀа        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys        | Phe        | Pro        | Glu        | Asp        | Gln        | Phe        | Gln        | Val        | Ala        | Leu        | Asp        | Gln        | Val        | Phe        |

|              | 450                     |              |              |            |            | 455        |            |            |            |            | 460        |            |            |            |            |
|--------------|-------------------------|--------------|--------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Glu<br>465   | Asn                     | Ile          | Glu          | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu          | Ser                     | Ser          | Ala          | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu          | Ile                     | Ala          | Val<br>500   | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile          | Ile                     | Lys<br>515   | Lys          | Thr        | Lys        | Lys        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly          | Val<br>530              | Thr          | Asn          | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |
| <211<br><212 | )> SE<br>L> LE<br>L> TY | ENGTH<br>PE: | 1: 53<br>PRT | 39         |            |            | 7          |            |            |            |            |            |            |            |            |
| <220         | )> FE                   | EATUR        | RE:          |            | ific:      |            | _          |            |            |            |            |            |            |            |            |
|              |                         |              |              |            | rion:      | : Syr      | itnet      | ic E       | ютАІ       | pept:      | Lae        |            |            |            |            |
|              | )> SE                   |              |              |            | Val        | T10        | T1.0       | Dho        | Cor        | T 011      | Lou        | T10        | The        | Dro        | Cl n       |
| 1            |                         |              |              | 5          |            |            |            |            | 10         |            |            |            |            | 15         |            |
|              |                         |              | 20           |            | Ser        |            |            | 25         |            |            | _          |            | 30         |            |            |
| Glu          | Gly                     | Tyr<br>35    | Leu          | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr          | Leu<br>50               | Pro          | Val          | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | 60<br>Gåa  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65    | Leu                     | Ile          | Lys          | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Leu        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu          | Lys                     | Thr          | Val          | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn          | Pro                     | Gly          | Ser<br>100   | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala          | Ala                     | Ala<br>115   | Ala          | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg          | Leu<br>130              | Glu          | Ser          | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145   | Glu                     | Ala          | Val          | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala          | Val                     | Arg          | Glu          | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile          | Asn                     | Lys          | Asn<br>180   | Lys        | CAa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe          | Ser                     | Gln<br>195   | Phe          | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp          | Asn<br>210              | Ala          | Gly          | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225   | Glu                     | Leu          | Ala          | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile          | Lys                     | Leu          | Met          | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly          | Ile                     | Leu          | Ile<br>260   | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu          | Pro                     | Ile          | Phe          | Gly        | Val        | Ile        | Asp        | Thr        | Pro        | Cys        | Trp        | Ile        | Val        | Lys        | Ala        |

285

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280

|                              |            | 2/5                             |                             |            |            |            | 280            |            |            |            |            | 285        |            |            |            |
|------------------------------|------------|---------------------------------|-----------------------------|------------|------------|------------|----------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Ala                          | Pro<br>290 | Ser                             | Cys                         | Ser        | Glu        | Lys<br>295 | Lys            | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305                   | Asp        | Gln                             | Gly                         | Trp        | Tyr<br>310 | Cas        | Gln            | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro                          | Asn        | Glu                             | Lys                         | Asp<br>325 | Сув        | Glu        | Thr            | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr                          | Ala        | Ala                             | Gly<br>340                  | Ile        | Asn        | Val        | Ala            | Glu<br>345 | Gln        | Ser        | ГЛа        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn                          | Ile        | Ser<br>355                      | Thr                         | Thr        | Asn        | Tyr        | Pro<br>360     | Cys        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro                          | Ile<br>370 | Ser                             | Met                         | Val        | Ala        | Leu<br>375 | Ser            | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385                   | Lys        | Gly                             | Val                         | Ser        | 390        | Ser        | Ile            | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| ГÀа                          | Gln        | Leu                             | Asn                         | Lys<br>405 | Gly        | CÀa        | Ser            | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr                          | Val        | Thr                             | Ile<br>420                  | Asp        | Asn        | Thr        | Val            | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu                          | Gln        | His<br>435                      | Val                         | Ile        | ГÀа        | Gly        | Arg<br>440     | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile                          | Lys<br>450 | Phe                             | Pro                         | Glu        | Asn        | Gln<br>455 | Phe            | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465                   | Asn        | Ile                             | Glu                         | Asn        | Ser<br>470 | Gln        | Ala            | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu                          | Ser        | Ser                             | Ala                         | Glu<br>485 | ГÀа        | Gly        | Asn            | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu                          | Ile        | Ala                             | Val<br>500                  | Leu        | Gly        | Ser        | Ser            | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile                          | Ile        | Lys<br>515                      | Lys                         | Thr        | Lys        | Lys        | Pro<br>520     | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly                          | Val<br>530 | Thr                             | Asn                         | Asn        | Gly        | Phe<br>535 | Ile            | Pro        | His        | Asn        |            |            |            |            |            |
| <213<br><213<br><213<br><220 | )> FI      | ENGTI<br>YPE :<br>RGAN<br>EATUI | H: 53<br>PRT<br>ISM:<br>RE: | 39<br>Art: |            |            | Seque<br>nthet |            | ?olyţ      | pept:      | ide        |            |            |            |            |
| < 400                        | )> SI      | EQUEI                           | ICE :                       | 93         |            |            |                |            |            |            |            |            |            |            |            |
| Met<br>1                     | Ser        | Trp                             | Lys                         | Val<br>5   | Val        | Ile        | Ile            | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                          | Gly        | Leu                             | Lys<br>20                   | Glu        | Ser        | Tyr        | Leu            | Glu<br>25  | Glu        | Ser        | Cys        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly        | Tyr<br>35                       | Leu                         | Ser        | Val        | Leu        | Arg<br>40      | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50  | Glu                             | Val                         | Gly        | Asp        | Val<br>55  | Glu            | Asn        | Leu        | Thr        | 60<br>Cys  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu        | Ile                             | Lys                         | Thr        | Glu<br>70  | Leu        | Asp            | Leu        | Leu        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | Lys        | Thr                             | Val                         | Ser<br>85  | Ala        | Asp        | Gln            | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn                          | Pro        | Gly                             | Ser                         | Gly        | Ser        | Phe        | Val            | Leu        | Gly        | Ala        | Ile        | Ala        | Leu        | Gly        | Val        |

|            |            |            | 100        |            |            |            |            | 105        |            |            |            |            | 110        |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
|            |            |            | 100        |            |            |            |            | 103        |            |            |            |            | 110        |            |            |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГÀа        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | ràs        | Asp        | Phe        | Val        | Ser<br>170 | ГÀз        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | rys        | Asn<br>180 | ГÀа        | CAa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | ГÀа        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | CÀa        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | CÀa        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Сув        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390<br>CAa | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | ГÀа        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | ГÀа        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu        | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile        | Ile        | Lys<br>515 | Lys        | Thr        | Lys        | Lys        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
|            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |

| Gly                          | Val<br>530                       | Thr        | Asn                         | Asn        | Gly            | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |
|------------------------------|----------------------------------|------------|-----------------------------|------------|----------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| <211<br><212<br><213<br><220 | L> LE<br>2> TY<br>3> OF<br>0> FE | EATUF      | H: 53<br>PRT<br>[SM:<br>RE: | 39<br>Art: | Lfici<br>TION: |            | _          |            | Polyp      | pepti      | _de        |            |            |            |            |
| < 400                        | )> SE                            | EQUEN      | ICE :                       | 94         |                |            |            |            |            |            |            |            |            |            |            |
| Met<br>1                     | Ser                              | Trp        | Lys                         | Val<br>5   | Val            | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                          | Gly                              | Leu        | Lys<br>20                   | Glu        | Ser            | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | CAa        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly                              | Tyr<br>35  | Leu                         | Ser        | Val            | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50                        | Glu        | Val                         | Gly        | Asp            | Leu<br>55  | Glu        | Asn        | Leu        | Thr        | Cys        | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu                              | Ile        | Lys                         | Thr        | Glu<br>70      | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | Lys                              | Thr        | Val                         | Ser<br>85  | Ala            | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn                          | Pro                              | Gly        | Ser<br>100                  | Gly        | Ser            | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala                          | Ala                              | Ala<br>115 | Ala                         | Ala        | Val            | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg                          | Leu<br>130                       | Glu        | Ser                         | Glu        | Val            | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145                   | Glu                              | Ala        | Val                         | Ser        | Thr<br>150     | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala                          | Val                              | Arg        | Glu                         | Leu<br>165 | Lys            | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile                          | Asn                              | Lys        | Asn<br>180                  | Lys        | Cys            | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe                          | Ser                              | Gln<br>195 | Phe                         | Asn        | Arg            | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp                          | Asn<br>210                       | Ala        | Gly                         | Ile        | Thr            | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225                   | Glu                              | Leu        | Ala                         | Arg        | Ala<br>230     | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile                          | Lys                              | Leu        | Met                         | Leu<br>245 | Glu            | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly                          | Ile                              | Leu        | Ile<br>260                  | Gly        | Val            | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu                          | Pro                              | Ile<br>275 | Phe                         | Gly        | Val            | Ile        | Asp<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala                          | Pro<br>290                       | Ser        | Cys                         | Ser        | Glu            | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305                   | Asp                              | Gln        | Gly                         | Trp        | Tyr<br>310     | Cys        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro                          | Asn                              | Glu        | Lys                         | Asp<br>325 | Cys            | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Cys<br>335 | Asp        |
| Thr                          | Ala                              | Ala        | Gly<br>340                  | Ile        | Asn            | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |

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Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His 360 Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys 375 Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile 485 490 Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser 520 Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 95 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 95 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln 10 His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Thr Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile 120 Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr 135 Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr 150 155 Ala Val Arg Glu Leu Lys Asp Phe Val Leu Lys Asn Leu Thr Arg Ala 170

Ile Asn Lys Asn Lys Cys Asp Ile Asp Asp Leu Lys Met Ala Val Ser Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser 200 Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln Leu Pro Ile Phe Gly Val Ile Asp Thr Pro Cys Trp Ile Val Lys Ala Ala Pro Ser Cys Ser Glu Lys Lys Gly Asn Tyr Ala Cys Leu Leu Arg 295 Glu Asp Gln Gly Trp Tyr Cys Gln Asn Ala Gly Ser Thr Val Tyr Tyr 310 315 Pro Asn Glu Lys Asp Cys Glu Thr Arg Gly Asp His Val Phe Cys Asp Thr Ala Ala Gly Ile Asn Val Ala Glu Gln Ser Lys Glu Cys Asn Ile 345 Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His 360 Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys 375 Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp 410 Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly 425 Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser 520 Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 96

<sup>&</sup>lt;211> LENGTH: 539

<sup>&</sup>lt;212> TYPE: PRT

<sup>&</sup>lt;213 > ORGANISM: Artificial Sequence

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Synthetic Polypeptide

| Met<br>1   | Ser        | Trp        | Lys        | Val<br>5   | Val        | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| His        | Gly        | Leu        | Lys<br>20  | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | CÀa        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu        | Gly        | Tyr<br>35  | Leu        | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr        | Leu<br>50  | Glu        | Val        | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | 60<br>CAa  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu        | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | ГЛа        | Lys        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Trp        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | CAa        | Aap        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | Cys        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | CAa        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | Cys        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | Сув        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | ГÀа        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Cys        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | Cys        | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | Cys        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |

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Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 97 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 97 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Leu Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Leu Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr Ala Val Arg Glu Leu Lys Asp Phe Val Leu Lys Asn Leu Trp Arg Ala 170 Ile Asn Lys Asn Lys Cys Asp Ile Asp Asp Leu Lys Met Ala Val Ser Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser 200 Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp 215 Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln

