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(54) Title: SUBUNIT VACCINE AGAINST FLAVIVIRUS INFECTION

(57) Abstract

The invention provides vaccines capable of eliciting the production of neutralizing antibodies against flavivirus containing, as an active ingredient, a secreted recombinantly produced envelope protein or subunit thereof. In illustrative embodiments, the B domain or the 60 % or 80 % N-terminal portions of the envelope protein (E) of dengue virus are secreted. The secreted product can easily be purified and prepared as a vaccine.

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SUBUNIT VACCINE AGAINST FLAVIVIRUS INFECTION

- 1 -

Technical Field

The invention relates to protection against and diagnosis of the conditions caused by flavivirus. More specifically, the invention concerns subunits of the flavivirus envelope protein secreted as mature recombinantly produced proteins from eucaryotic cells. Additional viral

- 10 proteins or subunits, also produced in this way, provide additional active ingredients. These E-proteins or subunits, alone or in combination including combination with additional viral-derived peptides are protective against infection by flavivirus, raise antibodies useful in passive
- 15 immunization, and are useful in diagnosis of infection by the virus.

Background Art

The flavivirus family, Flaviviridae, includes the 20 dengue viruses Japanese encephalitis (JEV) virus, Tick-borne encephalitis (TBE) virus, and the initially discovered prototype of this class, the yellow fever (YF) virus. The flaviviruses contain a single positive strand genomic RNA and are small enveloped viruses affecting animals, but

25 generally transmitted to vertebrates by chronically infected mosquito or tick vectors. Flaviviruses are enveloped by host cell membrane and contain the three structural proteins capsid (C), membrane (M), and envelope (E). The E and M proteins are found on the surface of the virion where they 30 are anchored in the membrane. Mature E is glycosylated, whereas M is not, although its precursor, prM, is a glycoprotein. Glycoprotein E, the largest structural protein, contains functional domains responsible for cell surface attachment and intraendosomal fusion activities. It 35 is also a major target of the host immune system, inducing neutralizing antibodies, protective immunity, as well as

antibodies which inhibit hemagglutination.

Dengue virus is the causative agent of dengue fever and is transmitted to man by Aedes mosquitoes, principally Aedes aegypti and Aedes albopictus. Classic dengue fever is an acute illness marked by fever, headache, aching muscles and joints, and rash. A fraction of cases, typically in children, results in more extreme forms of infection, i.e., dengue hemorrhagic fever (DHF) or dengue shock syndrome (DSS). Without diagnosis and prompt medical intervention, the sudden onset and rapid progress of DHF/DSS can be fatal.

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Dengue is one of the most important virus groups transmitted to man by arthropods in terms of global morbidity; it has been estimated that dengue is responsible for up to 100 million illnesses annually. With the advent of modern jet travel, dengue has spread globally in the tropics and subtropics, and multiple dengue serotypes in a region are common.

Every flavivirus genome is a single positive-stranded RNA of approximately 10,500 nucleotides 20 containing short 5' and 3' untranslated regions, a single long open reading frame (ORF), a 5' cap, and a nonpolyadenylated 3' terminus. The ten gene products encoded by the single, long open reading frame are contained in a polyprotein organized in the order, C (capsid), prM/M

- 25 (membrane), E (envelope), NS1 (nonstructural), NS2a, NS2b, NS3, NS4a, NS4b, and NS5 (Chambers, T.J. et al. <u>Ann Rev</u> <u>Microbiol</u> (1990) <u>44</u>:649-688). Processing of the encoded polyprotein is initiated cotranslationally, and full maturation requires both host and viral-specific proteases.
- 30 The sites of proteolytic cleavage in the YF virus have been determined by comparing the nucleotide sequence and the amino terminal sequences of the viral proteins. Subsequent to initial processing of the polyprotein, prM is converted to M during virus release (Wengler, G. et al. <u>J Virol</u> (1989)
 35 <u>63</u>:2521-2526), and anchored C is processed during virus maturation (Nowak et al. <u>Virology</u> (1987) <u>156</u>:127-137).

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There are four antigenically related dengue viruses which, however, can be recognized as distinct serotypes. The complete genomic sequence for at least one strain of each of the four dengue serotypes has been reported (DEN-1, Fu, J. et al. <u>Virology</u> (1992) <u>188</u>:953-958; DEN-2, Deubel, V. et al. <u>Virology</u> (1986) <u>155</u>:365-377; Hahn, Y.S. et al. <u>Virology</u> (1988) <u>162</u>:167-180; DEN-3, Osatomi, K. et al. <u>Virus Genes</u> (1988) <u>2</u>:99-108; Osatomi, K. et al. <u>Virology</u> (1990) <u>176</u>:643-647; DEN-4, Zhao, B.E. et al.

10 <u>Virology</u> (1986) <u>155</u>:77-88; Mackow, E. et al. <u>Virology</u> (1987) <u>159</u>:217-228). In addition, the compete genomic sequences of other flaviviruses are known (e.g., YF virus: Rice et al., <u>Science</u> (1985) <u>229</u>:726-733).

It does not appear that infection by one dengue serotype can confer long-term immunity on the individual with respect to other serotypes. In fact, secondary infections with heterologous serotypes are quite common. In general, antibody responses in infected subjects to primary infection are mostly IgM antibodies and these antibodies are directed against type-specific determinants. On the other hand, secondary infections by heterologous serotypes generate IgG antibodies which are flavivirus crossreactive.

Helpful reviews of the nature of the dengue disease, the history of attempts to develop suitable vaccines, and structural features of flaviviruses in general as well as the molecular structural features of the envelope protein of flaviviruses are found in Halstead, S.B. <u>Science</u> (1988) <u>239</u>:476-481; Brandt, W.E. <u>J Infect Disease</u> (1990) <u>162</u>:577-583; Chambers, T.J. et al. <u>Annual Rev Microbiol</u> (1990) <u>44</u>:649-688; Mandl, C.W. et al. <u>Virology</u> (1989) <u>63</u>:564-571; and Henchal, E.A. and J.R. Putnak, <u>Clin</u>

<u>Microbiol Rev</u> (1990) <u>3</u>:376-396.

Other flavivirus family members, i.e., JEV, TBE and YF appear presently to represent single serotypes -i.e., antibodies elicited in response to one strain are cross reactive with other known strains. **WO 96/37221**

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A successful vaccine for protection against flavivirus infection has never been developed. However, there have been a number of preliminary efforts, many of which focus on the envelope protein, since this protein is exposed at the surface and is believed to be responsible for eliciting immunity.

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Monoclonal antibodies (Mabs) directed against purified E of several flaviviruses, DEN-2 (Henchal *et al*. <u>Am</u> <u>J Trop Med Hyg</u> (1985) <u>34</u>:162-169, TBE (Heinz, F.X. *et al*.

Virology (1983) <u>126</u>:525-537), St. Louis encephalitis (SLE, Mathews, J.H. et al. <u>J Immunol</u> (1984) <u>132</u>:1533-1537), Murray Valley encephalitis (MVE, Hawkes, R.A. et al. <u>J Gen Virol</u> (1988) <u>69</u>:1105-1109), and JEV (Takegami, T. et al. <u>Acta</u> <u>Virologica</u> (1982) <u>26</u>:312-320) are neutralizing in vitro.

15 Some of these Mabs can also passively confer protection in vivo (Heinz, F.X. et al. (1983, supra)); Mathews, J.H. et al. (1984, supra)); Kimuro-Kuroda and Yasui, <u>J Immunol</u> (1988) <u>141</u>:3603-3610).

Although the primary amino acid sequence of the 20 flavivirus E glycoprotein is variable (45-80% identity), all have twelve conserved cysteine residues, forming six disulfide bridges, and hydrophilicity profiles are nearly superimposable, suggesting that they may all have similar secondary and tertiary structures. Based on the position of

- 25 the 12 conserved cysteines (determined for West Nile virus, Nowak and Wengler, <u>Virology</u> (1987) <u>156</u>:127-137), monoclonal antibody competitive binding studies, monoclonal antibody binding to purified proteolytic fragments, and analysis of neutralizing antibody escape mutants of Tick-Borne
- 30 Encephalitis Virus, glycoprotein E was divided into three antigenic domains (A, B, and C) and two transmembrane segments at its carboxy-terminus. See, for example, Mandl, C.W. et al. <u>J Virol</u> (1989) <u>63</u>:564-571. Figure 1, reproduced from this article, shows the locations of these domains.

Domain A was defined as a denaturation sensitive, highly folded, and disulfide stabilized discontinuous domain composed of the amino acids from 50-125 and 200-250

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containing five of the six disulfide bridges. Neutralization and hemagglutination inhibition epitopes are found within domain A, and, for dengue viruses, one of the two N-linked glycosylation sites. A conserved hydrophobic amino acid sequence within domain A has been postulated to provide fusogenic properties after low pH treatment. Amino acid sequences conserved among the flavivirus family are located within this region; thus, broadly

flavivirus-cross-reactive epitopes lie within this domain.