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Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe
                                   250
Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln
Leu Pro Ile Phe Gly Val Ile Asp Thr Pro Cys Trp Ile Val Lys Ala
Ala Pro Ser Cys Ser Glu Lys Lys Gly Asn Tyr Ala Cys Leu Leu Arg
Glu Asp Gln Gly Trp Tyr Cys Gln Asn Ala Gly Ser Thr Val Tyr Tyr
Pro Asn Glu Lys Asp Cys Glu Thr Arg Gly Asp His Val Phe Cys Asp
Thr Ala Ala Gly Ile Asn Val Ala Glu Gln Ser Lys Glu Cys Asn Ile
Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His
Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys
Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile
Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp
                                410
Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly
          420
                             425
Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro
                          440
Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe
                       455
Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile
           470
Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile
                                   490
Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile
Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser
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Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn
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His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr
Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe
                40
Thr Leu Pro Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro
```

| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Leu        | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | ГЛа        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | CAa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala        | Pro<br>290 | Ser        | Cys        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | CAa        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | 390<br>GAa | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| ГÀа        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu        | Lys        | Gly        | Asn        | Thr        | Gly        | Phe        | Ile        | Ile        | Val        | Ile        | Ile        |

| _                            |  |                                  |                             | 485        |            |            |            |            | 490        |            |            |            |            | 495        |            |
|------------------------------|--|----------------------------------|-----------------------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| _                            |  |                                  |                             |            |            | _          | _          |            |            | _          |            |            |            |            |            |
| Leu                          | Ile  | Ala                              | Val<br>500                  | Leu        | GIY        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | 11e<br>510 | Phe        | Ile        |
| Ile                          | Ile  | Lys<br>515                       | Lys                         | Thr        | Lys        | Lys        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly                          | Val<br>530   | Thr                              | Asn                         | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |
| <213<br><213<br><213<br><220 | 0 > SI<br>1 > LI<br>2 > TY<br>3 > OF<br>0 > FI<br>3 > OT | ENGTI<br>(PE :<br>RGAN:<br>EATUI | H: 50<br>PRT<br>ISM:<br>RE: | 39<br>Art: |            |            | -          |            | Polyj      | pept:      | ide        |            |            |            |            |
| < 40                         | 0> SI  | EQUEI                            | NCE:                        | 99         |            |            |            |            |            |            |            |            |            |            |            |
| Met<br>1                     | Ser  | Trp                              | ГЛа                         | Val<br>5   | Val        | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His                          | Gly  | Leu                              | Lys<br>20                   | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | Cha        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly  | Tyr<br>35                        | Leu                         | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50  | Glu                              | Val                         | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | 60<br>CÀa  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu  | Ile                              | Lys                         | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | ГÀа  | Thr                              | Val                         | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn                          | Pro  | Gly                              | Ser<br>100                  | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala                          | Ala  | Ala<br>115                       | Ala                         | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГЛа        | Thr        | Ile        |
| Arg                          | Leu<br>130   | Glu                              | Ser                         | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145                   | Glu  | Ala                              | Val                         | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala                          | Val  | Arg                              | Glu                         | Leu<br>165 | rys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile                          | Asn  | Lys                              | Asn<br>180                  | Lys        | CAa        | Asp        | Ile        | Pro<br>185 | Asp        | Leu        | ГÀа        | Met        | Ala<br>190 | Val        | Ser        |
| Phe                          | Ser  | Gln<br>195                       | Phe                         | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp                          | Asn<br>210   | Ala                              | Gly                         | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225                   | Glu  | Leu                              | Ala                         | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile                          | Lys  | Leu                              | Met                         | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly                          | Ile  | Leu                              | Ile<br>260                  | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu                          | Pro  | Ile<br>275                       | Phe                         | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Сув        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala                          | Pro<br>290   | Ser                              | CÀa                         | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | CÀa        | Leu        | Leu        | Arg        |
| Glu                          | Asp  | Gln                              | Gly                         | Trp        | Tyr        |            | Gln        | Asn        | Ala        | Gly        |            | Thr        | Val        | Tyr        | Tyr        |
|                              |  |                                  |                             |            |            |            |            |            |            |            |            |            |            |            |            |

| 305  |            |            |            |            | 310        |            |            |            |            | 315        |            |            |            |            | 320        |
|--|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Pro  | Asn        | Glu        | Lys        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr  | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn  | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro  | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Сув        |
| Tyr<br>385   | ГЛа        | Gly        | Val        | Ser        | 390<br>CAa | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys  | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr  | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | ГÀа        | Val<br>430 | Glu        | Gly        |
| Glu  | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile  | Lys<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465   | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu  | Ser        | Ser        | Ala        | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu  | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile  | Ile        | Lys<br>515 | Lys        | Thr        | ГÀа        | ГÀа        | Pro<br>520 | Thr        | Gly        | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly  | Val<br>530 | Thr        | Asn        | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |
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|  |            |            |            | ORMA'      | TION       | : Syı      | nthet      | ic E       | olyr       | ept:       | ide        |            |            |            |            |
| < 400  | )> SI      | EQUEI      | NCE :      | 100        |            |            |            |            |            |            |            |            |            |            |            |
| Met<br>1   | Ser        | Trp        | Lys        | Val<br>5   | Val        | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His  | Gly        | Leu        | Lys<br>20  | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | СЛа        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu  | Gly        | Tyr<br>35  | Leu        | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr  | Leu<br>50  | Glu        | Val        | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | Cys<br>60  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu  | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn  | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala  | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | Lys        | Thr        | Ile        |
| Ara  | Leu        | Glu        | Ser        | Glu        | Val        | Thr        | Ala        | Ile        | Asn        | Asn        | Ala        | Leu        | ГХа        | Lys        | Thr        |

|            | 130        |            |            |            |            | 135        |            |            |            |            | 140        |            |            |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | CAa        | Pro        | Ile        | Asp<br>185 | Asp        | Leu        | ГÀз        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
|            |            | 275        |            |            | Val        |            | 280        |            |            |            |            | 285        |            |            |            |
|            | 290        |            | -          |            | Glu        | 295        | -          | _          |            |            | 300        |            |            |            |            |
| 305        | _          |            | -          | _          | Tyr<br>310 |            |            |            |            | 315        |            |            |            | -          | 320        |
|            |            |            | -          | 325        | CAa        |            |            |            | 330        |            |            |            |            | 335        |            |
|            |            |            | 340        |            | Asn        |            |            | 345        |            |            |            |            | 350        |            |            |
|            |            | 355        |            |            | Asn        | -          | 360        | -          | -          |            |            | 365        | -          | _          |            |
|            | 370        |            |            |            | Ala        | 375        |            |            |            |            | 380        |            |            |            |            |
| 385        | -          | -          |            |            | 390<br>Gly |            |            | _          |            | 395        | _          |            | -          |            | 400        |
| _          |            |            |            | 405        | Asn        | -          |            | -          | 410        |            |            |            | _          | 415        | _          |
|            |            |            | 420        | _          | Lys        |            |            | 425        |            |            |            | -          | 430        |            | _          |
|            |            | 435        |            |            | Asp        |            | 440        |            |            |            |            | 445        |            |            |            |
|            | 450        |            |            |            | Ser        | 455        |            |            |            |            | 460        | _          |            |            |            |
| 465        |            |            |            |            | 470<br>Lys |            |            |            |            | 475        |            |            |            |            | 480        |
|            |            |            |            | 485        | Gly        | -          |            |            | 490        |            |            |            |            | 495        |            |
|            |            |            | 500        |            |            |            |            | 505        |            |            |            |            | 510        |            |            |
|            |            | 515        |            |            | Lys        |            | 520        |            |            |            | PTO        | Pro<br>525 | GIU        | ьец        | ser        |
| Gly        | Val<br>530 | Thr        | Asn        | Asn        | Gly        | Phe<br>535 | Ile        | Pro        | His        | Asn        |            |            |            |            |            |

| <pre>&lt;211&gt; LENGTH: 539 &lt;212&gt; TYPE: PRT</pre>         |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |
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| Met<br>1   | Ser        | Trp        | Lys        | Val<br>5   | Val        | Ile        | Ile        | Phe        | Ser<br>10  | Leu        | Leu        | Ile        | Thr        | Pro<br>15  | Gln        |
| His  | Gly        | Leu        | Lys<br>20  | Glu        | Ser        | Tyr        | Leu        | Glu<br>25  | Glu        | Ser        | Сув        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu  | Gly        | Tyr<br>35  | Leu        | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr  | Leu<br>50  | Glu        | Val        | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | 60<br>CÀa  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu  | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn  | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala  | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГЛа        | Thr        | Ile        |
| Arg  | Leu<br>130 | Pro        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | ГЛа        | Lys        | Thr        |
| Asn<br>145   | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala  | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Aap        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile  | Asn        | Lys        | Asn<br>180 | Lys        | Сув        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | ГÀа        | Met        | Ala<br>190 | Val        | Ser        |
| Phe  | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp  | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225   | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile  | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly  | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu  | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala  | Pro<br>290 | Ser        | Cys        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305   | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro  | Asn        | Glu        | Lys        | Asp<br>325 | Cys        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr  | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | ГÀа        | Glu        | 350        | Asn        | Ile        |
| Asn  | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro  | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Сув        |

Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile 395 Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Asp Pro Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser 520 Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn 530 <210> SEQ ID NO 102 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 102 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Thr Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr 135 Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr 150 Ala Val Arg Glu Leu Lys Asp Phe Val Ser Lys Asn Leu Thr Arg Ala Ile Asn Lys Asn Lys Cys Asp Ile Asp Asp Leu Lys Met Ala Val Ser 185 Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser 200

Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp 215 Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln 230 Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln Leu Pro Ile Phe Gly Val Ile Asp Thr Pro Cys Trp Ile Val Lys Ala Ala Pro Ser Cys Ser Glu Lys Lys Gly Asn Tyr Ala Cys Leu Leu Arg Glu Asp Gln Gly Trp Tyr Cys Gln Asn Ala Gly Ser Thr Val Tyr Tyr Pro Asn Glu Lys Asp Cys Glu Thr Arg Gly Asp His Val Phe Cys Asp 330 Thr Ala Ala Gly Ile Asn Val Ala Glu Gln Ser Lys Glu Cys Asn Ile 345 Asn Ile Ser Thr Thr Asn Tyr Pro Cys Lys Val Ser Thr Gly Arg His 360 Pro Ile Ser Met Val Ala Leu Ser Pro Leu Gly Ala Leu Val Ala Cys Tyr Lys Gly Val Ser Cys Ser Ile Gly Ser Asn Arg Val Gly Ile Ile 390 395 Lys Gln Leu Asn Lys Gly Cys Ser Tyr Ile Thr Asn Gln Asp Ala Asp 405 410 Thr Val Thr Ile Asp Asn Thr Val Tyr Gln Leu Ser Lys Val Glu Gly Glu Gln His Val Ile Lys Gly Arg Pro Val Ser Ser Ser Phe Pro Pro 440 Ile Lys Phe Pro Glu Asp Gln Phe Gln Val Ala Leu Asp Gln Val Phe 455 Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 103 <211> LENGTH: 539 <212> TYPE: PRT <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEQUENCE: 103 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln 10

His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr 20 25 30

| Glu        | Gly        | Tyr<br>35  | Leu        | Ser        | Val        | Leu        | Arg<br>40  | Thr        | Gly        | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Thr        | Leu<br>50  | Glu        | Val        | Gly        | Asp        | Val<br>55  | Glu        | Asn        | Leu        | Thr        | Cys        | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65  | Leu        | Ile        | Lys        | Thr        | Glu<br>70  | Leu        | Asp        | Leu        | Thr        | Lys<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu        | Lys        | Thr        | Val        | Ser<br>85  | Ala        | Asp        | Gln        | Leu        | Ala<br>90  | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГÀа        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | ГÀа        | ГÀа        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | CÀa        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Phe        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | ГÀЗ        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | CAa        | Trp        | Ile<br>285 | Val        | ГÀз        | Ala        |
| Ala        | Pro<br>290 | Ser        | Сув        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | CAa        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Сув        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | CÀa        |
| Tyr<br>385 | ГÀа        | Gly        | Val        | Ser        | 390<br>CAa | Ser        | Ile        | Gly        | Ser        | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | CÀa        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |

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Ile Lys Phe Pro Glu Asn Gln Phe Gln Val Ala Leu Asp Gln Val Phe 455 Glu Asn Ile Glu Asn Ser Gln Ala Leu Val Asp Gln Ser Asn Arg Ile Leu Ser Ser Ala Glu Lys Gly Asn Thr Gly Phe Ile Ile Val Ile Ile Leu Ile Ala Val Leu Gly Ser Ser Met Ile Leu Val Ser Ile Phe Ile 505 Ile Ile Lys Lys Thr Lys Lys Pro Thr Gly Ala Pro Pro Glu Leu Ser Gly Val Thr Asn Asn Gly Phe Ile Pro His Asn <210> SEQ ID NO 104 <211> LENGTH: 539 <212> TYPE: PRT <213 > ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polypeptide <400> SEOUENCE: 104 Met Ser Trp Lys Val Val Ile Ile Phe Ser Leu Leu Ile Thr Pro Gln His Gly Leu Lys Glu Ser Tyr Leu Glu Glu Ser Cys Ser Thr Ile Thr 25 Glu Gly Tyr Leu Ser Val Leu Arg Thr Gly Trp Tyr Thr Asn Val Phe Thr Leu Glu Val Gly Asp Val Glu Asn Leu Thr Cys Ser Asp Gly Pro Ser Leu Ile Lys Thr Glu Leu Asp Leu Thr Lys Ser Ala Leu Arg Glu Leu Lys Thr Val Ser Ala Asp Gln Leu Ala Arg Glu Glu Gln Ile Glu Asn Pro Gly Ser Gly Ser Phe Val Leu Gly Ala Ile Ala Leu Gly Val 105 Ala Ala Ala Ala Val Thr Ala Gly Val Ala Ile Ala Lys Thr Ile Arg Leu Glu Ser Glu Val Thr Ala Ile Asn Asn Ala Leu Lys Lys Thr Asn Glu Ala Val Ser Thr Leu Gly Asn Gly Val Arg Val Leu Ala Thr Ala Val Arg Glu Leu Lys Asp Phe Val Ser Lys Asn Leu Thr Arg Ala Ile Asn Lys Asn Lys Cys Asp Ile Asp Asp Leu Lys Met Ala Val Ser Phe Ser Gln Phe Asn Arg Arg Phe Leu Asn Val Val Arg Gln Phe Ser 200 Asp Asn Ala Gly Ile Thr Pro Ala Ile Ser Leu Asp Leu Met Thr Asp Ala Glu Leu Ala Arg Ala Val Pro Asn Met Pro Thr Ser Ala Gly Gln 230 Ile Lys Leu Met Leu Glu Asn Arg Ala Met Val Arg Arg Lys Gly Phe 250 Gly Ile Leu Ile Gly Val Tyr Gly Ser Ser Val Ile Tyr Met Val Gln 265

|                              |                                  |                                  |             |            |            |            |                |            |                   |            |            | COII       | CIII       | aca        |            |
|------------------------------|----------------------------------|----------------------------------|-------------|------------|------------|------------|----------------|------------|-------------------|------------|------------|------------|------------|------------|------------|
| Leu                          | Pro                              | Ile<br>275                       | Phe         | Gly        | Val        | Ile        | Asp<br>280     | Thr        | Pro               | Cys        | Trp        | Ile<br>285 | Val        | Lys        | Ala        |
| Ala                          | Pro<br>290                       | Ser                              | CAa         | Ser        | Glu        | Lys<br>295 | Lys            | Gly        | Asn               | Tyr        | Ala<br>300 | CAa        | Leu        | Leu        | Arg        |
| Glu<br>305                   | Asp                              | Gln                              | Gly         | Trp        | Tyr<br>310 | CAa        | Gln            | Asn        | Ala               | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro                          | Asn                              | Glu                              | Lys         | Asp<br>325 | CÀa        | Glu        | Thr            | Arg        | Gly<br>330        | Asp        | His        | Val        | Phe        | Cys        | Asp        |
| Thr                          | Ala                              | Ala                              | Gly<br>340  | Ile        | Asn        | Val        | Ala            | Glu<br>345 | Gln               | Ser        | ГÀа        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn                          | Ile                              | Ser<br>355                       | Thr         | Thr        | Asn        | Tyr        | Pro<br>360     | Cys        | Lys               | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro                          | Ile<br>370                       | Ser                              | Met         | Val        | Ala        | Leu<br>375 | Ser            | Pro        | Leu               | Gly        | Ala<br>380 | Leu        | Val        | Ala        | Cys        |
| Tyr<br>385                   | Lys                              | Gly                              | Val         | Ser        | Cys        | Ser        | Ile            | Gly        | Ser               | Asn<br>395 | Arg        | Val        | Gly        | Ile        | Ile<br>400 |
| Lys                          | Gln                              | Leu                              | Asn         | Lys<br>405 | Gly        | CAa        | Ser            | Tyr        | Ile<br>410        | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr                          | Val                              | Thr                              | Ile<br>420  | Asp        | Asn        | Thr        | Val            | Tyr<br>425 | Gln               | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu                          | Gln                              | His<br>435                       | Val         | Ile        | Lys        | Gly        | Arg<br>440     | Pro        | Val               | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile                          | Lys<br>450                       | Phe                              | Pro         | Gln        | Asp        | Gln<br>455 | Phe            | Gln        | Val               | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465                   | Asn                              | Ile                              | Glu         | Asn        | Ser<br>470 | Gln        | Ala            | Leu        | Val               | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu                          | Ser                              | Ser                              | Ala         | Glu<br>485 | Lys        | Gly        | Asn            | Thr        | Gly<br>490        | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu                          | Ile                              | Ala                              | Val<br>500  | Leu        | Gly        | Ser        | Ser            | Met<br>505 | Ile               | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile                          | Ile                              | Lys<br>515                       | Lys         | Thr        | Lys        | Lys        | Pro<br>520     | Thr        | Gly               | Ala        | Pro        | Pro<br>525 | Glu        | Leu        | Ser        |
| Gly                          | Val<br>530                       | Thr                              | Asn         | Asn        | Gly        | Phe<br>535 | Ile            | Pro        | His               | Asn        |            |            |            |            |            |
| <211<br><212<br><213<br><220 | L> LE<br>2> TY<br>3> OF<br>0> FE | ENGTI<br>(PE :<br>RGAN)<br>EATUI | ISM:<br>RE: | 39<br>Art: |            |            | Seque<br>nthet |            | ?oly <sub>l</sub> | pept:      | ide        |            |            |            |            |
|                              |                                  | ~                                | NCE:        |            | Val        | Tle        | Ile            | Phe        | Ser               | Len        | Len        | Tle        | Thr        | Pro        | Gln        |
| 1                            |                                  |                                  |             | 5          |            |            |                |            | 10                |            |            |            |            | 15         |            |
| His                          | Gly                              | Leu                              | Lув<br>20   | Glu        | Ser        | Tyr        | Leu            | Glu<br>25  | Glu               | Ser        | Cys        | Ser        | Thr<br>30  | Ile        | Thr        |
| Glu                          | Gly                              | Tyr<br>35                        | Leu         | Ser        | Val        | Leu        | Arg<br>40      | Thr        | Gly               | Trp        | Tyr        | Thr<br>45  | Asn        | Val        | Phe        |
| Thr                          | Leu<br>50                        | Glu                              | Val         | Gly        | Asp        | Val<br>55  | Glu            | Asn        | Leu               | Thr        | 60<br>Cys  | Ser        | Asp        | Gly        | Pro        |
| Ser<br>65                    | Leu                              | Ile                              | Lys         | Thr        | Glu<br>70  | Leu        | Asp            | Leu        | Thr               | Lуs<br>75  | Ser        | Ala        | Leu        | Arg        | Glu<br>80  |
| Leu                          | Lys                              | Thr                              | Val         | Ser<br>85  | Ala        | Asp        | Gln            | Leu        | Ala<br>90         | Arg        | Glu        | Glu        | Gln        | Ile<br>95  | Glu        |

85 90 95

| _          |            |            |            |            |            |            |            |            |            |            |            |            |            |            |            |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Asn        | Pro        | Gly        | Ser<br>100 | Gly        | Ser        | Phe        | Val        | Leu<br>105 | Gly        | Ala        | Ile        | Ala        | Leu<br>110 | Gly        | Val        |
| Ala        | Ala        | Ala<br>115 | Ala        | Ala        | Val        | Thr        | Ala<br>120 | Gly        | Val        | Ala        | Ile        | Ala<br>125 | ГÀа        | Thr        | Ile        |
| Arg        | Leu<br>130 | Glu        | Ser        | Glu        | Val        | Thr<br>135 | Ala        | Ile        | Asn        | Asn        | Ala<br>140 | Leu        | Lys        | Lys        | Thr        |
| Asn<br>145 | Glu        | Ala        | Val        | Ser        | Thr<br>150 | Leu        | Gly        | Asn        | Gly        | Val<br>155 | Arg        | Val        | Leu        | Ala        | Thr<br>160 |
| Ala        | Val        | Arg        | Glu        | Leu<br>165 | Lys        | Asp        | Phe        | Val        | Ser<br>170 | Lys        | Asn        | Leu        | Thr        | Arg<br>175 | Ala        |
| Ile        | Asn        | Lys        | Asn<br>180 | Lys        | СЛа        | Asp        | Ile        | Asp<br>185 | Asp        | Leu        | Lys        | Met        | Ala<br>190 | Val        | Ser        |
| Phe        | Ser        | Gln<br>195 | Trp        | Asn        | Arg        | Arg        | Phe<br>200 | Leu        | Asn        | Val        | Val        | Arg<br>205 | Gln        | Phe        | Ser        |
| Asp        | Asn<br>210 | Ala        | Gly        | Ile        | Thr        | Pro<br>215 | Ala        | Ile        | Ser        | Leu        | Asp<br>220 | Leu        | Met        | Thr        | Asp        |
| Ala<br>225 | Glu        | Leu        | Ala        | Arg        | Ala<br>230 | Val        | Pro        | Asn        | Met        | Pro<br>235 | Thr        | Ser        | Ala        | Gly        | Gln<br>240 |
| Ile        | Lys        | Leu        | Met        | Leu<br>245 | Glu        | Asn        | Arg        | Ala        | Met<br>250 | Val        | Arg        | Arg        | Lys        | Gly<br>255 | Phe        |
| Gly        | Ile        | Leu        | Ile<br>260 | Gly        | Val        | Tyr        | Gly        | Ser<br>265 | Ser        | Val        | Ile        | Tyr        | Met<br>270 | Val        | Gln        |
| Leu        | Pro        | Ile<br>275 | Phe        | Gly        | Val        | Ile        | Asp<br>280 | Thr        | Pro        | Càa        | Trp        | Ile<br>285 | Val        | ГÀЗ        | Ala        |
| Ala        | Pro<br>290 | Ser        | Сув        | Ser        | Glu        | Lys<br>295 | Lys        | Gly        | Asn        | Tyr        | Ala<br>300 | Cys        | Leu        | Leu        | Arg        |
| Glu<br>305 | Asp        | Gln        | Gly        | Trp        | Tyr<br>310 | Cys        | Gln        | Asn        | Ala        | Gly<br>315 | Ser        | Thr        | Val        | Tyr        | Tyr<br>320 |
| Pro        | Asn        | Glu        | Lys        | Asp<br>325 | CAa        | Glu        | Thr        | Arg        | Gly<br>330 | Asp        | His        | Val        | Phe        | Сув<br>335 | Asp        |
| Thr        | Ala        | Ala        | Gly<br>340 | Ile        | Asn        | Val        | Ala        | Glu<br>345 | Gln        | Ser        | Lys        | Glu        | Сув<br>350 | Asn        | Ile        |
| Asn        | Ile        | Ser<br>355 | Thr        | Thr        | Asn        | Tyr        | Pro<br>360 | Cys        | Lys        | Val        | Ser        | Thr<br>365 | Gly        | Arg        | His        |
| Pro        | Ile<br>370 | Ser        | Met        | Val        | Ala        | Leu<br>375 | Ser        | Pro        | Leu        | Gly        | Ala<br>380 | Leu        | Val        | Ala        | CAa        |
| Tyr<br>385 | Lys        | Gly        | Val        | Ser        | Сув<br>390 |            | Ile        | Gly        |            | Asn<br>395 |            | Val        | Gly        | Ile        | Ile<br>400 |
| Lys        | Gln        | Leu        | Asn        | Lys<br>405 | Gly        | Cys        | Ser        | Tyr        | Ile<br>410 | Thr        | Asn        | Gln        | Asp        | Ala<br>415 | Asp        |
| Thr        | Val        | Thr        | Ile<br>420 | Asp        | Asn        | Thr        | Val        | Tyr<br>425 | Gln        | Leu        | Ser        | Lys        | Val<br>430 | Glu        | Gly        |
| Glu        | Gln        | His<br>435 | Val        | Ile        | Lys        | Gly        | Arg<br>440 | Pro        | Val        | Ser        | Ser        | Ser<br>445 | Phe        | Asp        | Pro        |
| Ile        | Lys<br>450 | Phe        | Pro        | Glu        | Asp        | Gln<br>455 | Phe        | Gln        | Val        | Ala        | Leu<br>460 | Asp        | Gln        | Val        | Phe        |
| Glu<br>465 | Asn        | Ile        | Glu        | Asn        | Ser<br>470 | Gln        | Ala        | Leu        | Val        | Asp<br>475 | Gln        | Ser        | Asn        | Arg        | Ile<br>480 |
| Leu        | Ser        | Ser        | Ala        | Glu<br>485 | Lys        | Gly        | Asn        | Thr        | Gly<br>490 | Phe        | Ile        | Ile        | Val        | Ile<br>495 | Ile        |
| Leu        | Ile        | Ala        | Val<br>500 | Leu        | Gly        | Ser        | Ser        | Met<br>505 | Ile        | Leu        | Val        | Ser        | Ile<br>510 | Phe        | Ile        |
| Ile        | Ile        | ГЛа        | Lys        | Thr        | ГЛа        | Lys        | Pro        | Thr        | Gly        | Ala        | Pro        | Pro        | Glu        | Leu        | Ser        |

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                                                                     240
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                                                                     480
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                                                                     540
                                                                     600
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acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc
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tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac
                                                                     960
cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga
                                                                    1020
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tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc
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                                                                    1200
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gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga
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gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc
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ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg
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| accggctggt acaccaacgt gttcacactg gaagtgggcg acgtcgagaa tctgacatgc   | 180  |
| tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
| ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca   | 360  |
| ggcgtggcca tctgcaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480  |
| gccgtgcgcg agctgaagga cttcgtgtcc aagaacctga cacgggccat taacaagaac   | 540  |
| aagtgcgaca tcgacgacct gaagatggcc gtgtccttta gccagttcaa ccggcggttt   | 600  |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660  |
| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag   | 720  |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgtgt   | 780  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840  |
| acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc   | 900  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020 |
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| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc   | 1140 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200 |
| aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc   | 1260 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320 |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgagc accagtggca tgtggccctg   | 1380 |
| gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc   | 1440 |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg   | 1500 |
| ctgggcagct ccatgatect ggtgtecate tteateatta teaagaagae caagaagee  | 1560 |
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| accggctggt acaccaacgt gttcacactg gaagtgggcg acgtcgagaa tctgacatgc   | 180  |
| tetgatggce ctageetgat caagacegag etggatetge teaagagege eetgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
| ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca   | 360  |
| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
|   |      |

480

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| ggcgtgtacg   | gcagcagcgt  | gatctatatg  | gtgcagctgc  | ctatcttcgg   | cgtgatcgac   | 840  |
| acaccctgct   | ggattgtgaa  | ggeegeteet  | agctgtagcg  | agaagaaggg   | caattacgcc   | 900  |
| tgcctgctga   | gagaggacca  | aggctggtat  | tgtcagaacg  | ccggcagcac   | cgtgtactac   | 960  |
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| accggctggt<br>tctgatggcc   | aggtggtcat<br>tggaagagtc<br>acaccaacgt<br>ctagcctgat  | ctgcagcacc<br>gttcacactg<br>caagaccgag  | atcacagagg<br>gaagtgggcg<br>ctggatctgc  | caceteagea<br>getacetgte<br>aegtegagaa<br>teaagagege   | tgtgctgaga<br>tctgacatgc<br>cctgagagaa   | 120<br>180<br>240  |
| accggctggt<br>tctgatggcc<br>ctcaagaccg   | aggtggtcat<br>tggaagagtc<br>acaccaacgt<br>ctagcctgat<br>tgtctgccga  | ctgcagcacc<br>gttcacactg<br>caagaccgag<br>tcagctggcc  | atcacagagg<br>gaagtgggcg<br>ctggatctgc<br>agagaggaac  | caceteagea<br>getacetgte<br>aegtegagaa<br>teaagagege<br>agategagaa   | tgtgctgaga<br>tctgacatgc<br>cctgagagaa<br>tcctggcagc   | 120<br>180<br>240<br>300   |
| accggctggt<br>tctgatggcc<br>ctcaagaccg<br>ggcagctttg   | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgtctgccga tgctgggagc   | ctgcagcacc<br>gttcacactg<br>caagaccgag<br>tcagctggcc<br>cattgctctt  | atcacagagg<br>gaagtgggcg<br>ctggatctgc<br>agagaggaac<br>ggagtggctg  | cacctcagca<br>gctacctgtc<br>acgtcgagaa<br>tcaagagcgc<br>agatcgagaa<br>ctgctgcagc   | tgtgctgaga tctgacatgc cctgagagaga tcctggcagc tgttacagca  | 120<br>180<br>240<br>300<br>360                                    |
| accggctggt tctgatggcc ctcaagaccg ggcagctttg ggcgtggcca   | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgtctgccga tgctgggagc tcgctaagac  | ctgcagcacc<br>gttcacactg<br>caagaccgag<br>tcagctggcc<br>cattgctctt<br>catcagactg  | atcacagagg<br>gaagtgggcg<br>ctggatctgc<br>agagaggaac<br>ggagtggctg<br>gaaagcgaag  | caceteagea<br>getacetgte<br>aegtegagaa<br>teaagagege<br>agategagaa<br>etgetgeage<br>tgacegeeat   | tgtgctgaga tctgacatgc cctgagagaa tcctggcagc tgttacagca caacaacgcc  | 120<br>180<br>240<br>300<br>360<br>420                             |
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| accggctggt tetgatggcc ctcaagaccg ggcagctttg ggcgtggcca ctgaagaaga gccgtgcgcg   | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgctgccga tgctgggagc tcgctaagac caaacgaggc                                  | ctgcagcacc gttcacactg caagaccgag tcagctggcc cattgctctt catcagactg cgtcagcaca cttcgtgtcc   | atcacagagg gaagtgggcg ctggatctgc agagaggaac ggagtggctg gaaagcgaag ctcggcaatg aagaacctga   | caceteagea<br>getacetgte<br>aegtegagaa<br>teaagagege<br>agategagaa<br>etgetgeage<br>tgacegeeat<br>gegttagagt   | tgtgctgaga tctgacatgc cctgagagaaa tcctggcagc tgttacagca caacaacgcc gctggccaca taacaagaac   | 120<br>180<br>240<br>300<br>360<br>420<br>480                      |
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| accggctggt tctgatggcc ctcaagaccg ggcagctttg ggcgtggcca ctgaagaaga gccgtgcgcg aagtgcgaca                                  | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgctgccga tgctgggagc tcgctaagac caaacgaggc                                  | ctgcagcacc gttcacactg caagaccgag tcagctggcc cattgctctt catcagactg cgtcagcaca cttcgtgtcc gaagatggcc                                  | atcacagagg gaagtgggcg ctggatctgc agagaggaac ggagtggctg gaaagcgaag ctcggcaatg aagaacctga   | cacctcagca gctacctgtc acgtcgagaa tcaagagcgc agatcgagaa ctgctgcagc tgaccgccat gcgttagagt cacgggccat gcacgtcaa   | tgtgctgaga tctgacatgc cctgagagaaa tcctggcagc tgttacagca caacaacgcc gctggccaca taacaagaac ccggcggttt                                  | 120<br>180<br>240<br>300<br>360<br>420<br>480                      |
| accggctggt tctgatggcc ctcaagaccg ggcagctttg ggcgtggcca ctgaagaaga gccgtgcgcg aagtgcgaca ctgaacgtcg                       | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgtctgccga tgctgggagc tcgctaagac caaacgaggc agctgaagga                      | ctgcagcacc gttcacactg caagaccgag tcagctggcc cattgctctt catcagactg cgtcagcaca cttcgtgtcc gaagatggcc tagcgacaac                       | atcacagagg gaagtgggcg ctggatctgc agagaggaac ggagtggctg gaaagcgaag ctcggcaatg aagaacctga gtgtccttta gccggaatca                       | cacctcagca gctacctgtc acgtcgagaa tcaagagcgc agatcgagaa ctgctgcagc tgaccgccat gcgttagagt cacgggccat gccagttcaa caccagccat                               | tgtgctgaga tctgacatgc cctgagagaaa tcctggcagc tgttacagca caacaacgcc gctggccaca taacaagaac ccggcggttt cagcctggac                       | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540               |
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| accggctggt tctgatggcc ctcaagaccg ggcagctttg ggcgtggcca ctgaagaaga gccgtgcgcg aagtgcgaca ctgaacgtcg ctgatgacag ctgatgacag | aggtggtcat tggaagagtc acaccaacgt ctagcctgat tgctgccga tgctgggagc tcgctaagac caaacgaggc agctgaagga tccctgacct tgcggcagtt | ctgcagcacc gttcacactg caagaccgag tcagctggcc cattgctctt catcagactg cgtcagcaca cttcgtgtcc gaagatggcc tagcgacaac ggctagagcc tagagccatg | atcacagagg gaagtgggcg ctggatctgc agagaggaac ggagtggctg gaaagcgaag ctcggcaatg aagaacctga gtgtccttta gccggaatca gtgcctaaca gtccgacgga | cacctcagca gctacctgtc acgtcgagaa tcaagagcgc agatcgagaa ctgctgcagc tgaccgccat gcgttagagt cacgggccat gccagttcaa caccagccat tgcctacatc                    | tgtgctgaga tctgacatgc cctgagagaaa tcctggcagc tgttacagca caacaacgcc gctggccaca taacaagaac ccggcggttt cagcctggac tgccggccag cattctgatt | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660 |

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|--|--|------|
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| tgcanagts; concegging gracectatt tetatigstag cittistetee tetaggagee 1140 ctggrigett strataagge optstoctst ageatoegga gracecate 1200 aageagetga acaagggetg cagetacate aceaaccagg acescogtate optstacete 1200 gacaacacg tstateaget gageaagsta gaaggegaa accasteat cageagstaga 1320 cetsgricea genagitega cectateaag teecetgag tegeacagte canacagaate 1440 ctgctetagog cogagaacga cagaattee caggetetg tgcaccagte canacagaate 1440 ctgctetagog cogagaagga aacaccagg teecateate genacacte 1200 aceggeget cetegateet ggtgeeate teetatata teaaggaage caagaagee 1560 aceggegete cetegateet ggtgeeate teetateatta teaaggaage caagaagee 1660 aceggegete cetegateet ggtgeeate teetateatta teaaggaage caagaagee 1670 ctgggeaget cetegateet ggtgeeate teetateatta teaaggaage caagaagee 1670 aceggeggete cetegateet gageggagt aceaacaatg getteateee teaacacc 1617 <pre> <pre> <pre> <pre></pre></pre></pre></pre>  | cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga                              | 1020 |
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| asgocagotta acasagacta caquacate accasecaga acquegatae egiqaccate gacaacaccag tytatcagot gaquasgig gaaggogaac agcacqigat caagggcaga 1320 cotgigicaca gaagottega coctatoaag ticootgaga accagitica ggiggocotg gaccaacaccag tytatcagot gagaastica caggactigg gagacgogat caacaggaat cotgotagog cogagaaggg acaacacggg ticatcatag tygaccate tracacaga 1440 ctgotagog cogagaaggg acaacacggg ticatcatag tygaccate tracacaga 1500 ctgggcagot coatgatect ggigtocate troatcata toaagaagac caagaagac caagaagac ctgggcagot coatgatect ggigtocate troatcata toaagaagac caagaagac caagaagac caggacgote otccagaact gagoggagtg accaacaatg gotteatcoc toacaac ctgggcagot coatgatect ggigtocate troatcata toaagaagac caagaagac cataacaaca coggocote otccagaact gagoggagtg accaacaatg gotteatcoc toacaac ctgggcagot coatgatect gagoggagga accaacaatg gotteatcoc toacaac ctggcaga aggigtotat taitical Sequence ctgotagaagato tratificial Sequence ctgotagaga aggigtotat catottcago ctgotgatac cacctcagca cggcotgaaa ctgagotgga aggigtotat ctgcagcacc atcacagaag gotacctgc tygigotgaa totgatgga aggigtotat ctgcagcacc atcacagaag gotacctgc tygigotgaga totgatggc ctagacgaga tracacatg gaagtggga acgtcagaaa totgacaaca totgaagaaga gagacgaag cattgatgca gaagaagaa gaacgaaga tygitacacaacagac totgaagaaga catagaagaag cattgatgaga gaacgaaga tygitaagag tygitacaacaacagac ggoggggcaa togctagaagac cattgatgatg gaagaagaag gaacgcagaa tacaacaacaga ggogggcaa togctagaagaa catcagacag gaaagaagaag gaacgccat taacaacagaa 400 gcoggacgaa gagaagaagaagaagaagaagaagaagaagaagaag  | tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc                              | 1140 |
| gacacacacog tgtatcagot gagacaaggt gaaggegaac ageacgtgat caagggcaga 1320 cctgtgtoca geagettega cectatcaag tteectgaga accagtteca ggtggecetg 1380 gaccaggtgt togagaacat ogagaattec caggatetga gagacagte caacagaate 1440 ctgtctagog eegagaaggg aaacacegge tteatcateg tgateatect gategeetgg 1500 ctgggcaget categatect ggtgtccate tteatcatat toaagaagac caagaagec 1560 accggcoget etccagaact gageggagg accacaaatg getteatece teacaac 1617 <pre> &lt;210</pre>   | ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc                              | 1200 |
| cetgtgteca geagettega cectateaag ttecetgaga accagtteca ggtggecetg gaccaggtgt tegagaacat egagaattee caggatetgt ggaccagte caacagaate 1440 etgtetageg cegagaaggg aaacacegge tteateateg tgateateet gategeegtg 1500 etgggcaget ecatgateet ggtgtecate tteateatta teaagaagac caagaagece 1560 accgggegte etceagaact gageggagtg accaccagt getteateee teacaac 1517  **210-SEO ID NO 110 **2115-ISMNTH: 1617 **212-TFFE: DNA **213-ORGAINSH: Artificial Sequence **223-OFBATURE: **223-OF | aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc                              | 1260 |
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| aceggegete etceagaact gageggagtg aceaacaatg getteatece teacaac 1617  | ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg                              | 1500 |
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| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt 780 ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac 840 acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc 900 tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac 960 cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga 1020 atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc 1080 tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc 1140 ctggtggctt gttataaggg cgtgtcctgt agcatcgga gcaacaggg gggcatcatc 1200 aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260   | ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac                              | 660  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac 840  acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc 900  tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac 960  cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga 1020  atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc 1080  tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc 1140  ctggtggctt gttataaggg cgtgtcctgt agcatcgca gcaacagagt gggcatcatc 1200  aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260   | ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag                              | 720  |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee 900 tgeetgetga gagaggacea aggetggtat tgteagaaeg eeggeageae egtgtaetae 960 eetaaegaga aggaetgega gacaagagge gaceaegtgt tetgtgatae egeegetgga 1020 ateaatgtgg eegageagag caaagagtge aacateaaea teageaceae eaactateee 1080 tgeaaggtgt eeaeeggeag geaeeetatt tetatggtgg etetgtetee tetgggagee 1140 etggtggett gttataaggg egtgteetgt ageateggea geaaeagagt gggeateate 1200 aageagetga acaagggetg eagetaeate aceaaeeagg aegeegatae egtgaceate 1260   | atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt                              | 780  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac 960  cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga 1020  atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc 1080  tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc 1140  ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc 1200  aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260  | ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac                              | 840  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tetgtgatac egecgetgga 1020 atcaatgtgg ccgagcagag caaagagtgc aacatcaaca teagcaccac caactatece 1080 tgcaaggtgt ccaceggcag gcaccetatt tetatggtgg etetgtetee tetgggagec 1140 ctggtggett gttataaggg egtgteetgt agcateggca gcaacagagt gggcateate 1200 aagcagetga acaagggetg cagetacate accaaccagg aegecgatac egtgaccate 1260   | acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee                              | 900  |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc 1080 tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc 1140 ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc 1200 aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260  | tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac                              | 960  |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc 1140 ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc 1200 aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260   | cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga                              | 1020 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc 1200 aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260  | atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc                              | 1080 |
| aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc 1260   | tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc                              | 1140 |
|  | ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc                              | 1200 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga 1320   | aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc                              | 1260 |
|  | gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga                              | 1320 |

cctgtgtcca gcagcttcga ccctatcaag ttccctgagg atcagttcca ggtggccctg 1380

| gaccaggtgt tcgagaacat  | cgagaattcc | caggetetgg | tggaccagtc | caacagaatc | 1440 |
|--|------------|------------|------------|------------|------|
| ctgtctagcg ccgagaaggg  |            |            |            |            | 1500 |
| ctgggcaget ccatgatect  |            |            |            |            | 1560 |
| accggcgctc ctccagaact  |            |            |            |            | 1617 |
| 33 3   | 3 3 33 3 3 | 3          | 3          |            |      |
| <pre>&lt;210&gt; SEQ ID NO 111 &lt;211&gt; LENGTH: 1617 &lt;212&gt; TYPE: DNA &lt;213&gt; ORGANISM: Artif &lt;220&gt; FEATURE: &lt;223&gt; OTHER INFORMATION</pre> |            |            | eotide     |            |      |
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| gagagctacc tggaagagtc  | ctgcagcacc | atcacagagg | gctacctgtc | tgtgctgaga | 120  |
| accggctggt acaccaacgt  | gttcacactg | gaagtgggcg | acgtcgagaa | tctgacatgc | 180  |
| totgatggcc ctagcotgat  | caagaccgag | ctggatctgc | tcaagagcgc | cctgagagaa | 240  |
| ctcaagaccg tgtctgccga  | tcagctggcc | agagaggaac | agatcgagaa | tectggcage | 300  |
| ggcagctttg tgctgggagc  | cattgctctt | ggagtggctg | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca tcgctaagac  | catcagactg | gaaagcgaag | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga caaacgaggc  | cgtcagcaca | ctcggcaatg | gcgttagagt | gctggccaca | 480  |
| gccgtgcgcg agctgaagga  | cttcgtgctt | aagaacctga | cacgggccat | taacaagaac | 540  |
| aagtgcgaca tccctgacct  | gaagatggcc | gtgtccttta | gccagttcaa | ccggcggttt | 600  |
| ctgaacgtcg tgcggcagtt  | tagcgacaac | gccggaatca | caccagccat | cagcctggac | 660  |
| ctgatgacag atgctgagct  | ggctagagcc | gtgcctaaca | tgcctacatc | tgccggccag | 720  |
| atcaagctga tgctcgagaa  | tagagccatg | gtccgacgga | aaggcttcgg | cattctgatt | 780  |
| ggcgtgtacg gcagcagcgt  | gatctatatg | gtgcagctgc | ctatcttcgg | cgtgatcgac | 840  |
| acaccctgct ggattgtgaa  | ggccgctcct | agctgtagcg | agaagaaggg | caattacgcc | 900  |
| tgcctgctga gagaggacca  | aggctggtat | tgtcagaacg | ccggcagcac | cgtgtactac | 960  |
| cctaacgaga aggactgcga  | gacaagaggc | gaccacgtgt | tctgtgatac | cgccgctgga | 1020 |
| atcaatgtgg ccgagcagag  | caaagagtgc | aacatcaaca | tcagcaccac | caactatccc | 1080 |
| tgcaaggtgt ccaccggcag  | gcaccctatt | tctatggtgg | ctctgtctcc | tetgggagee | 1140 |
| ctggtggctt gttataaggg  | cgtgtcctgt | agcatcggca | gcaacagagt | gggcatcatc | 1200 |
| aagcagctga acaagggctg  | cagctacatc | accaaccagg | acgccgatac | cgtgaccatc | 1260 |
| gacaacaccg tgtatcagct  | gagcaaggtg | gaaggcgaac | agcacgtgat | caagggcaga | 1320 |
| cctgtgtcca gcagcttcga  | ccctatcaag | ttccctgaga | accagttcca | ggtggccctg | 1380 |
| gaccaggtgt tcgagaacat  | cgagaattcc | caggetetgg | tggaccagtc | caacagaatc | 1440 |
| ctgtctagcg ccgagaaggg  | aaacaccggc | ttcatcatcg | tgatcatcct | gatcgccgtg | 1500 |
| ctgggcagct ccatgatcct  | ggtgtccatc | ttcatcatta | tcaagaagac | caagaagccc | 1560 |
| accggcgctc ctccagaact  | gagcggagtg | accaacaatg | gcttcatccc | tcacaac    | 1617 |
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| accggctggt acaccaacgt gttcacactg cctgtgggcg acgtcgagaa tctgacatgc   | 180  |
| tctgatggcc ctagcctgat caagaccgag ctggatctgc tcaagagcgc cctgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
| ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca   | 360  |
| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480  |
| gccgtgcgcg agctgaagga cttcgtgtcc aagaacctga cacgggccat taacaagaac   | 540  |
| aagtgcgaca tcgacgacct gaagatggcc gtgtccttta gccagttcaa ccggcggttt   | 600  |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660  |
| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag   | 720  |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt   | 780  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840  |
| acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc   | 900  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020 |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc   | 1080 |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc   | 1140 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200 |
| aagcagetga acaagggetg cagetacate accaaccagg acgeegatae egtgaccate   | 1260 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320 |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgagg atcagttcca ggtggccctg   | 1380 |
| gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc   | 1440 |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg   | 1500 |
| ctgggcagct ccatgatcct ggtgtccatc ttcatcatta tcaagaagac caagaagccc   | 1560 |
| accggcgctc ctccagaact gagcggagtg accaacaatg gcttcatccc tcacaac  | 1617 |
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| accggctggt acaccaacgt gttcacactg cctgtgggcg acgtcgagaa tctgacatgc   | 180  |