Domain B was identified as a continuous domain composed of amino acids 301-395 (an approximate region between amino acids 300-400 for all flaviviruses). The domain can be isolated as a single immunoreactive proteolytic fragment. It has been postulated that this

- 15 domain forms part of the receptor binding site (Heinz, F.X. et al. <u>APMIS</u> (1993) <u>101</u>:735-745), and attenuating mutations have been mapped to sequences within domain B Heinz et al. (supra). A variety of neutralizing antibodies have been shown to specifically map to Domain B (Heinz et al. (1983, supra)); Trirawatanapong et al., 1992; Megret et al., 1992;
- Lin et al., 1994). The binding of these neutralizing
 monoclonal antibodies is dependent on formation of a
 disulfide bond, and in some cases also is sensitive to
 detergent denaturation. Species-specific monoclonal
 antibodies bind this domain.

Domain C represents a hypervariable loop between the two segments of Domain A. Its antigenicity is insensitive to denaturation or reducing agents, and contains one N-linked glycosylation site. Predominantly sub-type specific monoclonal antibodies react with this domain. No specific activity has been assigned to this domain.

Many strategies are currently under investigation to develop an effective and safe dengue vaccine; however, to date, no single strategy has proven completely satisfactory. Attempts to generate live attenuated dengue vaccine strains have not been entirely successful, although research into this area continues. In the absence of effective, live

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attenuated dengue vaccines, a significant effort has been invested in the development of recombinant, dengue subunit or viral-vectored vaccines.

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- Recombinant dengue proteins have been expressed in several systems to date (see Putnak, R.A. (1994) Modern 5 Vaccinology, E. Kurstak ed., Plenum Medical, New York, pp. 231-252, for review). Most efforts using Escherichia coli have yielded poor immunogen unable to elicit neutralizing This may reflect nonnative conformation of antibodies. 10 dengue proteins expressed in the bacteria and the necessity to process the viral proteins through the secretion pathway in order to form the proper disulfide bonds, glycosylate the proteins, or both.
- Several reports have described vaccinia-flavivirus 15 recombinants expressing envelope protein as part of a polyprotein (e.g. C-prM-E-NS1; [Dengue] Zhao, B.G. et al. J <u>Virol</u> (1987) <u>61</u>:4019-4022; Deubel, V. et al. <u>J Gen Virol</u> (1988) <u>69</u>:1921-1929; Bray, M. et al. <u>J Virol</u> (1991) 63:2853-2856; [YF] Hahn, Y.S. et al. Arch Virol (1990)
- 20 115:251-265), as a single protein (e.g. 100%E; [Denque] Bray, M. et al., <u>J Virol</u> (1989) <u>63</u>:2853-2856), or as polypeptides (e.g. 79%E-RKG; Men, R. et al. J Virol (1991) 65:1400-1407). The most successful recombinant vaccinia viruses, those capable of inducing neutralizing antibodies
- 25 and protecting mice from virus challenge, were those which were secreted E extracellularly or accumulated E on the cell surface.

Men, R. et al. (1991, supra) described the recombinant production of various C-terminal truncations of the DEN-4 envelope protein using a recombinant Vaccinia 30 virus vector and infecting mammalian CV1 cells. The results showed that the recombinants that contain greater than 79% of the coding sequence produced an intracellular protein that could be immunoprecipitated with anti-dengue virus 35 antibodies contained in hyperimmune mouse ascitic fluid (HMAF). Although there was a reduced level of detection for protein which contained 79% of envelope or less, this did

not appear to result from reduced production of the protein. It was also found that only truncations which contained 79% of E or less were secreted efficiently; E polypeptides equal to or larger than 81%E were not secreted efficiently.

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Men et al. (1991, supra) constructed additional Cterminal truncations between 79%E and 81%E to map the amino acids responsible for the difference in secretion and immunoreactivity with HMAF of these two truncated E polypeptides. The results demonstrated that 79% E containing the additional tripeptide sequence RKG was also 10 secreted. Although both 59% E and 79% E-RKG were secreted, only 79% E-RKG was detected at the cells' surface. The recombinant Vaccinia viruses containing various truncations were also used to immunize mice. Mice immunized with recombinants expressing 79%E-RKG or larger portions of the envelope protein were protected. However, except for 59% E, mice immunized with 79% E or a smaller product were only partially protected. The 59%E elicited high protection rates (>90%) comparable to 79%E-RKG and larger C-terminal 20 truncated E polypeptides. Protection correlated with

binding to HMAF.

Combinations of immunogenic structural and nonstructural JEV virus, DEN-1, DEN-2, and DEN-4 proteins have been expressed by baculovirus recombinants (Matsuura, Y. et al. <u>Virology</u> (1989) <u>173</u>:674-682; Putnak, R.A. et al. Am J

<u>Trop Med Hyg</u> (1991) <u>45</u>:159-167; Deubel, V. et al. <u>Virology</u> (1991) <u>180</u>:442-447). Baculovirus-expressed dengue and JE E glycoprotein elicited neutralizing antibodies, protected mice from a lethal dengue virus challenge, or both. In

30 spite of these successes, the expression levels reported in baculovirus are low and the recombinant protein is less immunogenic than the viral protein (Putnak, R.A. et al. Am J Trop Med Hyq (1991) supra).

Research with purified polypeptides released by 35 proteolysis of flavivirus envelope proteins, with recombinant polypeptides, and with synthetic peptides has indicated where protective epitopes may map. The isolated

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9000 dalton domain B trypsin fragment from TBE virus spontaneously refolds and is stabilized by disulfide bridges (Winkler, G. et al. J Gen Virol (1987) <u>68</u>:2239-2244). This disulfide stabilized fragment elicited neutralizing antibodies in mice (Heinz, F.X. et al. <u>Virology</u> (1984) <u>130</u>:485-501). In contrast, a 28,000 dalton trypsin fragment from WN virus containing domain B sequences was unable to spontaneously refold and did not elicit neutralizing antibodies (Wengler and Wengler, 1989).

10 A cyanogen bromide-cleaved 8 kD fragment (amino acids 375-456) overlapping domain B from JEV envelope protein was found to induce neutralizing antibodies in mice (Srivastava, A.K. et al. <u>Acta Virol</u> (1990) <u>34</u>:228-238). Immunization of mice with a larger polypeptide (JE E amino 15 acid 319 to NS1 amino acid 65) spanning the 8 kD peptide expressed in Escherichia coli as a fusion to protein A elicited neutralizing antibodies and protected mice from lethal virus challenge (Srivastava, A.K. et al. Microbiol Immunol (1991) 35:863-870). This polypeptide begins between 20 the two cysteines within domain B, and, therefore, cannot form the stabilizing disulfide bond that earlier reports suggest is necessary for presentation of protective epitopes.

Immunization of mice with synthetic peptides 25 corresponding to amino acids within domain B, aa 356-376 from MVE (Roehrig, J.T. et al. <u>Virology</u> (1989) <u>171</u>:49-60) or aa 352-368 from DEN-2 virus (Roehrig, J.T. et al. <u>Virology</u> (1990) <u>177</u>:668-675), elicited low levels of neutralizing antibodies in mice, suggesting the presence in domain B of a 30 weak linear neutralizing epitope (Roehrig, J.T. et al. 1989 and 1990, supra).

Mason, P.W. et al. <u>J Gen Virol</u> (1990) <u>71</u>:2107-2114 identified two domains of the DEN-1 envelope protein: domain I which includes amino acids 76-93 of the E protein and domain II (equivalent to domain B) which includes amino acids 293-402. These domains were identified from deletion analysis using recombinant fusion proteins expressed in E. coli and reacted with antiviral monoclonal antibodies. Recombinant fusion proteins containing E. coli trpE sequences fused to the envelope protein (amino acids 1 to 412) elicited antibodies in mice which reacted with a portion of the protein containing domain II.

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In addition, Mason, P.W. et al. (<u>J Gen Virol</u> (1989) <u>70</u>:2037-2049) expressed a collection of *E. coli trpE* fusion proteins to segments of JEV virus envelope protein spanning domain B. The *trpE* fusion proteins containing the smallest JEV E fragments that retained immunoreactivity with a panel of neutralizing monoclonal antibodies included amino acid residues from methionine 303 through tryptophan 396. However, animals immunized with immunoreactive *trpE* fusion polypeptides did not produce neutralizing antibodies nor were they protected from lethal challenge.

Trirawatanapong, T. et al. <u>Gene</u> (1992) <u>116</u>:139-150 prepared several truncated forms of dengue 2 envelope proteins in *E. coli* for epitope mapping, and used the neutralizing monoclonal antibody 3H5 to locate its 20 corresponding epitope. This was first localized between amino acids 255 and 422. Targeted gene deletions in the plasmid constructs encoding the truncated proteins permitted mapping of the binding site to the 12 amino acids between positions 386 and 397. The mapping was apparently confirmed

25 by the ability of a synthetic peptide containing E protein amino acids 386-397 to bind 3H5 specifically.