| tetgatggce etageetgat caagacegag etggatetge teaagagege eetgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
|   | 2.60 |

ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca

| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
|---|------|
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480  |
| geegtgegeg agetgaagga ettegtgtee aagaacetga caegggeeat taacaagaac   | 540  |
| aagtgcgaca tegaegaeet gaagatggee gtgteettta geeagtteaa eeggeggttt   | 600  |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660  |
| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag   | 720  |
| atcaagetga tgetegagaa tagageeatg gteegaegga aaggettegg eattetgatt   | 780  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840  |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee   | 900  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020 |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc   | 1080 |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc   | 1140 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200 |
| aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc   | 1260 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320 |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgaga accagttcca ggtggccctg   | 1380 |
| gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc   | 1440 |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg   | 1500 |
| ctgggcagct ccatgatcct ggtgtccatc ttcatcatta tcaagaagac caagaagccc   | 1560 |
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| accggctggt acaccaacgt gttcacactg gaagtgggcg acgtcgagaa tctgacatgc   | 180  |
| tetgatggce etageetgat caagacegag etggatetge teaagagege eetgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
| ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca   | 360  |
| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480  |
| gccgtgcgcg agctgaagga cttcgtgtcc aagaacctga cacgggccat taacaagaac   | 540  |
| aagtgegaca tegaegaeet gaagatggee gtgteettta gecagtteaa eeggeggttt   | 600  |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660  |
| ctgatgacag atgctgaget ggctagagec gtgcctaaca tgcctacate tgccggccag   | 720  |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt   |      |
|   | 780  |

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| -continued  |              |
|---|--------------|
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840          |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee   | 900          |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960          |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020         |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc   | 1080         |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc   | 1140         |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200         |
| aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc   | 1260         |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320         |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgagg atcagttcca ggtggccctg   | 1380         |
| gaccaggtgt tegagaacat egagaattee eaggetetgg tggaccagte eaacagaate   | 1440         |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg   | 1500         |
| ctgggcagct ccatgatcct ggtgtccatc ttcatcatta tcaagaagac caagaagcc  | 1560         |
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| accggctggt acaccaacgt gttcacactg gaagtgggcg acctcgagaa tctgacatgc   | 180          |
| tctgatggcc ctagcctgat caagaccgag ctggatctga ccaagagcgc cctgagagaa   | 240          |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300          |
| ggcagctttg tgctgggagc cattgctctt ggagtggctg ctgctgcagc tgttacagca   | 360          |
| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420          |
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480          |
| gccgtgcgcg agctgaagga cttcgtgtcc aagaacctga cacgggccat taacaagaac   | 540          |
| aagtgcgaca tcgacgacct gaagatggcc gtgtccttta gccagttcaa ccggcggttt   | 600          |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660          |
| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag   | 720          |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt   | 780          |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840          |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee   | 900          |
| tacctactas dagaagacca agactaatet tatcegeeca agagegeec agtatectes  |              |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960          |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 960<br>1020  |
|   |              |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020         |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc                         | 1020<br>1080 |

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<211> LENGTH: 1617 <212> TYPE: DNA <213> ORGANISM: Ar

<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

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<213> ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

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| ctcaagaccg  | tgtctgccga                                | tcagctggcc | agagaggaac          | agatcgagaa | tcctggcagc | 300  |
|-------------|---|------------|---------------------|------------|------------|------|
| ggcagctttg  | tgctgggagc                                | cattgctctt | ggagtggctg          | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca  | tcgctaagac                                | catcagactg | gaaagcgaag          | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga  | caaacgaggc                                | cgtcagcaca | ctcggcaatg          | gcgttagagt | gctggccaca | 480  |
| geegtgegeg  | agctgaagga                                | cttcgtgctt | aagaacctgt          | ggcgggccat | taacaagaac | 540  |
| aagtgcgaca  | tcgacgacct                                | gaagatggcc | gtgtccttta          | gccagttcaa | ccggcggttt | 600  |
| ctgaacgtcg  | tgcggcagtt                                | tagcgacaac | gccggaatca          | caccagccat | cagcctggac | 660  |
| ctgatgacag  | atgctgagct                                | ggctagagcc | gtgcctaaca          | tgcctacatc | tgccggccag | 720  |
| atcaagctga  | tgctcgagaa                                | tagagccatg | gtccgacgga          | aaggettegg | cattctgatt | 780  |
| ggcgtgtacg  | gcagcagcgt                                | gatctatatg | gtgcagctgc          | ctatcttcgg | cgtgatcgac | 840  |
| acaccctgct  | ggattgtgaa                                | ggeegeteet | agctgtagcg          | agaagaaggg | caattacgcc | 900  |
| tgcctgctga  | gagaggacca                                | aggctggtat | tgtcagaacg          | ccggcagcac | cgtgtactac | 960  |
| cctaacgaga  | aggactgcga                                | gacaagaggc | gaccacgtgt          | tctgtgatac | cgccgctgga | 1020 |
| atcaatgtgg  | ccgagcagag                                | caaagagtgc | aacatcaaca          | tcagcaccac | caactatccc | 1080 |
| tgcaaggtgt  | ccaccggcag                                | gcaccctatt | tctatggtgg          | ctctgtctcc | tctgggagcc | 1140 |
| ctggtggctt  | gttataaggg                                | cgtgtcctgt | agcatcggca          | gcaacagagt | gggcatcatc | 1200 |
| aagcagctga  | acaagggctg                                | cagctacatc | accaaccagg          | acgccgatac | cgtgaccatc | 1260 |
| gacaacaccg  | tgtatcagct                                | gagcaaggtg | gaaggcgaac          | agcacgtgat | caagggcaga | 1320 |
| cctgtgtcca  | gcagcttcga                                | ccctatcaag | ttccctgagg          | atcagttcca | ggtggccctg | 1380 |
| gaccaggtgt  | tcgagaacat                                | cgagaattcc | caggetetgg          | tggaccagtc | caacagaatc | 1440 |
| ctgtctagcg  | ccgagaaggg                                | aaacaccggc | ttcatcatcg          | tgatcatcct | gatcgccgtg | 1500 |
| ctgggcagct  | ccatgatcct                                | ggtgtccatc | ttcatcatta          | tcaagaagac | caagaagccc | 1560 |
| accggcgctc  | ctccagaact                                | gagcggagtg | accaacaatg          | gcttcatccc | tcacaac    | 1617 |
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| gagagctacc  | tggaagagtc                                | ctgcagcacc | atcacagagg          | gctacctgtc | tgtgctgaga | 120  |
| accggctggt  | acaccaacgt                                | gttcacactg | cctgtgggcg          | acgtcgagaa | tctgacatgc | 180  |
| tctgatggcc  | ctagcctgat                                | caagaccgag | ctggatctga          | ccaagagcgc | cctgagagaa | 240  |
| ctcaagaccg  | tgtctgccga                                | tcagctggcc | agagaggaac          | agatcgagaa | tcctggcagc | 300  |
| ggcagctttg  | tgctgggagc                                | cattgctctt | ggagtggctg          | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca  | tcgctaagac                                | catcagactg | gaaagcgaag          | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga  | caaacgaggc                                | cgtcagcaca | ctcggcaatg          | gcgttagagt | gctggccaca | 480  |
| gccgtgcgcg  | agctgaagga                                | cttcgtgtcc | aagaacctga          | cacgggccat | taacaagaac | 540  |
| aagtgcgaca  | tcgacgacct                                | gaagatggcc | gtgtccttta          | gccagttcaa | ccggcggttt | 600  |
| -           |   |            |                     |            | •          |      |

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| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag  | 720   |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt  | 780   |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac  | 840   |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee  | 900   |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac  | 960   |
| cetaacgaga aggaetgega gaeaagagge gaeeaegtgt tetgtgatae egeegetgga  | 1020  |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc  | 1080  |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc  | 1140  |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc  | 1200  |
| aagcagetga acaagggetg cagetacate accaaccagg acgeegatac egtgaccate  | 1260  |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga  | 1320  |
| octgtgtoca gcagottoga occtatoaag ttooctgagg atcagttoca ggtggcoctg  | 1380  |
| gaccaggtgt tegagaacat egagaattee caggetetgg tggaccagte caacagaate  | 1440  |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg  | 1500  |
| ctgggcagct ccatgatect ggtgtccate tteateatta teaagaagae caagaageee  | 1560  |
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| Cloop Blgomed. 120   |   |
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|  | 60<br>120   |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa  |   |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa<br>gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga   | 120   |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge  | 120<br>180  |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ceaagagege eetgagagaa  | 120<br>180<br>240   |
| atgagetgga aggtggteat catetteage etgetgatea eaceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagageg tgtetgeega teagetggee agagaggaac agategagaa teetggeage  | 120<br>180<br>240<br>300                                    |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea  | 120<br>180<br>240<br>300<br>360                             |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagageg tgtetgeega teagetggee aggaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae cateagaetg gaaagegaag tgacegeeat caacaaegee   | 120<br>180<br>240<br>300<br>360<br>420                      |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeca tegetaagac cateagactg gaaagegaag tgacegecat caacaacgee etgaagaaga caaacgagge egteageaca eteggeaatg gegttagagt getggecaca  | 120<br>180<br>240<br>300<br>360<br>420                      |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgaagaa etceaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage cattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagaceg gaaagegaag tgacegeeat eaacaacgee etgaagaaga caaacgage egteageaca eteggeaatg gegttagagt getggeeaca geegtgeege agetgaagga ettegtgtee aagaacetga eaceggeeat taacaagaac geegtgeege agetgaagga ettegtgtee aagaacetga eaceggeeat taacaagaac   | 120<br>180<br>240<br>300<br>360<br>420<br>480               |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etagaetgat caagaeegag etggatetga ecaagagege eetgagagaa etcaagaeeg ggeagetttg tgetgeega teagetggee agagaggaae agategagaa teetggeage ggeagetttg tgetggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagaetg gaaagegaag tgacegeeat eaacaaegee etgaagaaga eaaaegagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgeege agetgaagga ettegtgtee aagaaeetga eaegggeeat taacaagaac aagtgegaca teeetgacet gaagatggee gtgteettta geeagtteaa eeggeggttt   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540        |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etagategge etggatetga ecaagagege eetgagagaa eteagagagaa eteagagagaa eteagagagaa eteagagagaa eteagagagaa eteagagagaa eteagagagaa eteagagagaa eteagagaga etggatetga eagageggegggegggeggttt tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eatcagaetg gaaagegaag tgacegeeat eaacaacgee etgaagaagaa eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgegge agetgaagga ettegtgtee aagaacetga eacgggeeat taacaagaacaagtgegaaca teeetgacet gaagatggee gtgteettta geeagtteaa eeggeggttt etgaacgteg tgeggeagtt tagegacaac geeggaatea eaccageeat eageetggae  | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600 |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteegagagaa eteegagagaa eteegagagaa eteegagagaa eteegagagaa eteegagagaa eteegagagaa eteegagagaa etageegageeg   | 120 180 240 300 360 420 480 540 600 660                     |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etceaagaceg tgtetgeega teagetggee aggagggaac agategagaa teetggeage ggeagetttg tgetgggage cattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagaceg gaaagegaag tgacegeeat caacaacegee etgaagaaa eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca etgaagaaga etteetggeage etgaagaaga geegtgeege agetgaagga etteetgtee aagaacetga eaceggeeat taacaagaac aagtgegaca teeetgacet gaagatggee gtgteettta geeagtteaa eeggeggttt etgaacgteg tgeggeagtt tagegacaac geeggaatea eaceageeat eageetggac etgatgacag tgeetgagee tagegacaga etgatgaee gtgeetaaca tgeetacate tgeeggeeag ateaagetga tgetegagaa tagageeag gtgeetaaca aaggettegg eattetgatt  | 120 180 240 300 360 420 480 540 600 660 720                 |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etagaegg etggatetga ecaagagege eetgagagaa eteegagagaa eteegagagagagegagegegegegegegegegegegege  | 120 180 240 300 360 420 480 540 600 660 720 780             |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcegagagaa etggagegegggegggegggegggegggggggggg  | 120 180 240 300 360 420 480 540 600 660 720 780 840         |
| atgagetgga aggtggteat catetteage etgetgatea caceteagea eggeetgaaa gagaagetace tggaaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgg gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etagetgat caagacegag etggatetga ecaagagege ectgagagaa etctgagagaa etcaagaceg tgtetgeega teagetggee aggagggaac agategagaa teetggeage ggeagetttg tgetgggage cattgetett ggagtggetg etgetgage tgttacagea ggegtggeea tegetaagae eateagactg gaaagegaag tgacegeeat eaacaacgee etgaagaaa etcetggagae etgaagaag etgaaggaag tgacegeeat eaacaacgee etgaagaaga etgetgagea etgetaagae ettegtgee aagaacetga gegttagagt getggeeaca geegtgegeg agetgaagga ettegtgtee aagaacetga eacgggeeat taacaagaac aagtgegaca teeetgacet gaagatggee gtgteettta geeagtteaa eeggeggttt etgaacgteg tgeggeagtt tageggacaac geeggaatea eacacageeat eageetggae etgatgacag atgetgaget ggetagagee gtgeetaaca tgeetacate tgeeggeeag ateaagetga tgetegagaa tagageeatg gteegaacga aaggettegg eattetgatt ggegtgtaceg geageagetg gatetatatg gtgeagetge etatettegg egtgategae acaceetget ggattgtaa ggeegeteet agetgagee etatettegg egtgategae acaceetget ggattgtgaa ggeegeteet agetgageg agaagaaggg caattaegee tgeetgeetga gagaggacea aggeetget tgeetgetga egtgategae egtgategae egtgetgetge gagaggagae egtgategae acaceetget ggattgtgaa ggeegeteet agetgageg agaagaaggg caattaegee tgeetgetga gagaggacea aggeetgeteet agetgagea egtgeageae egtgatetae | 120 180 240 300 360 420 480 540 600 660 720 780 840 900     |

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| ctggtggctt  | gttataaggg                                | cgtgtcctgt | agcatcggca          | gcaacagagt | gggcatcatc | 1200 |
|-------------|---|------------|---------------------|------------|------------|------|
| aagcagctga  | acaagggctg                                | cagctacatc | accaaccagg          | acgccgatac | cgtgaccatc | 1260 |
| gacaacaccg  | tgtatcagct                                | gagcaaggtg | gaaggcgaac          | agcacgtgat | caagggcaga | 1320 |
| cctgtgtcca  | gcagcttcga                                | ccctatcaag | ttccctgagg          | atcagttcca | ggtggccctg | 1380 |
| gaccaggtgt  | tcgagaacat                                | cgagaattcc | caggetetgg          | tggaccagtc | caacagaatc | 1440 |
| ctgtctagcg  | ccgagaaggg                                | aaacaccggc | ttcatcatcg          | tgatcatcct | gatcgccgtg | 1500 |
| ctgggcagct  | ccatgatcct                                | ggtgtccatc | ttcatcatta          | tcaagaagac | caagaagccc | 1560 |
| accggcgctc  | ctccagaact                                | gagcggagtg | accaacaatg          | gcttcatccc | tcacaac    | 1617 |
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| gagagctacc  | tggaagagtc                                | ctgcagcacc | atcacagagg          | gctacctgtc | tgtgctgaga | 120  |
| accggctggt  | acaccaacgt                                | gttcacactg | gaagtgggcg          | acgtcgagaa | tctgacatgc | 180  |
| tctgatggcc  | ctagcctgat                                | caagaccgag | ctggatctga          | ccaagagcgc | cctgagagaa | 240  |
| ctcaagaccg  | tgtctgccga                                | tcagctggcc | agagaggaac          | agatcgagaa | tcctggcagc | 300  |
| ggcagctttg  | tgctgggagc                                | cattgctctt | ggagtggctg          | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca  | tcgctaagac                                | catcagactg | gaaagcgaag          | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga  | caaacgaggc                                | cgtcagcaca | ctcggcaatg          | gcgttagagt | gctggccaca | 480  |
| gccgtgcgcg  | agctgaagga                                | cttcgtgtcc | aagaacctga          | cacgggccat | taacaagaac | 540  |
| aagtgcccta  | tcgacgacct                                | gaagatggcc | gtgtccttta          | gccagttcaa | ccggcggttt | 600  |
| ctgaacgtcg  | tgcggcagtt                                | tagcgacaac | gccggaatca          | caccagccat | cagcctggac | 660  |
| ctgatgacag  | atgctgagct                                | ggctagagcc | gtgcctaaca          | tgcctacatc | tgccggccag | 720  |
| atcaagctga  | tgctcgagaa                                | tagagccatg | gtccgacgga          | aaggcttcgg | cattctgatt | 780  |
| ggcgtgtacg  | gcagcagcgt                                | gatctatatg | gtgcagctgc          | ctatcttcgg | cgtgatcgac | 840  |
| acaccctgct  | ggattgtgaa                                | ggccgctcct | agctgtagcg          | agaagaaggg | caattacgcc | 900  |
| tgcctgctga  | gagaggacca                                | aggctggtat | tgtcagaacg          | ccggcagcac | cgtgtactac | 960  |
| cctaacgaga  | aggactgcga                                | gacaagaggc | gaccacgtgt          | tctgtgatac | cgccgctgga | 1020 |
| atcaatgtgg  | ccgagcagag                                | caaagagtgc | aacatcaaca          | tcagcaccac | caactatccc | 1080 |
| tgcaaggtgt  | ccaccggcag                                | gcaccctatt | tctatggtgg          | ctctgtctcc | tctgggagcc | 1140 |
| ctggtggctt  | gttataaggg                                | cgtgtcctgt | agcatcggca          | gcaacagagt | gggcatcatc | 1200 |
| aagcagctga  | acaagggctg                                | cagctacatc | accaaccagg          | acgccgatac | cgtgaccatc | 1260 |
| gacaacaccg  | tgtatcagct                                | gagcaaggtg | gaaggcgaac          | agcacgtgat | caagggcaga | 1320 |
| cctgtgtcca  | gcagcttcga                                | ccctatcaag | ttccctgagg          | atcagttcca | ggtggccctg | 1380 |
| gaccaggtgt  | tcgagaacat                                | cgagaattcc | caggctctgg          | tggaccagtc | caacagaatc | 1440 |
| ctgtctagcg  | ccgagaaggg                                | aaacaccggc | ttcatcatcg          | tgatcatcct | gatcgccgtg | 1500 |
|             |   |            |                     |            |            |      |

ctgggcagct ccatgatect ggtgtecate tteateatta teaagaagae caagaagee 1560

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                                                                     480
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aagtgcgaca tcgacgacct gaagatggcc gtgtccttta gccagttcaa ccggcggttt
                                                                     600
ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac
                                                                     660
                                                                     720
ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccgqccag
                                                                     780
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ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac
                                                                     840
acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee
                                                                     900
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                                                                     960
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                                                                    1020
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tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc
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                                                                    1200
aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc
                                                                    1260
gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga
                                                                    1320
cctgtgtcca gcagcttcga ccctatcaag ttccctgagg atcagttcca ggtggccctg
                                                                    1380
gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc
                                                                    1440
ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg
                                                                    1500
ctgggcagct ccatgatcct ggtgtccatc ttcatcatta tcaagaagac caagaagcc
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| accggctggt | acaccaacgt  | gttcacactg                   | gaagtgggcg | acgtcgagaa | tetgacatge | 180  |
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| tctgatggcc | ctagcctgat  | caagaccgag                   | ctggatctga | ccaagagcgc | cctgagagaa | 240  |
| ctcaagaccg | tgtctgccga  | tcagctggcc                   | agagaggaac | agatcgagaa | tcctggcagc | 300  |
| ggcagctttg | tgctgggagc  | cattgctctt                   | ggagtggctg | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca | tegetaagae  | catcagactg                   | gaaagcgaag | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga | caaacgaggc  | cgtcagcaca                   | ctcggcaatg | gcgttagagt | gctggccaca | 480  |
| gccgtgcgcg | agctgaagga  | cttcgtgtcc                   | aagaacctga | cacgggccat | taacaagaac | 540  |
| aagtgcgaca | tegaegaeet  | gaagatggcc                   | gtgtccttta | gccagttcaa | ccggcggttt | 600  |
| ctgaacgtcg | tgcggcagtt  | tagcgacaac                   | gccggaatca | caccagccat | cagcctggac | 660  |
| ctgatgacag | atgctgagct  | ggctagagcc                   | gtgcctaaca | tgcctacatc | tgccggccag | 720  |
| atcaagctga | tgctcgagaa  | tagagccatg                   | gtccgacgga | aaggettegg | cattctgatt | 780  |
| ggcgtgtacg | gcagcagcgt  | gatctatatg                   | gtgcagctgc | ctatcttcgg | cgtgatcgac | 840  |
| acaccctgct | ggattgtgaa  | ggccgctcct                   | agctgtagcg | agaagaaggg | caattacgcc | 900  |
| tgcctgctga | gagaggacca  | aggctggtat                   | tgtcagaacg | ccggcagcac | cgtgtactac | 960  |
| cctaacgaga | aggactgcga  | gacaagaggc                   | gaccacgtgt | tctgtgatac | cgccgctgga | 1020 |
| atcaatgtgg | ccgagcagag  | caaagagtgc                   | aacatcaaca | tcagcaccac | caactatccc | 1080 |
| tgcaaggtgt | ccaccggcag  | gcaccctatt                   | tctatggtgg | ctctgtctcc | tctgggagcc | 1140 |
| ctggtggctt | gttataaggg  | cgtgtcctgt                   | agcatcggca | gcaacagagt | gggcatcatc | 1200 |
| aagcagctga | acaagggctg  | cagctacatc                   | accaaccagg | acgccgatac | cgtgaccatc | 1260 |
| gacaacaccg | tgtatcagct  | gagcaaggtg                   | gaaggcgaac | agcacgtgat | caagggcaga | 1320 |
| cctgtgtcca | gcagcttccc  | acctatcaag                   | ttccctgagg | atcagttcca | ggtggccctg | 1380 |
| gaccaggtgt | tcgagaacat  | cgagaattcc                   | caggetetgg | tggaccagtc | caacagaatc | 1440 |
| ctgtctagcg | ccgagaaggg  | aaacaccggc                   | ttcatcatcg | tgatcatcct | gatcgccgtg | 1500 |
| ctgggcagct | ccatgatcct  | ggtgtccatc                   | ttcatcatta | tcaagaagac | caagaagccc | 1560 |
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| gagagetace | tggaagagtc  | ctgcagcacc                   | atcacagagg | gctacctgtc | tgtgctgaga | 120  |
| accggctggt | acaccaacgt  | gttcacactg                   | gaagtgggcg | acgtcgagaa | tctgacatgc | 180  |
| tctgatggcc | ctagcctgat  | caagaccgag                   | ctggatctga | ccaagagcgc | cctgagagaa | 240  |
| ctcaagaccg | tgtctgccga  | tcagctggcc                   | agagaggaac | agatcgagaa | tcctggcagc | 300  |
| ggcagctttg | tgctgggagc  | cattgctctt                   | ggagtggctg | ctgctgcagc | tgttacagca | 360  |
| ggcgtggcca | tcgctaagac  | catcagactg                   | gaaagcgaag | tgaccgccat | caacaacgcc | 420  |
| ctgaagaaga | caaacgaggc  | cgtcagcaca                   | ctcggcaatg | gcgttagagt | gctggccaca | 480  |
|            |   |                              |            |            |            |      |

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| aagtgegaca tegaegacet gaagatggee gtgteettta geeagtteaa eeggeggttt  | 600  |
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| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag  | 720  |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt  | 780  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac  | 840  |
| acaccetget ggattgtgaa ggeegeteet agetgtageg agaagaaggg caattaegee  | 900  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac  | 960  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga  | 1020   |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc  | 1080   |
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc  | 1140   |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc  | 1200   |
| aagcagctga acaagggctg cagctacatc accaaccagg acgccgatac cgtgaccatc  | 1260   |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga  | 1320   |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgaga accagttcca ggtggccctg  | 1380   |
| gaccaggtgt tcgagaacat cgagaattcc caggctctgg tggaccagtc caacagaatc  | 1440   |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg  | 1500   |
| ctgggcagct ccatgatcct ggtgtccatc ttcatcatta tcaagaagac caagaagcc   | 1560   |
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|  |  |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga  | 120  |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga<br>aceggetggt acaccaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge   | 120<br>180   |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga<br>aceggetggt acaccaacgt gttcacactg gaagtgggeg acgtegagaa tetgacatge<br>tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa  | 120<br>180<br>240  |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage  | 120<br>180<br>240<br>300   |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea  | 120<br>180<br>240<br>300<br>360                                    |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaccaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagactg gaaagegaag tgacegeeat eaacaacgee  | 120<br>180<br>240<br>300<br>360<br>420                             |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagaceg tgtetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagac eateagactg gaaagegaag tgacegeeat eaacaacgee etgaagaaga eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca  | 120<br>180<br>240<br>300<br>360<br>420                             |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaccaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa etcaagagege ggeagetttg tgetgeega teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagac eateagactg gaaagegaag tgacegeeat eaacaacgee etgaagaag eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgeege agetgaagga ettegtgtee aagaacetga eacgggeeat taacaagaac  | 120<br>180<br>240<br>300<br>360<br>420<br>480                      |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteeagagaga eteagagaga agategagaa teetggeage ggeagetttg tgetggegag eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagaetg gaaagegaag tgacegeeat eaacaaegee etgaagaaga eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgegeg agetgaagga ettegtgtee aagaacetga eacgggeeat taacaagaac aagtgegaca tegacgacet gaagatggee gtgteettta geeagtteaa eeggeggttt   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600        |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteeagagaga eteeagagaga etgagagaa teetgagagaa eteeagagaga etgageetggeggeggeggetggeeggeggeggeggeggeggeg   | 120<br>180<br>240<br>300<br>360<br>420<br>480<br>540<br>600<br>660 |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteeagagaga eteeagagae etgagetgge etgetgagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgaga tgttacagea ggeggtggeca tegetaagae eateagaetg gaaagegaag tgacegeat eaacaaegee etgaagaaga eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgegeg agetgaagga ettegtgtee aagaacetga eacgggeeat taacaagaac aagtgegaca tegacgacet gaagatggee gtgteettta geeagteaa eeggeggttt etgaacgteg tgeggeagtt tagegacaac geeggaatea eaceageeat eageetggae etgatgacaga atgetgaget ggetagagee gtgeetaaca tegeetagae etgatgaea etgatgaea ggetgaagee gtgeetaaca tgeetacate tgeeggeeage etgatgaeag atgetgaget ggetagagee gtgeetaaca tgeetacate tgeeggeeag   | 120 180 240 300 360 420 480 540 600 660                            |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaccaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteeagagege ggeagetttg tgetggegag teagetggee agagaggaac agategagaa teetggeage ggeagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea ggegtggeea tegetaagae eateagaetg gaaagegaag tgacegeeat eaacaacgee etgaagagaa eaaacgagge egteageaca eteggeaatg gegttagagt getggeeaca geegtgegeg agetgaagga ettegtgee aagaacetga eacegggeeat taacaagaac aagtgegaca tegacgaect gaagatggee gtgteettta geeagtteaa eeggeggttt etgaacgteg tgeeggaate tageagaece etgaagaega tegacegeat eageetggae etgaacgteg tgeeggaate tageagaece gtgteettta geeagtteaa eeggeggttt etgaacgteg tgeeggaatt tagegacaac geeggaatea eaceageeat eageetggae etgatgaeagae etgatgaeaga tegetgagae tegetgagae etgaagaea tegaegaea tageetgagae etgaagaea tagageeaga atgeetagaea tagageeaga atgeetgagaa tagageeaga gteegaegga aaggettegg eattetgatt | 120 180 240 300 360 420 480 540 600 660 720                        |
| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga aceggetggt acaceaacgt gtteacactg gaagtgggeg acgtegagaa tetgacatge tetgatggee etageetgat caagacegag etggatetga ecaagagege eetgagagaa eteeagagaga eteeagagagagegeggeggeggeggeggeggeggegggggg   | 120 180 240 300 360 420 480 540 600 660 720 780                    |

cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga 1020

| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc   | 1080 |
|---|------|
| tgcaaggtgt ccaccggcag gcaccctatt tctatggtgg ctctgtctcc tctgggagcc   | 1140 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200 |
| aagcagetga acaagggetg cagetacate accaaccagg acgeegatac egtgaccate   | 1260 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320 |
| cetgtgteca geagettega ceetateaag tteeeteagg ateagtteca ggtggeeetg   | 1380 |
| gaccaggtgt togagaacat ogagaattoo caggototgg tggaccagto caacagaato   | 1440 |
| ctgtctagcg ccgagaaggg aaacaccggc ttcatcatcg tgatcatcct gatcgccgtg   | 1500 |
| ctgggcagct ccatgatect ggtgtccate ttcateatta tcaagaagae caagaageee   | 1560 |
| accggcgctc ctccagaact gagcggagtg accaacaatg gcttcatccc tcacaac  | 1617 |
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| gagagetace tggaagagte etgeageace ateacagagg getacetgte tgtgetgaga   | 120  |
| accggctggt acaccaacgt gttcacactg gaagtgggcg acgtcgagaa tctgacatgc   | 180  |
| tetgatggcc etageetgat caagacegag etggatetga eeaagagege eetgagagaa   | 240  |
| ctcaagaccg tgtctgccga tcagctggcc agagaggaac agatcgagaa tcctggcagc   | 300  |
| ggcagetttg tgetgggage eattgetett ggagtggetg etgetgeage tgttacagea   | 360  |
| ggcgtggcca tcgctaagac catcagactg gaaagcgaag tgaccgccat caacaacgcc   | 420  |
| ctgaagaaga caaacgaggc cgtcagcaca ctcggcaatg gcgttagagt gctggccaca   | 480  |
| geogtgegeg agetgaagga ettegtgtee aagaaeetga eaegggeeat taacaagaae   | 540  |
| aagtgegaca tegaegaeet gaagatggee gtgteettta geeagtggaa eeggeggttt   | 600  |
| ctgaacgtcg tgcggcagtt tagcgacaac gccggaatca caccagccat cagcctggac   | 660  |
| ctgatgacag atgctgagct ggctagagcc gtgcctaaca tgcctacatc tgccggccag   | 720  |
| atcaagctga tgctcgagaa tagagccatg gtccgacgga aaggcttcgg cattctgatt   | 780  |
| ggcgtgtacg gcagcagcgt gatctatatg gtgcagctgc ctatcttcgg cgtgatcgac   | 840  |
| acaccctgct ggattgtgaa ggccgctcct agctgtagcg agaagaaggg caattacgcc   | 900  |
| tgcctgctga gagaggacca aggctggtat tgtcagaacg ccggcagcac cgtgtactac   | 960  |
| cctaacgaga aggactgcga gacaagaggc gaccacgtgt tctgtgatac cgccgctgga   | 1020 |
| atcaatgtgg ccgagcagag caaagagtgc aacatcaaca tcagcaccac caactatccc   | 1080 |
| tgcaaggtgt ccaceggcag gcacectatt tetatggtgg etetgtetee tetgggagee   | 1140 |
| ctggtggctt gttataaggg cgtgtcctgt agcatcggca gcaacagagt gggcatcatc   | 1200 |
| aagcagetga acaagggetg cagetacate accaaccagg acgeegatac egtgaccate   | 1260 |
| gacaacaccg tgtatcagct gagcaaggtg gaaggcgaac agcacgtgat caagggcaga   | 1320 |
| cctgtgtcca gcagcttcga ccctatcaag ttccctgagg atcagttcca ggtggccctg   | 1380 |
| gaccaggtgt togagaacat ogagaattoo caggototgg tggaccagto caacagaato   | 1440 |
|   |      |

|             |   |                              |            | -0011011   |            |      |  |
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| ctgtctagcg  | ccgagaaggg                                | aaacaccggc                   | ttcatcatcg | tgatcatcct | gatcgccgtg | 1500 |  |
| ctgggcagct  | ccatgatcct                                | ggtgtccatc                   | ttcatcatta | tcaagaagac | caagaagccc | 1560 |  |
| accggcgctc  | ctccagaact                                | gagcggagtg                   | accaacaatg | gcttcatccc | tcacaac    | 1617 |  |
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| gagagcuacc  | uggaagaguc                                | cugcagcacc                   | aucacagagg | gcuaccuguc | ugugcugaga | 120  |  |
| accggcuggu  | acaccaacgu                                | guucacacug                   | gaagugggcg | acgucgagaa | ucugacaugc | 180  |  |
| ucugauggcc  | cuagecugau                                | caagaccgag                   | cuggaucuga | ccaagagcgc | ccugagagaa | 240  |  |
| cucaagaccg  | ugucugccga                                | ucagcuggcc                   | agagaggaac | agaucgagaa | uccuggcagc | 300  |  |
| ggcagcuuug  | ugcugggagc                                | cauugcucuu                   | ggaguggcug | cugcugcagc | uguuacagca | 360  |  |
| ggcguggcca  | ucugcaagac                                | caucagacug                   | gaaagcgaag | ugaccgccau | caacaacgcc | 420  |  |
| cugaagaaga  | caaacgaggc                                | cgucagcaca                   | cucggcaaug | gcguuagagu | gcuggccuuu | 480  |  |
| gccgugcgcg  | agcugaagga                                | cuucgugucc                   | aagaaccuga | cacgggcccu | gaacaagaac | 540  |  |
| aagugcgaca  | ucgacgaccu                                | gaagauggcc                   | guguccuuua | gccaguucaa | ccggcgguuu | 600  |  |
| cugaacgucg  | ugcggcaguu                                | uagcgacaac                   | gccggaauca | caccagccau | cagccuggac | 660  |  |
| cugaugacag  | augcugagcu                                | ggcuagagcc                   | gugccuaaca | ugccuacauc | ugccggccag | 720  |  |
| aucaagcuga  | ugcucgagaa                                | uagagccaug                   | guccgacgga | aaggcuucgg | cauucugugu | 780  |  |
| ggcguguacg  | gcagcagcgu                                | gaucuauaug                   | gugcagcugc | cuaucuucgg | cgugaucgac | 840  |  |
| acacccugcu  | ggauugugaa                                | ggccgcuccu                   | agcuguagcg | agaagaaggg | caauuacgcc | 900  |  |
| ugccugcuga  | gagaggacca                                | aggcugguau                   | ugucagaacg | ccggcagcac | cguguacuac | 960  |  |
| ccuaacgaga  | aggacugcga                                | gacaagaggc                   | gaccacgugu | ucugugauac | cgccgcugga | 1020 |  |
| aucaaugugg  | ccgagcagag                                | caaagagugc                   | aacaucaaca | ucagcaccac | caacuauccc | 1080 |  |
| ugcaaggugu  | ccaccggcag                                | gcacccuauu                   | ucuauggugg | cucugucucc | ucugggagcc | 1140 |  |
| cugguggcuu  | guuauaaggg                                | cguguccugu                   | agcaucggca | gcaacagagu | gggcaucauc | 1200 |  |
| aagcagcuga  | acaagggcug                                | cagcuacauc                   | accaaccagg | acgccgauac | cgugaccauc | 1260 |  |
| gacaacaccg  | uguaucagcu                                | gagcaaggug                   | gaaggcgaac | agcacgugau | caagggcaga | 1320 |  |