Megret, F. et al. <u>Virology</u> (1992) <u>187</u>:480-491 prepared 16 overlapping fragments of DEN-2 envelope protein as trpE fusion products in *E. coli* for epitope mapping. The fusion proteins are produced intracellularly and obtained from the lysates. These products were used to map epitopes defined by a panel of 20 monoclonal antibodies. Six antigenic domains were described: nonneutralizing

antibodies bound to peptides containing amino acids 22-58, 35 amino acids 304-332, amino acids 60-97, and amino acids 298-397. Neutralizing antibodies bound to peptides containing amino acids 60-135, 60-205, and 298-397.

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Significantly, Megret et al. (1992, supra) demonstrated that all MAbs (including 3H5), with two exceptions (below), that recognize "full-length" domain B (amino acids 298-397) are unable to recognize slightly shorter polypeptides. For example, in contrast to the findings of Trirawatanapong et al. Gene (1992, supra), MAb 3H5 was unable to bind to trpE fusion proteins containing DEN-2 E amino acids 304-397, 298-385, or 366-424. The two exceptional MAbs in the findings of Megret et al. are MAbs 5A2 and 9D12. The pattern of binding of MAb 5A2 suggests that it recognizes a linear epitope between amino acids 304 to 332, while MAb 9D12 binds to a polypeptide, amino acids 298-385, which is slightly shorter than the smallest polypeptide (amino acids 298-397) to which MAb 3H5 binds. These results indicate that both the disulfide bond in domain B and the domain B C-terminal amino acids are involved in forming the immunodominant domain B epitopes.

U.S. Patent No. 5,494,671, issued 27 February 1996 discloses C-terminally truncated dengue and JEV envelope 20 proteins for use in vaccines. This patent is the U.S. counterpart of PCT Application WO 92/03161 published 5 March The truncated E proteins were either generated in 1992. situ using viral vectors, or in insect cells, also using viral vectors.

25 Although it appears established from the art that the B domain and subunits representing at least about 80% E (truncated at the C-terminus), of the flavivirus envelope protein contain epitopes which bind neutralizing antibodies, problems have arisen with respect to producing recombinant 30 polypeptides, all of which contain the B domain, in a form which mimics the native protein and is thus capable of eliciting an immune response. The only recombinantly produced E polypeptides containing the B domain that elicited a protective immune response in mice were expressed 35 from Vaccinia and baculovirus vectors. Generally, recombinantly produced proteins lack the appropriate

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glycosylation, folding, and disulfide bond formation for producing a proper immune response.

It has now been found that the B domain and other shortened forms of the envelope protein can be successfully secreted from yeast and from *Drosophila* in a form which elicits the production of neutralizing antibodies. This permits, for the first time, the production of a successful recombinantly produced subunit flavivirus vaccine.

10 <u>Disclosure of the Invention</u>

The invention provides vaccines containing, as an active ingredient, a secreted recombinantly produced the flavivirus envelope protein or a subunit thereof. The vaccines are capable of eliciting the production of

- 15 neutralizing antibodies against the relevant flavivirus. In the illustrations below, the B domain of the envelope protein (E) is secreted from yeast by producing it in an expression vector containing the α-mating factor prepropeptide leader sequence (preproMFa_L). Peptide
- 20 subunits representing 60%E and 80%E are secreted from Drosophila cells using the human tissue plasminogen activator secretion signal sequence for the propeptide (tPAL) or from the homologous premembrane (prM) leader. 100%E can also be so produced. The secreted products can 25 easily be purified and prepared as a vaccine.

Thus, in one aspect, the invention is directed to a vaccine for protection of a subject against infection by flavivirus. The vaccine contains, as active ingredient, the envelope protein of a flavivirus serotype or a subunit

30 thereof. The E or subunit is secreted as a recombinantly produced protein from eucaryotic cells under the control of a nonviral promoter. The vaccine preferably contains portions of more than one flavivirus serotype E protein similarly produced, and may further include additional
35 flavivirus proteins, particularly NS-1, which can be similarly recombinantly produced.

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In other aspects, the invention is directed to methods to protect subjects against infection by administering the invention vaccines, to antibodies generated in a mammalian subject administered an immunogenic amount of the vaccine; immortalized B cell lines which generate monoclonal antibodies of this type; methods to effect passive immunization by administering the antibodies of the invention; methods to detect the presence or absence of antiflavivirus immunoglobulins utilizing the secreted recombinantly produced peptides of the invention and the recombinant materials important in, methods for, their production.

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Brief Description of the Drawings

Figure 1 is a drawing reproduced from Mandl, et al. (supra) showing a model of the envelope protein of flaviviruses.

Figure 2 (SEQ ID NO:1) shows the partial nucleotide sequence for DEN-2 PR159 S1 mutant strain and differences from the wild-type strain reported by Hahn (1988, *supra*).

Figure 3 (SEQ ID NO:2 and SEQ ID NO:3) shows the partial nucleotide sequence and deduced amino acid sequence of the genome of DEN-2 PR159/S1 strain in comparison with wild-type.

Figure 4 (Nucleotides 1693-1752 of SEQ ID NO:2 and SEQ ID NO:4 through SEQ ID NO:7) shows the oligonucleotide used to mutagenize an 80%E cDNA clone to obtain the domain B coding sequence.

Figure 5 shows the construction of a cloning vector containing the nucleotide sequence encoding domain B.

Figure 6 shows the insertion of the domain B coding sequence into the yeast expression vector pLS5.

Figure 7 (SEQ ID NO:8 and SEQ ID NO:9) shows the 35 $\,$ preproMFa_L/domain B fusion protein.

Figure 8 shows the construction of an expression vector for domain B in Pichia.

Figure 9 (SEQ ID NO:10 through SEQ ID NO:12) shows the nucleotide and deduced amino acid sequence for the tPA_L-40 DomB fusion protein.



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Figure 10 shows the survival times of mice immunized with recombinant domain B and challenged with Dengue-2.

Figure 11 shows the construction of an expression 5 vector for production of domain B in *Drosophila melanogaster* tissue cultured Schneider cells.

Figure 12 shows the standard curve for a DomB sandwich assay.

Figure 13 shows the survival times of mice 10 immunized with *D. melanogaster* Schneider cell-secreted 80%E.

Modes of Carrying Out the Invention

The invention provides, for the first time, a subunit vaccine that can be efficiently produced

- 15 recombinantly and secreted that is effective in protecting subjects against infection with flavivirus. Although many attempts have been made to obtain such a subunit vaccine, either the subunit itself is resistant to recombinant production techniques which permit it to be secreted in a
- 20 processed form so as to render it effective as an immunogen, or, if its recombinant production is facile, it fails to elicit neutralizing antibodies. The present applicants have found that certain portions of the envelope protein of dengue virus type 2, such as domain B representing
- 25 approximately 100 amino acids of the envelope protein extending approximately from the Gly at position 296 to the Gly at position 395, and optionally including additional E sequence through position 413 of the protein, and other portions of E, i.e., 60%E and 80%E as well as 100%E are
- 30 effectively secreted by certain convenient eucaryotic recombinant hosts, in a form that permits processing to mimic the native conformation of the protein. The hosts generally utilize nonviral control sequences. The secretion of the protein into the culture medium facilitates 35 purification. Furthermore, this form is able, when administered, especially in the presence of adjuvant, to raise neutralizing antibodies in animals. Thus, this

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subunit represents a useful component of a vaccine for protecting subjects against dengue infection.

As used herein, "B domain" refers to a peptide which spans from approximately Gly 296 to Gly 395 of the DEN-2 envelope protein, and optionally including additional 5 E sequence through position 413 of the envelope protein. The corresponding positions in other flaviviruses define the B domain generally. These positions are approximate; for example, Mandl (1989, supra) describes the generation of a tryptic fragment containing domain B which spans the amino acids of the TBE E protein from position 301 to 396. The sequences illustrated in the present application represent the envelope protein from dengue Type 2 virus (DEN-2); three additional distinct dengue serotypes have been recognized.

- Therefore, "Domain B" also refers to the corresponding peptide region of the envelope protein of these serotypes, of other flaviviruses and to any naturally occurring In addition, B domain includes extended forms of variants. the about 100-120 amino acid peptides, wherein the
- 20 extensions do not interfere with the immunogenic effectiveness or secretion of the B domain. In one embodiment, such extensions are minimal -- i.e., not more than six additional amino acids -- at either the N-terminus or the C-terminus, or distributed between these termini;
- 25 preferably no more than four total additional amino acids, and most preferably no more than two.

The form of domain B which spans positions of about 296-395 is referred to herein as "classical" domain B. When the B domain includes at least portions of the region extending to amino acid 413, the additional region may confer additional functions, e.g., enhancing immunogenicity by providing a helper T cell epitope. The form of domain B which includes positions about 296-413 (of DEN2) and corresponding positions in other flaviviruses, is referred The domain B of the invention includes to herein as DomB+T. these two specific embodiments, "classical" domain B and

DomB+T, as well as those forms which span positions

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approximately 296 to a position between position 395 and 413.