| ccugugucca  | gcagcuucga                                | cccuaucaag                   | uucccugagg | aucaguucaa | cguggcccug | 1380 |  |
| gaccaggugu  | ucgagaacau                                | cgagaauucc                   | caggcucugg | uggaccaguc | caacagaauc | 1440 |  |
| cugucuagcg  | ccgagaaggg                                | aaacaccggc                   | uucaucaucg | ugaucauccu | gaucgccgug | 1500 |  |
| cugggcagcu  | ccaugauccu                                | gguguccauc                   | uucaucauua | ucaagaagac | caagaagccc | 1560 |  |
| accggcgcuc  | cuccagaacu                                | gagcggagug                   | accaacaaug | gcuucauccc | ucacaac    | 1617 |  |
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<sup>&</sup>lt;211> LENGTH: 1617 <212> TYPE: RNA

<sup>&</sup>lt;213> ORGANISM: Artificial Sequence <220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Synthetic Polynucleotide

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| gagagcuacc  | uggaagaguc                                | cugcagcacc                   | aucacagagg | gcuaccuguc | ugugcugaga | 120  |
| accggcuggu  | acaccaacgu                                | guucacacug                   | gaagugggcg | acgucgagaa | ucugacaugc | 180  |
| ucugauggcc  | cuagccugau                                | caagaccgag                   | cuggaucuga | ccaagagcgc | ccugagagaa | 240  |
| cucaagaccg  | ugucugccga                                | ucagcuggcc                   | agagaggaac | agaucgagaa | uccuggcagc | 300  |
| ggcagcuuug  | ugcugggagc                                | cauugcucuu                   | ggaguggcug | cugcugcagc | uguuacagca | 360  |
| ggcguggcca  | ucugcaagac                                | caucagacug                   | gaaagcgaag | ugaccgccau | caacaacgcc | 420  |
| cugaagaaga  | caaacgaggc                                | cgucagcaca                   | cucggcaaug | gcguuagagu | gcuggccaca | 480  |
| geegugegeg  | agcugaagga                                | cuucgugucc                   | aagaaccuga | cacgggccau | uaacaagaac | 540  |
| aagugcgaca  | ucgacgaccu                                | gaagauggcc                   | guguccuuua | gccaguucaa | ccggcgguuu | 600  |
| cugaacgucg  | ugcggcaguu                                | uagcgacaac                   | gccggaauca | caccagccau | cagccuggac | 660  |
| cugaugacag  | augcugagcu                                | ggcuagagcc                   | gugccuaaca | ugccuacauc | ugccggccag | 720  |
| aucaagcuga  | ugcucgagaa                                | uagagccaug                   | guccgacgga | aaggcuucgg | cauucugugu | 780  |
| ggcguguacg  | gcagcagcgu                                | gaucuauaug                   | gugcagcugc | cuaucuucgg | cgugaucgac | 840  |
| acacccugcu  | ggauugugaa                                | ggeegeueeu                   | agcuguagcg | agaagaaggg | caauuacgcc | 900  |
| ugccugcuga  | gagaggacca                                | aggcugguau                   | ugucagaacg | ccggcagcac | cguguacuac | 960  |
| ccuaacgaga  | aggacugcga                                | gacaagaggc                   | gaccacgugu | ucugugauac | cgccgcugga | 1020 |
| aucaaugugg  | ccgagcagag                                | caaagagugc                   | aacaucaaca | ucagcaccac | caacuauccc | 1080 |
| ugcaaggugu  | ccaccggcag                                | gcacccuauu                   | ucuauggugg | cucugucucc | ucugggagcc | 1140 |
| cugguggcuu  | guuauaaggg                                | cguguccugu                   | agcaucggca | gcaacagagu | gggcaucauc | 1200 |
| aagcagcuga  | acaagggcug                                | cagcuacauc                   | accaaccagg | acgccgauac | cgugaccauc | 1260 |
| gacaacaccg  | uguaucagcu                                | gagcaaggug                   | gaaggcgaac | agcacgugau | caagggcaga | 1320 |
| ccugugucca  | gcagcuucga                                | cccuaucaag                   | uucccugagc | accaguggca | uguggcccug | 1380 |
| gaccaggugu  | ucgagaacau                                | cgagaauucc                   | caggcucugg | uggaccaguc | caacagaauc | 1440 |
| cugucuagcg  | ccgagaaggg                                | aaacaccggc                   | uucaucaucg | ugaucauccu | gaucgccgug | 1500 |
| cugggcagcu  | ccaugauccu                                | gguguccauc                   | uucaucauua | ucaagaagac | caagaagccc | 1560 |
| accggcgcuc  | cuccagaacu                                | gagcggagug                   | accaacaaug | gcuucauccc | ucacaac    | 1617 |
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| gagagcuacc  | uggaagaguc                                | cugcagcacc                   | aucacagagg | gcuaccuguc | ugugcugaga | 120  |
| accggcuggu  | acaccaacgu                                | guucacacug                   | gaagugggcg | acgucgagaa | ucugacaugc | 180  |
| ucugauggcc  | cuagccugau                                | caagaccgag                   | cuggaucugc | ucaagagcgc | ccugagagaa | 240  |
| cucaagaccg  | ugucugccga                                | ucagcuggcc                   | agagaggaac | agaucgagaa | uccuggcagc | 300  |
| ggcagcuuug  | ugcugggagc                                | cauugcucuu                   | ggaguggcug | cugcugcagc | uguuacagca | 360  |

ggcguggcca ucgcuaagac caucagacug gaaagcgaag ugaccgccau caacaacgcc

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|                   |                  |            | -contir    | nued       |      |  |
|-------------------|------------------|------------|------------|------------|------|--|
| cugaagaaga caaac  | Jaggc cgucagcaca | cucggcaaug | gcguuagagu | gcuggccaca | 480  |  |
| gccgugcgcg agcuga | aagga cuucgugucc | aagaaccuga | cacgggccau | uaacaagaac | 540  |  |
| aagugegaea ueeeuq | gaccu gaagauggcc | guguccuuua | gccaguucaa | ccggcgguuu | 600  |  |
| cugaacgucg ugcgg  | aguu uagcgacaac  | gccggaauca | caccagccau | cagccuggac | 660  |  |
| cugaugacag augcu  | gagcu ggcuagagcc | gugccuaaca | ugccuacauc | ugccggccag | 720  |  |
| aucaagcuga ugcuc  | gagaa uagagccaug | guccgacgga | aaggcuucgg | cauucugauu | 780  |  |
| ggcguguacg gcagca | agcgu gaucuauaug | gugcagcugc | cuaucuucgg | cgugaucgac | 840  |  |
| acacccugcu ggauu  | jugaa ggccgcuccu | agcuguagcg | agaagaaggg | caauuacgcc | 900  |  |
| ugccugcuga gagag  | jacca aggcugguau | ugucagaacg | ccggcagcac | cguguacuac | 960  |  |
| ccuaacgaga aggacı | ıgcga gacaagaggc | gaccacgugu | ucugugauac | cgccgcugga | 1020 |  |
| aucaaugugg ccgago | agag caaagagugc  | aacaucaaca | ucagcaccac | caacuauccc | 1080 |  |
| ugcaaggugu ccacc  | ggcag gcacccuauu | ucuauggugg | cucugucucc | ucugggagcc | 1140 |  |
| cugguggcuu guuau  | aaggg cguguccugu | agcaucggca | gcaacagagu | gggcaucauc | 1200 |  |
| aagcagcuga acaag  | ggcug cagcuacauc | accaaccagg | acgccgauac | cgugaccauc | 1260 |  |
| gacaacaccg uguau  | agcu gagcaaggug  | gaaggcgaac | agcacgugau | caagggcaga | 1320 |  |
| ccugugucca gcagc  | ucga cccuaucaag  | uucccugagg | aucaguucca | gguggcccug | 1380 |  |
| gaccaggugu ucgaga | aacau cgagaauucc | caggcucugg | uggaccaguc | caacagaauc | 1440 |  |
| cugucuageg eegaga | aaggg aaacaccggc | uucaucaucg | ugaucauccu | gaucgccgug | 1500 |  |
| cugggcagcu ccauga | auccu gguguccauc | uucaucauua | ucaagaagac | caagaagccc | 1560 |  |
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| accggcuggu acacca | acgu guucacacug  | gaagugggcg | acgucgagaa | ucugacaugc | 180  |  |
| ucugauggcc cuagco | ugau caagaccgag  | cuggaucugc | ucaagagcgc | ccugagagaa | 240  |  |
| cucaagaccg ugucu  | geega ueageuggee | agagaggaac | agaucgagaa | uccuggcagc | 300  |  |
| ggcagcuuug ugcug  | ggagc cauugcucuu | ggaguggcug | cugcugcagc | uguuacagca | 360  |  |
| ggcguggcca ucgcua | agac caucagacug  | gaaagcgaag | ugaccgccau | caacaacgcc | 420  |  |
| cugaagaaga caaac  | gagge egueageaea | cucggcaaug | gcguuagagu | gcuggccaca | 480  |  |
| gccgugcgcg agcuga | agga cuucgugucc  | aagaaccuga | cacgggccau | uaacaagaac | 540  |  |
| aagugegaea ueeeug | gaccu gaagauggcc | guguccuuua | gccaguucaa | ccggcgguuu | 600  |  |
| cugaacgucg ugcgg  | caguu uagcgacaac | gccggaauca | caccagccau | cagccuggac | 660  |  |
| cugaugacag augcu  | gagcu ggcuagagcc | gugccuaaca | ugccuacauc | ugccggccag | 720  |  |
| aucaagcuga ugcuc  | gagaa uagagccaug | guccgacgga | aaggcuucgg | cauucugauu | 780  |  |
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840

900

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| ugccugcuga gagaggacca aggcugguau ugucagaacg ccggcagcac cguguacuac   | 960  |
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| ccuaacgaga aggacugcga gacaagaggc gaccacgugu ucugugauac cgccgcugga   | 1020 |
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| ugcaaggugu ccaccggcag gcacccuauu ucuauggugg cucugucucc ucugggagcc   | 1140 |
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| gaccaggugu ucgagaacau cgagaauucc caggcucugg uggaccaguc caacagaauc   | 1440 |
| cugucuageg cegagaaggg aaacaeegge uucaucaueg ugaucauecu gauegeegug   | 1500 |
| cugggcagcu ccaugauccu gguguccauc uucaucauua ucaagaagac caagaagccc   | 1560 |
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| accggcuggu acaccaacgu guucacacug gaagugggcg acgucgagaa ucugacaugc   | 180  |
| ucugauggcc cuagccugau caagaccgag cuggaucugc ucaagagcgc ccugagagaa   | 240  |
| cucaagaccg ugucugccga ucagcuggcc agagaggaac agaucgagaa uccuggcagc   | 300  |
| ggcagcuuug ugcugggagc cauugcucuu ggaguggcug cugcugcagc uguuacagca   | 360  |
| ggcguggcca ucgcuaagac caucagacug gaaagcgaag ugaccgccau caacaacgcc   | 420  |
| cugaagaaga caaacgaggc cgucagcaca cucggcaaug gcguuagagu gcuggccaca   | 480  |
| gccgugcgcg agcugaagga cuucgugcuu aagaaccuga cacgggccau uaacaagaac   | 540  |
| aagugcgaca ucccugaccu gaagauggcc guguccuuua gccaguucaa ccggcgguuu   | 600  |
| cugaacgucg ugcggcaguu uagcgacaac gccggaauca caccagccau cagccuggac   | 660  |
| cugaugacag augcugagcu ggcuagagcc gugccuaaca ugccuacauc ugccggccag   | 720  |
| aucaagcuga ugcucgagaa uagagccaug guccgacgga aaggcuucgg cauucugauu   | 780  |
| ggcguguacg gcagcagcgu gaucuauaug gugcagcugc cuaucuucgg cgugaucgac   | 840  |
| acacccugcu ggauugugaa ggccgcuccu agcuguagcg agaagaaggg caauuacgcc   | 900  |
| ugccugcuga gagaggacca aggcugguau ugucagaacg ccggcagcac cguguacuac   | 960  |
| ccuaacgaga aggacugcga gacaagaggc gaccacgugu ucugugauac cgccgcugga   | 1020 |
| aucaaugugg ccgagcagag caaagagugc aacaucaaca ucagcaccac caacuauccc   | 1080 |
| ugcaaggugu ccaccggcag gcacccuauu ucuauggugg cucugucucc ucugggagcc   | 1140 |
| cugguggcuu guuauaaggg cguguccugu agcaucggca gcaacagagu gggcaucauc   | 1200 |
| aagcagcuga acaagggcug cagcuacauc accaaccagg acgccgauac cgugaccauc   | 1260 |
| gacaacaccg uguaucagcu gagcaaggug gaaggcgaac agcacgugau caagggcaga   | 1320 |

| ccugugucca gcagcuucga cccuaucaag uucccugagg aucaguucca gguggcccug   | 1380 |
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| gaccaggugu ucgagaacau cgagaauucc caggcucugg uggaccaguc caacagaauc   | 1440 |
| cugucuageg cegagaaggg aaacacegge uucaucaueg ugaucauecu gaucgeegug   | 1500 |
| cugggcageu ccaugauceu gguguccauc uucaucauua ucaagaagac caagaagece   | 1560 |
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| accggcuggu acaccaacgu guucacacug gaagugggcg acgucgagaa ucugacaugc   | 180  |
| ucugauggcc cuagccugau caagaccgag cuggaucugc ucaagagcgc ccugagagaa   | 240  |
| cucaagaccg ugucugccga ucagcuggcc agagaggaac agaucgagaa uccuggcagc   | 300  |
| ggcagcuuug ugcugggagc cauugcucuu ggaguggcug cugcugcagc uguuacagca   | 360  |
| ggcguggcca ucgcuaagac caucagacug gaaagcgaag ugaccgccau caacaacgcc   | 420  |
| cugaagaaga caaacgaggc cgucagcaca cucggcaaug gcguuagagu gcuggccaca   | 480  |
| gccgugcgcg agcugaagga cuucgugcuu aagaaccuga cacgggccau uaacaagaac   | 540  |
| aagugegaca ueeeugaeeu gaagauggee gugueeuuua geeaguucaa eeggegguuu   | 600  |
| cugaacgucg ugcggcaguu uagcgacaac gccggaauca caccagccau cagccuggac   | 660  |
| cugaugacag augcugagcu ggcuagagcc gugccuaaca ugccuacauc ugccggccag   | 720  |
| aucaagcuga ugcucgagaa uagagccaug guccgacgga aaggcuucgg cauucugauu   | 780  |
| ggcguguacg gcagcagcgu gaucuauaug gugcagcugc cuaucuucgg cgugaucgac   | 840  |
| acacccugcu ggauugugaa ggccgcuccu agcuguagcg agaagaaggg caauuacgcc   | 900  |
| ugccugcuga gagaggacca aggcugguau ugucagaacg ccggcagcac cguguacuac   | 960  |
| ccuaacgaga aggacugcga gacaagaggc gaccacgugu ucugugauac cgccgcugga   | 1020 |
| aucaaugugg ccgagcagag caaagagugc aacaucaaca ucagcaccac caacuauccc   | 1080 |
| ugcaaggugu ccaccggcag gcacccuauu ucuauggugg cucugucucc ucugggagcc   | 1140 |
| cugguggcuu guuauaaggg cguguccugu agcaucggca gcaacagagu gggcaucauc   | 1200 |
| aagcagcuga acaagggcug cagcuacauc accaaccagg acgccgauac cgugaccauc   | 1260 |
| gacaacaccg uguaucagcu gagcaaggug gaaggcgaac agcacgugau caagggcaga   | 1320 |
| ccugugucca gcagcuucga cccuaucaag uucccugaga accaguucca gguggcccug   | 1380 |
| gaccaggugu ucgagaacau cgagaauucc caggcucugg uggaccaguc caacagaauc   | 1440 |
| cugucuageg cegagaaggg aaacacegge uucaucaueg ugaucauecu gaucgeegug   | 1500 |
| cugggcagcu ccaugauccu gguguccauc uucaucauua ucaagaagac caagaagcc  | 1560 |
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300

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| cugaagaaga                 | caaacgaggc  | cgucagcaca | cucggcaaug | gcguuagagu | gcuggccaca | 480  |  |
| gccgugcgcg                 | agcugaagga  | cuucgugucc | aagaaccuga | cacgggccau | uaacaagaac | 540  |  |
| aagugcgaca                 | ucgacgaccu  | gaagauggcc | guguccuuua | gccaguucaa | ccggcgguuu | 600  |  |
| cugaacgucg                 | ugeggeaguu  | uagcgacaac | gccggaauca | caccagccau | cagccuggac | 660  |  |
| cugaugacag                 | augcugagcu  | ggcuagagcc | gugccuaaca | ugccuacauc | ugccggccag | 720  |  |
| aucaagcuga                 | ugcucgagaa  | uagagccaug | guccgacgga | aaggcuucgg | cauucugauu | 780  |  |
| ggcguguacg                 | gcagcagcgu  | gaucuauaug | gugcagcugc | cuaucuucgg | cgugaucgac | 840  |  |
| acacccugcu                 | ggauugugaa  | ggeegeueeu | agcuguagcg | agaagaaggg | caauuacgcc | 900  |  |
| ugccugcuga                 | gagaggacca  | aggcugguau | ugucagaacg | ccggcagcac | cguguacuac | 960  |  |
| ccuaacgaga                 | aggacugcga  | gacaagaggc | gaccacgugu | ucugugauac | cgccgcugga | 1020 |  |
| aucaaugugg                 | ccgagcagag  | caaagagugc | aacaucaaca | ucagcaccac | caacuauccc | 1080 |  |
| ugcaaggugu                 | ccaccggcag  | gcacccuauu | ucuauggugg | cucugucucc | ucugggagcc | 1140 |  |
| cugguggcuu                 | guuauaaggg  | cguguccugu | agcaucggca | gcaacagagu | gggcaucauc | 1200 |  |
| aagcagcuga                 | acaagggcug  | cagcuacauc | accaaccagg | acgccgauac | cgugaccauc | 1260 |  |
| gacaacaccg                 | uguaucagcu  | gagcaaggug | gaaggcgaac | agcacgugau | caagggcaga | 1320 |  |
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| gaccaggugu                 | ucgagaacau  | cgagaauucc | caggcucugg | uggaccaguc | caacagaauc | 1440 |  |
| cugucuagcg                 | ccgagaaggg  | aaacaccggc | uucaucaucg | ugaucauccu | gaucgccgug | 1500 |  |
| cugggcagcu                 | ccaugauccu  | gguguccauc | uucaucauua | ucaagaagac | caagaagccc | 1560 |  |
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|                            | ugucugccga  |            |            |            |            | 300  |  |
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| ggcguggcca                 | ucgcuaagac  | caucagacug | gaaagcgaag | ugaccgccau | caacaacgcc | 420  |  |
| cugaagaaga                 | caaacgaggc  | cgucagcaca | cucggcaaug | gcguuagagu | gcuggccaca | 480  |  |
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| aagugcgaca                 | ucgacgaccu  | gaagauggcc | guguccuuua | gccaguucaa | ccggcgguuu | 600  |  |
| cugaacgucg                 | ugcggcaguu  | uagcgacaac | gccggaauca | caccagccau | cagccuggac | 660  |  |
| cugaugacag                 | augcugagcu  | ggcuagagcc | gugccuaaca | ugccuacauc | ugccggccag | 720  |  |
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780

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| ugccugcuga                             | gagaggacca  | aggcugguau                             | ugucagaacg   | ccggcagcac   | cguguacuac   | 960                |
| ccuaacgaga                             | aggacugcga  | gacaagaggc                             | gaccacgugu   | ucugugauac   | cgccgcugga   | 1020               |
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| gaccaggugu                             | ucgagaacau  | cgagaauucc                             | caggcucugg   | uggaccaguc   | caacagaauc   | 1440               |
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| cugggcagcu                             | ccaugauccu  | gguguccauc                             | uucaucauua   | ucaagaagac   | caagaagccc   | 1560               |
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| gagagcuacc                             | uggaagaguc  | cugcagcacc                             | aucacagagg   | gcuaccuguc   | ugugcugaga   | 120                |
| accggcuggu                             | acaccaacgu  | guucacacug                             | gaagugggcg   | accucgagaa   | ucugacaugc   | 180                |
| ucugauggcc                             | cuagccugau  | caagaccgag                             | cuggaucuga   | ccaagagcgc   | ccugagagaa   | 240                |
| cucaagaccg                             | ugucugccga  | ucagcuggcc                             | agagaggaac   | agaucgagaa   | uccuggcagc   | 300                |
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| cugaagaaga                             | caaacgaggc  | cgucagcaca                             | cucggcaaug   | gcguuagagu   | gcuggccaca   | 480                |
| geegugegeg                             | agcugaagga  | cuucgugucc                             | aagaaccuga   | cacgggccau   | uaacaagaac   | 540                |
| aagugcgaca                             | ucgacgaccu  | gaagauggcc                             | guguccuuua   | gccaguucaa   | ccggcgguuu   | 600                |
| cugaacgucg                             | ugcggcaguu  | uagcgacaac                             | gccggaauca   | caccagccau   | cagccuggac   | 660                |
| cugaugacag                             | augcugagcu  | ggcuagagcc                             | gugccuaaca   | ugccuacauc   | ugccggccag   | 720                |
| aucaagcuga                             | ugcucgagaa  | uagagccaug                             | guccgacgga   | aaggcuucgg   | cauucugauu   | 780                |
| ggcguguacg                             |   |  |  |  |  | 840                |
|  | gcagcagcgu  | gaucuauaug                             | gugcagcugc   | cuaucuucgg   | cgugaucgac   | 010                |
| acacccugcu                             |   | gaucuauaug<br>ggccgcuccu               |  |  |  | 900                |
|  | ggauugugaa  |  | agcuguagcg   | agaagaaggg   | caauuacgcc   |                    |
| ugccugcuga                             | ggauugugaa<br>gagaggacca                                  | ggccgcuccu                             | agcuguagcg<br>ugucagaacg                             | agaagaaggg   | caauuacgcc   | 900                |