Other portions of the E protein illustrated below are self-explanatory. 80%E is the N-terminal 80% of the protein from residue 1 to approximately residue 395. 60%E represents the corresponding shorter sequence. These precentages are approximate and may include 1-4 amino acid residues more or less than that calculated precisely. These subunits are produced from vectors containing the DNA encoding the mature subunit or protein or may be included in a prM fusion which results in secretion of the 80% or 60%E per se.

For practical large-scale production of the subunits used as active ingredients in the vaccines of the 15 invention, recombinant techniques provide the most practical approach. However, to be useful as active ingredients, the subunits as produced must assume a conformation and undergo processing under conditions which render them similar to the native envelope portion as it exists in the envelope protein

20 of the virus. In order to achieve this, the recombinant production must be conducted in eucaryotic cells, preferably yeast or *Drosophila* cells. In addition to the *S. cerevisiae* and *P. pastoris* yeasts illustrated below, other yeasts, such as *Kluveromyces* sp Yarrowia lipolytica, and

25 Schizosaccharomyces pombe may be used. Other insect cells besides the Drosophila melanogaster Schneider cells illustrated below may also be employed. Additional appropriate eucaryotic cells include mammalian expression cells such as Chinese hamster ovary cells. The B domain or 30 60%E or 80%E must be produced as a correctly processed protein and secreted.

It has been found, as demonstrated hereinbelow, that particularly efficient secretion of biologically active mature protein can be achieved in several ways. First, this can be done by expressing the protein in yeast in operable linkage with the α -mating factor signal sequence. Constructs which place the nucleotide sequence encoding the

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E protein or subunit disposed so as to encode a fusion protein with an upstream α -mating factor signal sequence are therefore included within the scope of the invention. An additional preferred embodiment employs Drosophila cells and

- 5 the human tissue plasminogen activator leader sequence for secretion of 60%E or 80%E as well as domain B, and other E proteins and subunits. This general expression system is disclosed in EP 290261 published 11 September 1988. Envelope protein subunits that represent N-terminal portions
- of truncated protein may also be secreted from the 10 homologous prM fusion. Other secretion signal peptides or secretion leader pre/pro peptides, such as those associated with invertase or acid phosphatase of S. cerevisiae or with glucoamylase of C. albicans or of A. niger or the bovine
- 15 chymosin prepropeptide secretion leader can also be used. Secretion leaders in general that occur at the amino terminus of secreted proteins and function to direct the protein into the cellular secretion pathway generically can be used. In general, the invention includes expression
- 20 systems that are operable in eucaryotic cells and which result in the formation of envelope protein or a subunit secreted into the medium. Preferably, nonviral control sequences are employed. Thus, useful in the invention are cells and cell cultures which contain expression systems 25
- resulting in the production and secretion of the E protein or subunit.

The properly processed E protein or subunit is recovered from the cell culture medium, purified, and formulated into vaccines. Purification and vaccine

30 formulation employ standard techniques and are matters of routine optimization; however, particularly advantageous and novel purification methods are described below. Suitable formulations are found, for example, in <u>Remington's</u> Pharmaceutical Sciences, latest edition, Mack Publishing Company, Easton, PA. In particular, formulations will include an adjuvant, such as alum or other effective

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adjuvant. Alternatively, the active ingredient and the adjuvant may be coadministered in separate formulations.

The active vaccines of the invention can be used alone or in combination with other active vaccines such as those containing attenuated forms of the virus. The vaccines may contain only one subunit as an active ingredient, or additional isolated active components may be added. Corresponding or different subunits from one or several serotypes or flavivirus types may be included in a particular formulation.

More specifically, the subunits utilized in the vaccines of the invention can include the E protein or subunits thereof from each or a subset of the four dengue serotypes, and/or E protein or subunits from alternative flavivirus such as Japanese encephalitis virus, tick-borne encephalitis virus and the yellow fever virus. Multiple subunits of the E protein can also be included, as well as, optionally, other viral proteins, particularly the NS1 protein. These alternative subunits derived from other proteins of the virus can be recombinantly produced in a manner similar to the E protein and subunits described

> Thus, typical combinations might include: 80%E of the four dengue serotypes plus 60%E of

25 JEV;

above.

80%E of JEV;

80%E of TBE plus 60%E of dengue;

80%E of the four dengue serotypes plus NS1 protein of four dengue serotypes;

30 80%E of two dengue seroptypes plus NS1 protein or fragment from JEV;

Domain B of JEV plus 80%E of TBE;

Domain B of YF plus 80%E of JEV plus 80%E of dengue type 2;

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60%E of JEV plus domain B of four dengue seroptypes plus NS1 or fragment of both dengue and JEV; 80%E of JEV plus NS1 of JEV;

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80%E of TBE plus NS1 of TBE 80%E of YF plus NS1 of YF.

To immunize subjects against dengue fever or other flavivirus infections, the vaccines are administered to the subject in conventional immunization protocols involving, usually, multiple administrations of the vaccine. Administration is typically by injection, typically intramuscular or subcutaneous injection; however, other systemic modes of administration may also be employed.

10 Although the technology is not as well developed, transmucosal and transdermal formulations are included within the scope of the invention as are effective means of oral administration. The efficacy of these formulations is a function of the development of formulation technology 15 rather than the contribution of the present invention.

Since the subunit vaccines elicit the production of neutralizing antibodies, the antibodies thus raised can themselves be used as passive vaccines. For production of passive vaccine, a suitable mammalian subject is immunized

with the E protein or subunit of the invention and antibodies are either recovered directly as a composition from the antisera or indirectly as monoclonal antibodies from immortalized B cells of the subject. For production of monoclonal antibodies, the conventional techniques of Kohler

25 & Milstein, for example, or treatment with Epstein Barr virus, are used in immortalizing peripheral blood lymphocytes or spleen cells and screening for antibodies immunoreactive with the immunogen. These antibodies may further be screened for their ability to effect plaque 30 reduction in infected sera or cultures.

The polyclonal antisera are generally subjected to purification techniques such as standard size separation and chromatographic techniques to obtain purified immunoglobulins. The recombinantly produced proteins of the invention are particularly useful affinity ligands for chromatographic purification. A multiplicity of techniques is available for the purification of immunoglobulins for use

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in passive vaccines. If monoclonal antibodies are to be purified from small volumes of a medium ascites fluid, a protein A affinity column is particularly useful. For larger volumes, additional standard chemical techniques, such as ammonium sulfate precipitation and DEAE

5 such as ammonium sulfate precipitation and DEAE chromatography can be used. These immunoglobulins or the monoclonal antibodies generated by the immortalized B cells are then used in vaccine formulations as is understood in the art.

As is the case with active vaccines, the passive vaccines of the invention may be used in combination with additional antibodies or tandem administration of the antibodies of the invention and additional antibodies may be employed.

In the event that the passive vaccine is intended for use in a species other than that of the subject in which the antibodies were prepared, it may be desirable to modify the antibodies to minimize any immunogenic response. For example, it may be possible to use only the variable regions of these antibodies, such as the F_{ab} , $F_{ab'}$, or $F_{(ab')}$

regions. These fragments can be prepared either from polyclonal antisera or from the supernatants of hybridoma cultures by treating with proteolytic enzymes and recovering the desired fragments. The fragments are readily separated by using the relevant protein of the invention as an affinity reagent.

Alternatively, chimeric antibodies can be produced wherein the constant region corresponding to the species to be protected is substituted for the constant region

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characteristic of the species of antibody origin. The availability of recombinant techniques makes the production of chimeric antibodies a relatively trivial exercise. Briefly, a hybridoma or cell line producing the antibody of interest is used as a source for the genes encoding the antibody. The genes are recovered from, for example, the hybridoma using standard cloning procedures. The genes are then manipulated in vitro to remove the constant region and

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replace it with a constant region of a different species The modified genes are then ligated into expression origin. systems and expressed in recombinant host cells, such as CHO cells, monkey cells, yeast cells, and the like.

Further modifications in the variable regions can also reduce immunogenicity. Again, since recovery of the genes encoding the antibody is within the skill of the art, the variable regions, too, can be manipulated to replace the framework regions with framework regions more representative of the desired species, leaving intact the complementarily determining regions responsible for antigen specificity. In still another embodiment, the variable heavy chain and

variable light chain regions can be linked through a peptide linker and produced as a single chain F_v molecule.

Thus, if the passive vaccines are intended for humans, the foregoing various techniques of humanizing antibodies can be employed to minimize any immunogenic response even though the original antibodies are raised in nonhuman species.

In addition to use in vaccines or in the generation of passive vaccines, the mature recombinant E protein and subunits of the invention may be used as analytical reagents in assessing the presence or absence of antiflavivirus antibodies in samples. The interest in doing

this may be diagnosis of infection with flavivirus, 25 monitoring response to infection or may simply reside in the use of immunoassays as part of standard laboratory procedures in the study of the progress of antibody formation or in epitope mapping and the like. The antigen

30 is employed in standard immunoassay formats with standard detection systems such as enzyme-based, fluorescence-based, or isotope-based detection systems. Preferably, the antigen is used coupled to solid support or in sandwich assays, but a multiplicity of protocols is possible and standard in the 35 art.

Thus, the secreted protein, such as 60%E, 80%E or B domain may be adsorbed onto solid support and the support

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then treated with a sample to be tested for the presence of antiflavivirus antibodies. Unbound sample is removed, and any bound antibodies are detected using standard detection systems, for example, by treating the support with an antispecies antibody with respect to the species represented in the sample to be tested, the antispecies antibody coupled to a detection reagent, for example, horseradish peroxidase (HRP). The presence of the HRP conjugated antispecies antibody is then detected by supplying a suitable chromogenic substrate.

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Alternatively, the anti-subunit or antiflavivirus antibody may be adsorbed to the solid support and detected by treating the solid support with the recombinant domain B, either directly labeled, or labeled with an additional antibody in a sandwich-type assay.

In addition, both the mature peptides, such as domain B and 60%E or 80%E of the invention and the antibodies immunoreactive with it can be used in standard purification procedures as affinity reagents. Thus,

20 purification of the subunits from recombinantly produced cultures can be effected by affinity columns using antibodies raised against these antigens. Conversely, immunoglobulins useful in passive vaccines can be readily purified from antisera using the peptides of the invention.

The mature domain B or other subunit of the invention may be used to detect the presence or absence of antibodies of various isotypes, including IgG and IgM isotypes. As set forth above, detection of IgM isotypes is significant since this is an index of primary infection.

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In the examples below, particular subunits of the dengue Type 2 envelope protein, in particular 60%E, 80%E and domain B are illustrated as representative of effective subunits of the envelope protein in flavivirus in general. For the 60%E and 80%E constructs in general, secretion can be obtained from constructions designed to express the prME subunit fusion or under control of the human tissue plasminogen activator leader sequence. The mature

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N-terminus of the envelope protein is then secreted into the culture medium. Whether the N-terminus of the envelope protein subunits were fused to a heterologous leader, such as the human tissue plasminogen activator leader sequence, or to the homologous prM sequence or C-terminal portion thereof, the mature form of the truncated envelope protein

The secreted truncated Es are expressed at is secreted. high levels in Drosophila, efficiently processed, and secreted into the medium. The products are glycosylated and

- processed to an endo-H resistant form. 10 The secreted form of truncated E produced cotranslationally with prM generally represents about 20-30% of the total protein in the medium. Furthermore, based upon reactivity with conformationally sensitive monoclonal antibodies, using a ELISA and
- 15 immunofluorescence formats, the secreted E products are shown to have a native conformation. Immunization of mice with crude medium from transformed cells expressing prMtruncated E induces a potent virus-neutralizing response.

The following examples are intended to illustrate 20 but not to limit the invention.

Example 1

Preparation of Envelope Proteins in Saccharomyces cerevisiae

25 A cDNA clone derived from dengue serotype 2 (DEN-2) described by Hahn, Y.S. et al. Virology (1988, supra) was used as the starting material. This cDNA derives from strain PR159/S1. This strain has a small plaque, temperature-sensitive phenotype; the complete sequence of 30 the cDNA derived from the genomic RNA for PR159/S1 is set forth in this publication.

Figure 2 shows the sequence of the cDNA derived from genomic RNA of DEN-2 PR159/S1 for the Capsid, preMembrane, Envelope, and NS1 genes. Shown in bold at nucleotides 103, 1940, 1991, and 2025 are corrections to the Hahn published sequence. Differences in the S1 sequence from the wild-type sequence are noted above the wild-type

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There are no nucleotide differences in the Capsid sequence. and preMembrane protein-encoding portions and there are four in the E encoding portion.

Figure 3 shows the cDNA sequence of DEN-2 PR159/S1 5 for the Capsid, preMembrane, Envelope, and NS1 genes and the inferred translation of those four genes, which is part of the larger dengue polyprotein. The four differences between wild-type DEN-2 PR159 and the S1 strain are shown above the S1 nucleotide sequence. Also shown is the context of the 10 codons in which the nucleotide differences occur and the encoded amino acid.

In the E gene, three of the four mutations are silent; S1 has G instead of A at position 1314, T rather than G at position 1356, and C rather than T at position 2025. The mutation at position 1858, a G in S1 rather than A, results in a coding change from Ile in wild-type to Val There is, therefore, a one amino acid difference in in S1. the B domain in the S1 strain as compared to wild-type. Although the amino acid substitution conferred by the

- mutation at 1858 is conservative, mutagenesis studies of other viral structural proteins (Coller, B.G. et al. (1994) Mutagenesis Studies on the Predicted VP2 E-F Loop of Coxsackievirus B3, Abstract, 13th Annual Meeting of the American Society for Virology) have demonstrated that even relatively conservative single amino acid changes in surface 25
- loops can have profound effects on viral biology, including infectivity and viral stability. Thus, it is not without foundation that the conservative mutation in domain B of DEN-2 PR159/S1 may be involved in the attenuation of this 30 small-plaque, temperature-sensitive variant.

Subsequent determinations on the S1 strain used herein (S1/HBG) showed that the A at position 103 in wildtype was retained in position 103 in S1/HBG and C was substituted for T at position 2025 (a silent mutation). A deduced amino acid sequence from DEN-2 PR159/S1 and differences occurring in the wild-type are shown in Figure з.

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Various E gene subclones were obtained which represented the amino-terminal 90% of the envelope, 80% of the envelope, 60% of the envelope and classical domain B. Using the assignment of Mandl, C.W. et al. <u>J Virol</u> (1989)

<u>63</u>:564-571, classical domain B starts at Gly296 and ends at Gly395. Classical domain B contains only one disulfide bond and is encoded by the sequence following domains A and C. This classical domain B does not contain the potential T cell epitope described by Mandl *et al.* at its carboxy end
which can be included in some forms of the domain B of the invention, e.g., DomB+T.

The portion of the genome that encodes 80% of the envelope protein (80%E) was amplified using the Polymerase Chain Reaction, primers D2E937p and D2E2121m, and plasmid pC8 (Hahn et al. (1988, supra) as template.

In this notation of the primers, the virus serotype is first indicated (D2 for DEN-2), then the corresponding dengue gene -- i.e., in this case envelope, E, is noted. Then is noted the number in the dengue cloned 20 sequences of Figures 2 or 3 for the first dengue nucleotide in the 5'-3' direction of the oligonucleotide, i.e., using the numbering of Hahn *et al.* (1988, *supra*), and finally the notation shows whether the oligonucleotide primes the plus (p) or the minus (m) strand synthesis. The sequence in the 25 primers corresponding to dengue cDNA are written in uppercase letters; nondengue sequence is written in lowercase letters.

D2E937p2 (SEQ ID NO:13)

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<u>BglII</u> 5'-ct<u>tctagatctcgag</u>ta<u>cccggg</u>acc ATG CGC TGC ATA GGA ATA TC-3' XbaI XhoI SmaI

35 D2E2121m (SEQ ID NO:14)

<u>Sall</u> 5'-gc<u>tctaga</u>gtcga cta tta TCC TTT CTT GAA CCA G-3' 40 Xbal End End WO 96/37221

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The D2E2121m primer placed two stop codons after the 395th codon of E. The 80%E amplified cDNA fragment was digested at the XbaI sites in the cloning adapters and cloned into the NheI site of pBR322 to obtain p29D280E. Double-strand sequence for 80%E was determined, which identified a single silent PCR-introduced mutation at nucleotide 2001 (AAC/Asn to AAT/Asn).

A subclone representing domain B was obtained from the 80%E subclone by oligonucleotide-directed mutagenesis. 10 In the mutagenesis, stop codons and restriction endonuclease sites were inserted between domain C- and domain B-encoding sequences. The stop codons were positioned to terminate domain A+C translation and SalI and PvuII restriction sites were added to facilitate subcloning of domains A+C and domain B fragments, respectively, into yeast expression 15 vectors. As shown in Figure 4, to avoid a high AT content in the mutagenic oligonucleotide, the stop codons defining the carboxy-terminus of 60%E containing domains A and C were positioned four codons upstream of the beginning of domain

B, i.e., following Lys291. The original and altered nucleotide sequences of the mutagenized region and the corresponding amino acid translation are shown in Figure 4.

To perform the mutagenesis, a 580 bp BamHI fragment spanning domain B from the pBR322-80%E clone 25 p29D280E was subcloned into pGEM3Zf (Promega) to yield (See Figure 5.) This BamHI fragment encodes the p29GEB2. 3' end of domains A and C and all of domain B. The plasmid obtained following mutagenesis, p29GEB24PS, was confirmed by DNA sequencing. As shown in Figure 6, the introduced PvuII site permits direct subcloning of domain B cDNA into yeast expression vectors as a fusion with yeast secretion leader.

The cloned cDNA fragments encoding B domain and 80%E were inserted into expression vectors so as to maintain the translational frame of fusions to secretion leaders as described below. The sequence of the fusions were confirmed by DNA sequencing.