| ugccugcuga<br>ccuaacgaga               | ggauugugaa<br>gagaggacca<br>aggacugcga                    | ggccgcuccu                             | agcuguagcg<br>ugucagaacg<br>gaccacgugu               | agaagaaggg<br>ccggcagcac<br>ucugugauac               | caauuacgcc<br>cguguacuac<br>cgccgcugga               | 900<br>960         |
| ugccugcuga<br>ccuaacgaga<br>aucaaugugg | ggauugugaa<br>gagaggacca<br>aggacugcga<br>ccgagcagag      | ggccgcuccu<br>aggcugguau<br>gacaagaggc | agcuguagcg<br>ugucagaacg<br>gaccacgugu<br>aacaucaaca | agaagaaggg<br>ccggcagcac<br>ucugugauac<br>ucagcaccac | caauuacgcc<br>cguguacuac<br>cgccgcugga<br>caacuauccc | 900<br>960<br>1020 |

cugguggcuu guuauaaggg cguguccugu agcaucggca gcaacagagu gggcaucauc 1200

|  |                       |              | COILCIIIC    | 200        |      |  |
|--|-----------------------|--------------|--------------|------------|------|--|
| aagcagcuga acaagg  | gcug cagcuacaud       | accaaccagg   | acgccgauac c | gugaccauc  | 1260 |  |
| gacaacaccg uguauc  | agcu gagcaaggug       | gaaggcgaac   | agcacgugau c | aagggcaga  | 1320 |  |
| ccugugucca gcagcu  | ucga cccuaucaaç       | uucccugagg   | aucaguucca g | guggcccug  | 1380 |  |
| gaccaggugu ucgaga  | acau cgagaauuco       | caggcucugg   | uggaccaguc c | caacagaauc | 1440 |  |
| cugucuageg cegaga  | aggg aaacaccggo       | uucaucaucg   | ugaucauccu g | gaucgccgug | 1500 |  |
| cugggcagcu ccauga  | uccu gguguccaud       | uucaucauua   | ucaagaagac c | caagaagccc | 1560 |  |
| accggcgcuc cuccag  | aacu gagcggagug       | g accaacaaug | gcuucauccc u | ıcacaac    | 1617 |  |
| <210> SEQ ID NO 1 <211> LENGTH: 161 <212> TYPE: RNA <213> ORGANISM: A <220> FEATURE: <223> OTHER INFOR | 7<br>.rtificial Seque |              | eotide       |            |      |  |
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| augagcugga aggugg  | ucau caucuucago       | cugcugauca   | caccucagca c | ggccugaaa  | 60   |  |
| gagagcuacc uggaag  | aguc cugcagcaco       | aucacagagg   | gcuaccuguc u | ıgugcugaga | 120  |  |
| accggcuggu acacca  | acgu guucacacuo       | gaagugggcg   | acgucgagaa u | ıcugacaugc | 180  |  |
| ucugauggcc cuagco  | ugau caagaccgag       | g cuggaucuga | ccaagagcgc c | cugagagaa  | 240  |  |
| cucaagaccg ugucug  | ccga ucagcuggco       | agagaggaac   | agaucgagaa u | ıccuggcagc | 300  |  |
| ggcagcuuug ugcugg  | gage cauugeueui       | ı ggaguggcug | cugcugcage u | ıguuacagca | 360  |  |
| ggcguggcca ucgcua  | agac caucagacuo       | gaaagcgaag   | ugaccgccau c | caacaacgcc | 420  |  |
| cugaagaaga caaacg  | agge egueageaca       | cucggcaaug   | gcguuagagu g | gcuggccaca | 480  |  |
| gccgugcgcg agcuga  | agga cuucgugcui       | ı aagaaccuga | cacgggccau u | ıaacaagaac | 540  |  |
| aagugcgaca ucgacg  | accu gaagauggco       | guguccuuua   | gccaguucaa c | cggcgguuu  | 600  |  |
| cugaacgucg ugcggo  | aguu uagcgacaac       | geeggaauea   | caccagccau c | agccuggac  | 660  |  |
| cugaugacag augcug  | agcu ggcuagagco       | gugccuaaca   | ugccuacauc u | ıgccggccag | 720  |  |
| aucaagcuga ugcucg  | agaa uagagccaug       | guccgacgga   | aaggcuucgg c | auucugauu  | 780  |  |
| ggcguguacg gcagca  | gcgu gaucuauaug       | gugcagcugc   | cuaucuucgg c | gugaucgac  | 840  |  |
| acacccugcu ggauug  | ugaa ggccgcucci       | ı agcuguagcg | agaagaaggg c | caauuacgcc | 900  |  |
| ugccugcuga gagagg  | acca aggcugguau       | ı ugucagaacg | ccggcagcac c | guguacuac  | 960  |  |
| ccuaacgaga aggacu  | .gcga gacaagaggo      | gaccacgugu   | ucugugauac c | gccgcugga  | 1020 |  |
| aucaaugugg ccgago  | agag caaagagugo       | aacaucaaca   | ucagcaccac c | caacuauccc | 1080 |  |
| ugcaaggugu ccaccg  | gcag gcacccuaui       | ı ucuauggugg | cucugucucc u | ıcugggagcc | 1140 |  |
| cugguggcuu guuaua  | aggg cguguccugu       | ı agcaucggca | gcaacagagu g | ggcaucauc  | 1200 |  |
| aagcagcuga acaagg  | gcug cagcuacaud       | accaaccagg   | acgccgauac c | gugaccauc  | 1260 |  |
| gacaacaccg uguauc  | agcu gagcaaggug       | gaaggegaae   | agcacgugau c | aagggcaga  | 1320 |  |
| ccugugucca gcagcu  |                       |              |              |            | 1380 |  |
| gaccaggugu ucgaga  |                       |              |              |            | 1440 |  |
|  |                       |              |              |            | 1500 |  |
| cugucuageg cegaga  |                       |              |              |            |      |  |
| cugggcagcu ccauga  |                       |              |              |            | 1560 |  |
| accggcgcuc cuccag  | aacu gagcggagug       | g accaacaaug | gcuucauccc u | ıcacaac    | 1617 |  |

725 726

<210> SEQ ID NO 138

<211> LENGTH: 1617

<212> TYPE: RNA

<213 > ORGANISM: Artificial Sequence

<220> FEATURE:

<223> OTHER INFORMATION: Synthetic Polynucleotide

<400> SEQUENCE: 138

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<210> SEQ ID NO 139

<400> SEQUENCE: 139

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accggcuggu acaccaacgu guucacacug gaagugggg accucgagaa ucugacaugc 180

<sup>&</sup>lt;211> LENGTH: 1617

<sup>&</sup>lt;212> TYPE: RNA

<sup>&</sup>lt;213> ORGANISM: Artificial Sequence

<sup>&</sup>lt;220> FEATURE:

<sup>&</sup>lt;223> OTHER INFORMATION: Synthetic Polynucleotide

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| ucugauggcc cuagccugau caagaccgag cuggaucugc ucaagagcgc ccugagagaa   | 240  |  |  |  |  |
| cucaagaceg ugucugeega ucageuggee agagaggaac agauegagaa uccuggeage   | 300  |  |  |  |  |
| ggcagcuuug ugcugggagc cauugcucuu ggaguggcug cugcugcagc uguuacagca   | 360  |  |  |  |  |
| ggcguggcca ucgcuaagac caucagacug gaaagcgaag ugaccgccau caacaacgcc   | 420  |  |  |  |  |
| cugaagaaga caaacgaggc cgucagcaca cucggcaaug gcguuagagu gcuggccaca   | 480  |  |  |  |  |
| gccgugcgcg agcugaagga cuucgugcuu aagaaccugu ggcgggccau uaacaagaac   | 540  |  |  |  |  |
| aagugcgaca ucgacgaccu gaagauggcc guguccuuua gccaguucaa ccggcgguuu   | 600  |  |  |  |  |
| cugaacgucg ugcggcaguu uagcgacaac gccggaauca caccagccau cagccuggac   | 660  |  |  |  |  |
| cugaugacag augcugagcu ggcuagagcc gugccuaaca ugccuacauc ugccggccag   | 720  |  |  |  |  |
| aucaagcuga ugcucgagaa uagagccaug guccgacgga aaggcuucgg cauucugauu   | 780  |  |  |  |  |
| ggcguguacg gcagcagcgu gaucuauaug gugcagcugc cuaucuucgg cgugaucgac   | 840  |  |  |  |  |
| acacccugcu ggauugugaa ggccgcuccu agcuguagcg agaagaaggg caauuacgcc   | 900  |  |  |  |  |
| ugccugcuga gagaggacca aggcugguau ugucagaacg ccggcagcac cguguacuac   | 960  |  |  |  |  |
| ccuaacgaga aggacugcga gacaagaggc gaccacgugu ucugugauac cgccgcugga   | 1020 |  |  |  |  |
| aucaaugugg ccgagcagag caaagagugc aacaucaaca ucagcaccac caacuauccc   | 1080 |  |  |  |  |
| ugcaaggugu ccaccggcag gcacccuauu ucuauggugg cucugucucc ucugggagcc   | 1140 |  |  |  |  |
| cugguggcuu guuauaaggg cguguccugu agcaucggca gcaacagagu gggcaucauc   | 1200 |  |  |  |  |
| aagcagcuga acaagggcug cagcuacauc accaaccagg acgccgauac cgugaccauc   | 1260 |  |  |  |  |
| gacaacaccg uguaucagcu gagcaaggug gaaggcgaac agcacgugau caagggcaga   | 1320 |  |  |  |  |
| ccugugucca gcagcuucga cccuaucaag uucccugagg aucaguucca gguggcccug   | 1380 |  |  |  |  |
| gaccaggugu ucgagaacau cgagaauucc caggcucugg uggaccaguc caacagaauc   | 1440 |  |  |  |  |
| cugucuageg eegagaaggg aaacaeegge uucaucaueg ugaucaueeu gauegeegug   | 1500 |  |  |  |  |
| cugggcagcu ccaugauccu gguguccauc uucaucauua ucaagaagac caagaagccc   | 1560 |  |  |  |  |
| accggcgcuc cuccagaacu gagcggagug accaacaaug gcuucauccc ucacaac  | 1617 |  |  |  |  |
| <210> SEQ ID NO 140 <211> LENGTH: 1617 <212> TYPE: RNA <213> ORGANISM: Artificial Sequence <220> FEATURE: <223> OTHER INFORMATION: Synthetic Polynucleotide |      |  |  |  |  |
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| augageugga agguggueau caucuucage cugeugauca caccucagea eggeeugaaa   | 60   |  |  |  |  |
| gagagcuacc uggaagaguc cugcagcacc aucacagagg gcuaccuguc ugugcugaga   | 120  |  |  |  |  |
| accggcuggu acaccaacgu guucacacug ccugugggcg acgucgagaa ucugacaugc   | 180  |  |  |  |  |
| ucugauggcc cuagccugau caagaccgag cuggaucuga ccaagagcgc ccugagagaa   | 240  |  |  |  |  |
| cucaagaceg ugucugeega ucageuggee agagaggaac agauegagaa uccuggeage   | 300  |  |  |  |  |
| ggcagcuuug ugcugggagc cauugcucuu ggaguggcug cugcugcagc uguuacagca   | 360  |  |  |  |  |
| ggcguggcca ucgcuaagac caucagacug gaaagcgaag ugaccgccau caacaacgcc   | 420  |  |  |  |  |
| cugaagaaga caaacgagge egucagcaca cueggcaaug geguuagagu geuggceaca   | 480  |  |  |  |  |
| gccgugcgcg agcugaagga cuucgugucc aagaaccuga cacgggccau uaacaagaac   | 540  |  |  |  |  |
|   | 500  |  |  |  |  |

600

660

aagugcgaca ucgacgaccu gaagauggcc guguccuuua gccaguucaa ccggcgguuu

cugaacgucg ugcggcaguu uagcgacaac gccggaauca caccagccau cagccuggac

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| cugaugacag  | augcugagcu   | ggcuagagcc  | gugccuaaca  | ugccuacauc  | ugccggccag   | 720   |
|---|--|---|---|---|--|---|
| aucaagcuga  | ugcucgagaa   | uagagccaug  | guccgacgga  | aaggcuucgg  | cauucugauu   | 780   |
| ggcguguacg  | gcagcagcgu   | gaucuauaug  | gugcagcugc  | cuaucuucgg  | cgugaucgac   | 840   |
| acacccugcu  | ggauugugaa   | ggccgcuccu  | agcuguagcg  | agaagaaggg  | caauuacgcc   | 900   |
| ugccugcuga  | gagaggacca   | aggcugguau  | ugucagaacg  | ccggcagcac  | cguguacuac   | 960   |
| ccuaacgaga  | aggacugcga   | gacaagaggc  | gaccacgugu  | ucugugauac  | cgccgcugga   | 1020  |
| aucaaugugg  | ccgagcagag   | caaagagugc  | aacaucaaca  | ucagcaccac  | caacuauccc   | 1080  |
| ugcaaggugu  | ccaccggcag   | gcacccuauu  | ucuauggugg  | cucugucucc  | ucugggagcc   | 1140  |
| cugguggcuu  | guuauaaggg   | cguguccugu  | agcaucggca  | gcaacagagu  | gggcaucauc   | 1200  |
| aagcagcuga  | acaagggcug   | cagcuacauc  | accaaccagg  | acgccgauac  | cgugaccauc   | 1260  |
| gacaacaccg  | uguaucagcu   | gagcaaggug  | gaaggcgaac  | agcacgugau  | caagggcaga   | 1320  |
| ccugugucca  | gcagcuucga   | cccuaucaag  | uucccugagg  | aucaguucca  | gguggcccug   | 1380  |
| gaccaggugu  | ucgagaacau   | cgagaauucc  | caggcucugg  | uggaccaguc  | caacagaauc   | 1440  |
| cugucuageg  | ccgagaaggg   | aaacaccggc  | uucaucaucg  | ugaucauccu  | gaucgccgug   | 1500  |
| cugggcagcu  | ccaugauccu   | gguguccauc  | uucaucauua  | ucaagaagac  | caagaagccc   | 1560  |
| accggcgcuc  | cuccagaacu   | gagcggagug  | accaacaaug  | gcuucauccc  | ucacaac  | 1617  |
|   | NISM: Artifi   | icial Sequer  | nce   |   |  |   |
| <220> FEATU<br><223> OTHER<br><400> SEQUI   | R INFORMATIO   | ON: Syntheti  | ic Polynucle  | eotide  |  |   |
| <223> OTHER   | R INFORMATIO   |   |   |   | cggccugaaa   | 60  |
| <223> OTHER<br><400> SEQUI<br>augagcugga  | R INFORMATIO   | caucuucagc  | cugcugauca  | caccucagca  |  | 60<br>120   |
| <223> OTHEI<br><400> SEQUI<br>augagcugga<br>gagagcuacc  | R INFORMATIO   | caucuucagc<br>cugcagcacc  | cugcugauca<br>aucacagagg  | caccucagca<br>gcuaccuguc  | ugugcugaga   |   |
| <223> OTHER <400> SEQUI augagcugga gagagcuacc accggcuggu  | R INFORMATIO<br>ENCE: 141<br>agguggucau<br>uggaagaguc  | caucuucagc<br>cugcagcacc<br>guucacacug  | cugcugauca<br>aucacagagg<br>gaagugggcg  | caccucagca<br>gcuaccuguc<br>acgucgagaa  | ugugcugaga<br>ucugacaugc   | 120   |
| <223> OTHEI <400> SEQUI augagcugga gagagcuacc accggcuggu ucugauggcc   | R INFORMATIO<br>ENCE: 141<br>agguggucau<br>uggaagaguc<br>acaccaacgu  | caucuucagc<br>cugcagcacc<br>guucacacug<br>caagaccgag  | cugcugauca<br>aucacagagg<br>gaagugggcg<br>cuggaucuga  | caccucagca<br>gcuaccuguc<br>acgucgagaa<br>ccaagagcgc  | ugugcugaga<br>ucugacaugc<br>ccugagagaa   | 120<br>180  |
| <223> OTHER <400> SEQUI augagcugga gagagcuacc accggcuggu ucugauggcc cucaagaccg  | R INFORMATION SINCE: 141  agguggucau uggaagaguc acaccaacgu cuagccugau  | caucuucage<br>cugcageace<br>guucacacug<br>caagacegag<br>ucageuggee  | cugcugauca<br>aucacagagg<br>gaagugggcg<br>cuggaucuga<br>agagaggaac  | caccucagca<br>gcuaccuguc<br>acgucgagaa<br>ccaagagcgc<br>agaucgagaa  | ugugcugaga<br>ucugacaugc<br>ccugagagaa<br>uccuggcagc   | 120<br>180<br>240   |
| <223> OTHEI <400> SEQUI augagcugga gagagcuacc accggcuggu ucugauggcc cucaagaccg ggcagcuuug   | R INFORMATION SINCE: 141  agguggucau  uggaagaguc  acaccaacgu  cuagccugau  ugucugccga   | caucuucagc<br>cugcagcacc<br>guucacacug<br>caagaccgag<br>ucagcuggcc<br>cauugcucuu  | cugcugauca<br>aucacagagg<br>gaagugggcg<br>cuggaucuga<br>agagaggaac<br>ggaguggcug  | caccucagca<br>gcuaccuguc<br>acgucgagaa<br>ccaagagcgc<br>agaucgagaa<br>cugcugcagc  | ugugcugaga ucugacaugc ccugagagaaa uccuggcagc uguuacagca  | 120<br>180<br>240<br>300  |
| <223> OTHEI <400> SEQUI augagcugga gagagcuacc accggcuggu ucugauggcc cucaagaccg ggcagcuuug   | R INFORMATION Agguggucau uggaagaguc acaccaacgu cuagccugau ugucugccga ugcugggagc  | caucuucage cugcagcacc guucacacug caagaccgag ucagcuggcc cauugcucuu caucagacug  | cugcugauca aucacagagg gaagugggcg cuggaucuga agagaggaac ggaguggcug   | caccucagca<br>gcuaccuguc<br>acgucgagaa<br>ccaagagcgc<br>agaucgagaa<br>cugcugcagc<br>ugaccgccau  | ugugcugaga ucugacaugc ccugagagaa uccuggcagc uguuacagca caacaacgcc  | 120<br>180<br>240<br>300<br>360   |
| <223> OTHEI <400> SEQUI augagcugga gagagcuacc accggcuggu ucugauggcc cucaagaccg ggcagcuuug ggcguggcca cugaagaaga   | R INFORMATION AND STATE OF THE  | caucuucagc cugcagcacc guucacacug caagaccgag ucagcuggcc cauugcucuu caucagacug cgucagcaca   | cugcugauca aucacagagg gaaguggcg cuggaucuga agagaggaac ggaguggcug gaaagcgaag cucggcaaug  | caccucagca gcuaccuguc acgucgagaa ccaagagcgc agaucgagaa cugcugcagc ugaccgccau gcguuagagu   | ugugcugaga ucugacaugc ccugagagaaa uccuggcagc uguuacagca caacaacgcc gcuggccaca  | 120<br>180<br>240<br>300<br>360<br>420                                    |
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| gacaacaccg uguaucagcu gagcaaggug gaaggcgaac agcacgugau caagggcaga   | 1320 |
| ccugugucca gcagcuucga cccuaucaag uucccugagg aucaguucca gguggcccug   | 1380 |
| gaccaggugu ucgagaacau cgagaauucc caggcucugg uggaccaguc caacagaauc   | 1440 |
| cugucuageg cegagaaggg aaacaeegge uucaucaueg ugaucauecu gauegeegug   | 1500 |
| cugggcagcu ccaugauccu gguguccauc uucaucauua ucaagaagac caagaagcc  | 1560 |
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| accggcuggu acaccaacgu guucacacug gaaguggggg acgucgagaa ucugacaugc   | 240  |
| ucugauggcc cuagccugau caagaccgag cuggaucuga ccaagagcgc ccugagagaa cucaagaccg ugucugccga ucagcuggcc agagaggaac agaucgagaa uccuggcagc                         | 300  |
| ggcagcuuug ugcugggagc cauugcucuu ggaguggcug cugcugcagc uguuacagca   | 360  |
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| cugaacgucg ugcggcaguu uagcgacaac gccggaauca caccagccau cagccuggac   | 660  |
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| aucaagcuga ugcucgagaa uagagccaug guccgacgga aaggcuucgg cauucugauu   | 780  |
| ggcguguacg gcagcagcgu gaucuauaug gugcagcugc cuaucuucgg cgugaucgac   | 840  |
| acacccugcu ggauugugaa ggccgcuccu agcuguagcg agaagaaggg caauuacgcc   | 900  |
| ugccugcuga gagaggacca aggcugguau ugucagaacg ccggcagcac cguguacuac   | 960  |
| ccuaacgaga aggacugcga gacaagaggc gaccacgugu ucugugauac cgccgcugga   | 1020 |
| aucaaugugg ccgagcagag caaagagugc aacaucaaca ucagcaccac caacuauccc   | 1080 |
| ugcaaggugu ccaccggcag gcacccuauu ucuauggugg cucugucucc ucugggagcc   | 1140 |
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                                                                     780
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| gaccaggugu  | ucgagaacau                        | cgagaauucc | caggcucugg | uggaccaguc | caacagaauc | 1440 |
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| ucugauggcc  | cuagecugau                        | caagaccgag | cuggaucuga | ccaagagcgc | ccugagagaa | 240  |
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480

540

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## What is claimed is:

- 1. A method comprising administering to a subject a messenger ribonucleic acid (mRNA) comprising an open reading frame encoding a betacoronavirus (BetaCoV) S protein or S protein subunit formulated in a lipid nanoparticle in an effective amount to induce in the subject an 15 immune response to the BetaCoV S protein or S protein subunit, wherein the lipid nanoparticle comprises 20-60 mol % ionizable cationic lipid, 5-25 mol % neutral lipid, 25-55 mol % cholesterol, and 0.5-15 mol % PEG-modified lipid.
- 2. The method of claim 1, wherein the open reading frame  $_{20}$  encodes a BetaCoV S protein.
- 3. The method of claim 2, wherein the immune response is a neutralizing antibody response specific to the BetaCoV S protein.
- **4.** The method of claim **1**, wherein the open reading frame encodes a BetaCoV S protein subunit selected from an S1 subunit and an S2 subunit.
- **5**. The method of claim **4**, wherein the immune response is a neutralizing antibody response specific to the BetaCoV S protein subunit.
- **6**. The method of claim **1**, wherein the mRNA formulated in a lipid nanoparticle is administered intramuscularly.
- 7. The method of claim 1, wherein the mRNA further comprises a 5' untranslated region and a 3' untranslated region.
- 8. The method of claim 1, wherein the mRNA further comprises a poly(A) tail.

  15 frame encodes a BetaCoV S protein.

  19. The method of claim 18, wherein the mRNA further comprises a poly(A) tail.
- 9. The method of claim 1, wherein the mRNA further comprises a 5' cap analog.
- 10. The method of claim 9, wherein the 5' cap analog is 7mG(5')ppp(5')NlmpNp.
- 11. The method of claim 1, wherein the mRNA comprises a chemical modification.
- 12. The method of claim 11, wherein the chemical modification is a 1-methylpseudouridine modification or a 1-ethylpseudouridine modification.

- 13. The method of claim 11, wherein at least 80% of the uracil in the open reading frame of the mRNA has a chemical modification.
- 14. The method of claim 1, wherein the lipid nanoparticle comprises 50 mol % ionizable cationic lipid, 10 mol % neutral lipid, 38.5 mol % cholesterol, and 1.5 mol % PEG-modified lipid.
- **15**. The method of claim 1, wherein the ionizable cationic lipid is Compound 25.
- **16**. The method of claim **1**, wherein the neutral lipid is 1,2-distearoyl-sn-glycero-3-phosphocholine (DSPC), and the PEG-modified lipid is 1,2-dimyristoyl-rac-glycero-3-methoxypolyethylene glycol-2000 (PEG-DMG).
- 17. A method comprising administering to a subject an mRNA comprising a 5' cap analog, a 5' untranslated region, an open reading frame encoding a BetaCoV S protein or S protein subunit, a 3' untranslated region, and a poly(A) tail formulated in a lipid nanoparticle in an effective amount to induce in the subject an immune response to the BetaCoV S protein or S protein subunit, wherein the lipid nanoparticle comprises 20-60 mol % ionizable cationic lipid, 5-25 mol % neutral lipid, 25-55 mol % cholesterol, and 0.5-15 mol % PEG-modified lipid.
- **18**. The method of claim **17**, wherein the open reading frame encodes a BetaCoV S protein.
- **19**. The method of claim **18**, wherein the ionizable cationic lipid is Compound 25, the neutral lipid is DSPC, and the PEG-modified lipid is PEG-DMG.
- **20**. The method of claim **18**, wherein at least 80% of the uracil in the open reading frame of the mRNA has a 1-methylpseudouridine modification.
  - **21**. The method of claim **20**, wherein the ionizable cationic lipid is Compound **25**, the neutral lipid is DSPC, and the PEG-modified lipid is PEG-DMG.

\* \* \* \* \*