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Example 2

Secretion of Domain B from Saccharomyces cerevisiae

The expression vector constructed to secrete classical domain B from Saccharomyces cerevisiae to include 5 envelope protein amino acids 296-395 was constructed so that processing by the proteases normally involved in $preproMFa_L$ processing would yield a precisely processed domain B. One of these proteases, Kex2p, the gene product of the Kex2 gene, cleaves following dibasic peptides, such as LysArg, 10 LysLys, ArqArq, and ArqLys. A second protease involved, a dipeptidyl amino peptidase, removes GluAla and AspAla dipeptides from the amino terminus after Kex2p processing.

The MFa prepropeptide (MFa_{I}) -domain B fusion was operably linked to the constitutive promoter from the gene (TDH3) encoding glyceraldehyde phosphate dehydrogenase (GAPDH) contained in pLS5 and, alternatively, to the copper sulfate-inducible CUPI gene promoter contained in pLS6. These vectors were provided by SmithKline Beecham. Both contain the MF α_{I} , sequence, and StuI, BglII, and SalI cloning

- 20 sites, and use the TRP1 gene as a selectable marker. They contain sequences derived from pBR322 to provide an E. coli origin of replication, the ampicillin resistance gene, and sequences derived from the 2-micron plasmid of S. cerevisiae to enable replication in S. cerevisiae. The insertion of
- the classical domain B coding sequence into the appropriate reading frame in pLS5 was accomplished by digesting pLS5 with StuI and SalI and by digesting p29GEB24PS with PvuII and SalI and gel purifying the small PvuII-SalI domain B fragment and ligating it into the thus opened pLS5 as shown in Figure 6. A domain B expression vector derived from pLS6 30

The nucleotide and amino acid sequence of the $MF\alpha_L$ -domain B fusion is shown in Figure 7.

was constructed in a similar manner.

The resulting expression vectors were transfected 35 into spheroplasts of Saccharomyces cerevisiae strain GL43 (MATatrp1 Δ 1 ura3-52 proA::URA3) and transformants were

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selected by their ability to grow on minimal medium without tryptophan supplementation. S. cerevisiae strain GL43 was supplied by SmithKline Beecham. Strain GL43 pLS5-DomB and pLS6-DomB transformants were grown in small (<5ml) cultures i Casamino acid-supplemented minimal medium, and proteins

- secreted into the culture medium and cellular proteins were analyzed by Coomassie-stained SDS-PAGE and Western blot. Α novel Coomassie-staining band of approximately 12 kD, which was weakly immunoreactive with DEN-2 HMAF, was observed in
- the culture media of domain B transformants. No novel 10 Coomassie-staining bands and little or no immunoreactive protein not found in negative controls was observed in cellular protein extracts of domain B transformants. When cultured in Casamino acid supplemented minimal medium, 15
 - comparable levels of domain B were secreted into the medium by pLS5-DomB and pLS6-DomB transformants. About 20% of the total secreted protein was domain B as estimated by the intensity of Coomassie brilliant blue staining of protein in an SDS-PAGE polyacrylamide gel.

The composition of the medium was optimized; when S. cerevisiae containing pLS5-DomB were grown in minimal medium (SD) or Casamino acids (Difco)-supplemented minimal medium, the yields of protein were less than when cultured in rich (YEPD) medium. The pLS5-domain B transformant was preferred since this transformant can be cultured in rich medium whereas pLS6-domain B transformants require the use of the less preferred supplemented minimal medium in order to induce with copper sulfate.

The secreted protein was confirmed as domain B 30 having a Glu-Ala dipeptide appendage to the N-terminus, designated herein Glu-Ala-Domain B or Glu-Ala-DomB. For this demonstration, proteins secreted by a pLS5-DomB transformant were separated by SDS-PAGE and electroblotted to Immobilon P membrane (Millipore), and the amino terminal 35 amino acid sequence was determined by microsequencing. That sequence is: H2N-Glu Ala Gly Met Ser Tyr Ser Met Xxx Thr Gly Lys Phe Xxx Val Val (SEQ ID NO:15). The persistence of the GluAla

The domain B+stem cDNA fragment was constructed 40 in E. coli cloning vectors by combining the domain B cDNA

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dipeptide indicates that the dipeptidyl aminopeptidase incompletely processed the domain B N-terminus following proteolysis by Kex2p.

Example 3

Purification of Domain B

A. <u>Initial Purification Method</u>:

Pilot purifications of domain B from *S. cerevisiae* culture medium used size exclusion chromatography on 10 acrylamide Biogel P100.

For these purifications, an S. cerevisiae GL43 pLS5-DomB transformant was cultured in minimal medium overnight and then transferred to either Casamino acidsupplemented minimal (SD) medium or rich (YEPD) medium for

- 15 expression. After culturing at 30°C for 2-3 days and feeding daily with buffered glucose to final concentrations of 0.4% glucose and 5 mM sodium phosphate (pH 6.7), the cells were pelleted by centrifugation and the medium was clarified by filtration through a 0.45 µm pore filter. The filtered
- 20 medium was concentrated about 30-fold either by tangential flow or by centrifugal ultrafiltration using Minitan (Millipore) or Centricon (Amicon) devices, and the concentrate was exchanged into PBS azide buffer (2.7 mM KCl, 1.2 mM KH₂PO₄, 138 mM NaCl, 4.3 mM Na₂HPO₄•7H₂O, 0.02% azide) 25 by diafiltration or dialysis.

In one example, the concentrated secreted proteins from approximately 500 ml of culture of GL43 pLS5-DomB grown in SD plus Casamino acids were loaded onto a 2.5 x 75 cm Biogel P100 column and domain B was eluted from the column at approximately 1 bed volume. Domain B was pooled based on SDS-PAGE analysis of column fractions. The pooled domain B fractions were brownish in color and could not be decolorized by dialysis. The brown colored preparation also was not immunoreactive in ELISAs. DEAE chromatography of this sample resulted in binding of the brown color and a mostly colorless flow-through containing domain B. For DEAE

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chromatography, the brown-colored pooled fractions were dialyzed against 0.1 M acetic acid, pH 5.1 and loaded onto a 1.4 x 15 cm DEAE (Biorad) column. Domain B was eluted using a 0.01 M acetic acid, pH 5.1, 0.1 M NaCl step gradient.

The purified domain B was of high purity as assessed by silver stain on SDS-PAGE. The resulting Glu-Ala-DomB also gained weak reactivity with antidengue mouse polyclonal HMAF and with antidenque MAbs 3H5 and 9D12 in an ELISA format. The reactivities of earlier colorless preparations from smaller cultures were greater than this 10 initial large-scale preparation. For the ELISA, 96-well microtiter plates were coated with domain B by overnight incubation at 4°C and then blocked with BSA in 50 mM tris-HCl, pH 7.0, 0.15 M NaCl, 0.05% Tween for 1 hour at room 15 temperature. The plates were then treated with either DEN-2

HMAF or monoclonal antibody. Bound antibody was detected with alkaline phosphatase conjugated goat antimouse IgG and measured by the increase in optical density at 414 nm using the alkaline phosphatase substrate paranitrophenylphosphate.

The domain B frequently eluted from the Biogel P100 in two equal fractions, the first with the bulk of the secreted yeast proteins and the second as 70-90% pure domain Treatment with 1M NaCl, 1 and 2 M urea, and 1% DDAPS, a Β. zwitterionic detergent, were ineffective in completely

25 disaggregating domain B during size exclusion chromatography. Since the domain B of both fractions had the same electrophoretic mobility in the presence of detergent and absence of DTT, this polypeptide was neither covalently cross-linked to another protein nor self-

30 polymerized via disulfide bonds.

Β. Improved Purification Method:

A standard culture and purification method was developed that has reproducibly yielded immunoreactive 35 domain B expressed and secreted by S. cerevisiae GL43 pLS5-DomB. For that method, a primary culture is prepared by inoculating 60 ml of SD medium in a 250 ml baffled flask

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with a single colony of yeast strain GL43 pLS5-DomB grown on an SD plate. SD medium is prepared by autoclaving 0.67% Bacto yeast nitrogen base without amino acids, cooling and providing 2.0% w/v dextrose and 12 mM K_2HPO_4 using autoclave stock solutions. Solid SD medium is prepared by including Bacto agar at 1.8%. The culture is grown at 30°C for 18-24 hours with shaking at 240 rpm.

Secondary cultures are then grown in 300 ml of SD medium supplemented with 0.2% casamino acids in 2 L baffled flasks. The secondary cultures are inoculated with 30 ml of the primary culture. The secondary cultures are incubated at 30°C for 18-24 hours with shaking at 300 rpm.

Large-scale tertiary cultures are then prepared by inoculating 1 L of YEPD medium in a 4 L baffled flask with 15 100 ml of the secondary culture. YEPD is prepared by autoclaving 1% yeast extract plus 2% Bactopeptone, to which sterile dextrose is added to a final concentration of 2% after cooling. The large-scale cultures are incubated at 30°C for 48 hours with shaking at 300 rpm. After 24 hours, 20 the culture is supplemented with 0.01 vol of sterile 40% w/v glucose and 0.02 vol of sterile 1 M phosphate buffer, pH 6.7.

After growth in the 1 L culture, the cells are removed by centrifugation at 5,000 rpm at 4°C for 5 min. in a Sorvall GS-3 rotor. EDTA and EGTA are added to final concentrations of 1 mM each to the cleared medium, and the resulting solution is filter sterilized using a 045 μ m pore filter membrane (Millipack-20 or Opticap-50, Millipore). Glycerol is added to 10% v/v to the filtrate which is then concentrated 20-30 fold using tangential flow

ultrafiltration (Millipore Minitan System) with two membrane cartridges (four regenerated cellulose membranes) of a 10 kD MW cutoff. The retentate is kept on ice during ultrafiltration in the coldroom at 4°C for about 10-15 hours. The concentrated medium is directly used or

rapidly frozen on dry ice/ethanol in 10 ml aliquots in 15 ml

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polypropylene or polystyrene tubes with a screw cap and stored at -70° C.

Concentrated medium of the tertiary culture is dialyzed at 4°C using a membrane with a 6-8 kD cutoff (Spectra/Pore-1 Membrane) against 10 mM acetate, pH 4.5 (4X4 liters; 2-3 hours each; 1 overnight).

For purification, the concentrated, bufferexchanged medium is loaded onto a 2.5 x 22 cm CM Biogel A (Biorad) column at 0.8 ml/min. at 4°C. The column is washed with 150-250 ml of 10 mM acetate, pH 4.5 until the optical density at 280 nm of the eluent drops to baseline. Domain B is then eluted with 10 mM acetate, 300 mM NaCl, pH 4.5 at 0.8 ml/min. Fractions of approximately 8 ml are collected and analyzed by SDS-PAGE (15%) and visualized by silver staining. Domain B migrates on the trailing edge of the main peak of eluted material as monitored by 280 nm.

The domain B-containing fractions are pooled and concentrated about 11-fold by centrifugal ultrafiltration using, for example, a Centriprep-10 (Amicon) and loaded onto a 5 x 60 cm Sephadex G-75 (Superfine, Pharmacia) column at 4°C. The column is eluted using phosphate-buffered saline (PBS: 8 g/l NaCl, 0.2 g/l KCl, 1.44 g/l Na₂HPO₄, 0.24 g/l KH₂PO₄) at a flow rate of 0.8 ml/min. Nine ml fractions are collected and analyzed by SDS-PAGE as above and by indirect ELISA as above using Mab 9D12.

The Sephadex G-75 fractions containing pure immunoreactive domain B from 2-4 purification runs are concentrated about 50-fold by centrifugal ultrafiltration using a Centriprep-10 (Amicon; final volume 2-5 ml). The pooled material is stored at 4°C.

Glu-Ala-DomB purified as above typically is immunoreactive with the virus-neutralizing monoclonal antibodies 3H5 and 9D12. The above-described culturing and purification protocol has been performed several times with reproducible results. The number of cultures of the aboveindicated volumes can be increased, and we have grown

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simultaneously up to six 1 L tertiary cultures. The culture medium from the multiple cultures is pooled for clarification and concentration, and domain B from the concentrate is purified in batches and pooled following

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concentrate is purified in batches and pooled following verification of immunoreactivity. The amount of protein in the purified domain B preparations may be quantitated by total amino acid analysis using HCl hydrolysis with on-line ninhydrin reaction and optical detection. Table X presents the yield of protein in purified domain B from four cultures.

Table X Total Protein Yields in Purified Domain B Preparations				
Lot No.	Culture Volume	Final Yield*	Yield mg/l	
198.76	850 ml	4 mg .	4.7	
198.71	1500 ml	35 mg	23	
244.68	5750 ml	90 mg	16	
244.190	5000 ml	25 mg	5	

Based on amino acid composition of total protein.

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Example 4

Production of Domain B in Pichia pastoris

In addition to Saccharomyces cerevisiae as used in the previous examples, the methylotrophic budding yeast *P*. *pastoris* was used as a host. The domain B coding sequence was subcloned into the expression vector pPIC9 (Invitrogen, San Diego, CA) which contains the alcohol oxidase (AOX1) promoter sequence from *P*. *pastoris*, the MFa prepropeptide secretion leader from *S*. cerevisiae, the AOX1

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transcriptional termination sequence, and the HIS4 gene from S. cerevisiae as a selectable marker. To construct pPIC9-DomB, the domain B-encoding XhoI-SalI fragment from pLS5DomB was inserted into XhoI-digested pPIC9 as shown in Figure 8. XhoI cuts at identical positions within the MF α_L -encoding DNA of both pLS5 and pPIC9, and SalI cuts immediately

30 following the translational stop codons following domain Bencoding sequences. After verification by DNA sequence 5

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analysis, the expression cassette, consisting of the AOXI promoter-MF α_L -domain B-AOXI terminator, plus HIS4 was released from pPIC9-DomB by BglII digestion and the linear DNA was transformed into competent *P. pastoris* cells according to the supplier's instructions (Invitrogen, San Diego, CA). Transformants were selected for the ability to grow on medium lacking histidine (His[±] phenotype), and specific integration of the transforming DNA at the AOX1 locus was indicated in some transformants by slow growth on medium containing methanol (Mut[§] phenotype) as the sole carbon source.

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The expression and secretion of domain B from eleven P. pastoris pPIC9-DomB transformants in 3 ml cultures was surveyed by Coomassie-stained SDS-PAGE analysis. Α 15 protein of molecular weight equal to domain B secreted by S. cerevisiae was evident in the culture media of ten of the eleven P. pastoris transformants. Immunoblots of a comparable polyacrylamide gel demonstrated that the protein secreted by P. pastoris pPIC9-DomB transformants was 20 immunoreactive with polyclonal antibodies made to domain B secreted by S. cerevisiae (see Example 6). The SDS-PAGE analysis indicated that P. pastoris secreted domain B at a level comparable to or higher than that achieved in S. cerevisiae.

For further comparison of the secreted expression levels of domain B by S. cerevisiae and P. pastoris, the S. cerevisiae transformant and the two best P. pastoris transformants, Nos. 5-8 (Mut[±]) and 6-16 (Mut^B), were cultured in 100 ml shake flask cultures. S. cerevisiae pLS5-DomB was cultured 48 hrs in YEPD medium, and 100 ml precultures of the P. pastoris pPIC9-DomB transformants were grown in BMGY medium (1.34% yeast nitrogen base without amino acids, 2% Bacto peptone, 0.4 μ g/ml biotin, 1% glycerol, and 100 mM potassium phosphate, pH 6.8) for 24 hrs and then harvested and resuspended in 25 ml of BMMY heterologous proteininducing medium (the same as BMGY, except with 0.5% methanol replacing the glycerol) and cultured 48 hrs. Culture media 5

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with cells removed by sequential centrifugation and filtration through a 0.45 µm pore size membrane were buffer exchanged by diafiltration into TEEN (10 mM Tris, pH 8.0, 1 mM EDTA, 1 mM EGTA, and 150 mM NaCl). The relative levels of domain B secreted by the three cultures were estimated by size fractionating the proteins in 3, 6, 9, and 12 μ l of each sample on a 15% SDS-PAGE gel and staining with Coomassie blue. As estimated by the intensity of Coomassie staining, P. pastoris secreted at least four-fold more domain B per volume of induction culture medium than S. cerevisiae in YEPD. Correcting for the four-fold concentration of culture volume of the P. pastoris when transferred into the induction medium, we conclude that P. pastoris secretes slightly more than S. cerevisiae in shake flask cultures on a per volume basis. However, on a per gram cell dry weight basis, P. pastoris (0.80 gm total cell dry weight) secreted approximately 1.6-fold more domain B than S. cerevisiae (1.33 gm total cell dry weight).

Example 5

Effect of Extending Domain B

In a manner similar to that set forth in the preceding examples, two plasmid constructs for expression in S. cerevisiae were made that include additional domain B-25 carboxy terminal sequence downstream of amino acid 395 with the domain B coding sequence. One construct, designated DomB+stem, represents Gly296-Gly445 and was expressed as a fusion with the MFa_L . This construct includes those sequences that lie between domain B and the transmembrane 30 anchor of E. This region contains a potential T cell epitope (Mandl et al. <u>J Virol</u> (1989) <u>63</u>:564-571) and additional hydrophobic sequences, a peptide from this region elicits virus-recognizing antibodies (Roehrig et al. 1922 in <u>Vaccines</u> 92, pp. 2777-2281), and this region may contribute 35 to the proper folding and presentation of the domain B Bcell epitope to the immune system.
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fragment and the 3' end of a 90%E clone. As introduced in Example 1, an E gene subclone representing the amino terminal 90% of E was constructed from DEN-2 PR159/S1 cDNA plasmid pC8 of Hahn et al. (1988, supra) using the PCR. The 90%E polypeptide contains all of E except for the C-terminal membrane anchor comprising two transmembrane domains. The 90%E cDNA clone was made as follows. The 90%E fragment was amplified by the PCR using pC8 as template and primers D2E937p and D2E2271m. The sequence of D2E937p is given in Example 1. The

sequence of D2E2271m is:

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D2E2271m (SEQ ID NO:16):

<u>Sall</u> 5'-gc<u>tctaga</u>gtcga cta tta CCC GTA GAT TGC TCC G-3' *Xbal* End End

Primer D2E2271m placed two stop codons after Gly445 of #, and the two primers positioned useful restriction enzyme sites at both ends of the fragment. The PCR-amplified 90%E cDNA fragment was made blunt at both ends and cloned into the *SmaI* site of a modified pUC13 cloning vector, yielding pVZ90E.

Combining domain B and the 90%E 3' end made use of a unique AfIIII restriction enzyme site found in most pUC-like cloning vectors and a unique AfIIII site in domain B sequences. This combining was accomplished by first subcloning the 90%E fragment from pVZ90E into pBluescript to reverse the orientation of 90%E relative

- 30 to the vector sequences, yielding pBS90E. Then, p29GEB24PS, containing domain B sequences in pGEM (Example 1), and pBS90E were digested with Af1III, and the vector-domain-B5' fragment and the domain-B3'-stemvector fragment from the two digestions, respectively,
- 35 were purified, ligated, and recovered in *E. coli* yielding pBS-Bstem.

For expression in S. cerevisiae, the PvuII-SalI domain B+stem fragment from pBS-Bstem was subcloned into the

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pLS5 yeast expression vector at its StuI and SalI sites, putting the MF α_L and B+stem in translational frame. The sequence of the ligated junctions were verified by DNA sequence determination. The domain B+stem expression vector, pLS5-BStem, was transformed into *S. cerevisiae* strain GL43. The resulting transformants grew more slowly than did transformants with pLS5-DomB containing classical DomB and the plasmid appeared to be rapidly lost during growth under nonselective conditions. The DomB+stem product may be toxic to the cells, and the hydrophobicity of the stem may inhibit secretion.

However, a similar construct representing Gly296-Ala413 analogously inserted into pLS5 resulted in substantial amounts of secreted domain B protein. This form of domain B, designated herein DomB+T, contains the downstream T cell epitope. As stated above, the DomB proteins of the present invention include classical DomB and all or part of the amino acid sequence downstream from Gly395 to the Ala residue at position 413.

The DomB+T cDNA fragment was amplified by the PCR from pC8 plasmid DNA using oligonucleotide primers D2E1822p and D2E2175m. The primer sequences are:

D2E1822p KpnI XhoI (SEQ ID NO:17)

5'-cta gcg gta ccc tcg aga AAA GGG AGG CCG GGA TGT CAT ACT CCA TGT GC-3'

D2E2175m Sall NotI (SEQ ID NO:18)

5'-cgt gtg tcg acg cgg ccg cta tta GGC CAT TCT TTT CGC TCC-3'

The PCR product was cloned into pUC18 following KpnI and 35 SalI digestion of the product. The DomB+T cDNA fragment was then released from pUC18-DomB+T by XhoI-SalI digestion and cloned into the XhoI site in pLS5 within the MFα_L and Gly395 of domain B+T. S. cerevisiae strain GL43 secrets approximately ten-fold more domain B than domain B+T.

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<u>Example 6</u>

Preparation of Antibodies to the Recombinant Domain B Protein

The Glu-Ala-DomB peptide which had been column-5 purified, denatured by heating in SDS, and reduced by treatment with DTT was used to immunize Swiss Webster mice following removal of the excess SDS. The mice yielded hightitre antibodies that were highly immunoreactive when used to probe Western blots displaying the same antigen or DEN-2 10 viral envelope protein.

Example 7

Recombinantly Produced Glu-Ala-Domain B in ELISA Assays

15 The serological diagnosis for dengue infection is based on the detection of anti-dengue IgM and IgG in primary and secondary viral infection using standard Enzyme Linked Immunosorbent Assay (ELISA) techniques. Current assays are based on the ability of anti-dengue immunoglobulins to 20 recognize semi-purified virus. Primary and secondary infections can be distinguished by the IgM:IgG ratios. (Innis et al., 1989; Kuno et al., 1991).

The recombinantly produced Glu-Ala-domain B purified as in paragraph B of Example 3 was tested as an 25 antigen preparation in both IgM and IgG tests. In these ELISAs, the antigen was coated on plates, followed by sera positive for dengue antibodies and then detection by goat antihuman antibody.

Microtiter plates were coated overnight at 4° C
30 with 100 µl of a 5 µg/ml solution of domain B. After
blocking with 3% normal goat serum, primary infected dengue
2 positive serum at a 1:100 dilution was added per well.
The bound IgM was detected by the addition of goat antihuman IgM (Fc) conjugated to horseradish peroxidase. Under
35 these conditions, the recombinant Glu-Ala-domain B did not
provide positive results.

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For IgG, high titer secondary dengue 2 infected sera were supplied at a 1:270 dilution and detected with goat anti-human IgG (Fc) conjugated to horseradish peroxidase. Two of the five sera gave positive reactions in this assay.

The recombinant Glu-Ala-domain B purified by the improved procedure of Example 3 is not recognized by murine polyclonal hyperimmune ascitic fluid (HMAF) to other dengue serotypes and to other flaviviruses, when assayed using a 10 similar ELISA format. Flavivirus infected murine sera tested include, Japanese Encephalitis virus, Tick-Borne Encephalitis virus, Yellow Fever virus, Saint Louis Encephalitis virus, West Nile virus, three viral isolates of dengue serotype 1, two viral isolates of dengue serotype 3, 15 and two viral isolates of dengue serotype 4.

Sandwich assay for the detection of any domain B-containing envelope antigen.

In an alternative enzyme immunoassay format, antidomain B or anti-Dengue capture antibody, polyclonal or monoclonal, may be absorbed to the solid support, and sample containing an unknown quantity or serotype of dengue antigen may be added and then detected by reacting with a second anti-domain B or anti-Dengue antibody, either conjugated to

25 a signal-generating enzyme or to be detected using a appropriate signal generating system, of which there are a multitude. This immunoassay is useful for the quantitation of recombinantly produced envelope protein or whole virus by comparing the immunoreactivity of a known concentration of

30 domain B with that of the unknown antigen preparation. In addition, the conformational sensitivity or epitope binding site of the anti-domain B capture antibody can be varied to garner additional information regarding the conformation of dengue protein in the preparation.

To perform the sandwich enzyme immunoassay, 100 μ 1 of anti-Dengue monoclonal antibody 9D12 or 3H5 (Henchal, E.A. *et al.*, <u>Am J Trop Med Hyg</u> (1985) <u>34</u>:162-169) was used

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to coat microtiter wells. The monoclonal antibodies were purified by Protein-A affinity chromatography and used at 10 µg/ml concentration (diluted in PBS: 50 mM sodium Phosphate, pH 7.0, 0.15 M NaCl). After a one hour incubation, the wells were washed three times with TBS-T (50 mM Tris-HCl pH 7.0, 0.15 M NaCl, 0.05% Tween-20), and blocked with 200 μ l/well of 1% BSA in PBS for 1 hour at room temperature. Following three washes with TBS-T, 100 µl per well of a standard purified solution of domain B or an unknown sample was added to each well and captured by the bound monoclonal antibody. The domain B solution was allowed to bind the antibody for one hour at room temperature, and after three washes with TBS-T, 100 µl of Protein A affinity purified polyclonal rabbit anti-domain B serum (25 μ g/Ml) diluted in PBS-T (PBS, 0.25% BSA, 0.05% Tween-20) was added to each well and allowed to bind to the bound antigen. Following three washes with TBS-T, 100 µl of alkaline phosphataselabeled goat anti-rabbit IgG+IgM (H and L) conjugate

titration) was added to each well. After incubation for one 20 hour at room temperature, the plates were washed four times with TBS-T and 200 μ l/well of 1 mg/ml p-nitrophenylphosphate (pNPP) substrate in alkaline phosphatase substrate buffer (25 mM Trizma base, pH 9.5, 0.15 M NaCl, 5 mM MgCl₂; 0.02%

(Caltag) diluted 1:2500 in PBS-T (previously established by

25 NaN_2) was added. The plates were incubated for one hour at room temperature and the absorbance of each well was measured on a Dynatech MR5000 using a sample wavelength of 410 nm and a reference wavelength of 630 nm. A typical domain B standard curve is presented in Figure 12.

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Example 8

Immunization of Mice to Raise Neutralizing Antibodies

Crude DomB Immunogenicity in Mice: Media from

yeast cultures secreting DomB and a negative control yeast 35 culture were buffer-exchanged with PBS, and total secreted proteins were concentrated for injection into mice.

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groups of five mice each (Outbred Swiss strain, Simonsen) were inoculated with the crude DomB preparation, negative control secreted protein, or saline, as listed below.

Group	Immunogen	Adjuvant
1	Saline	Freund's
	Saline	none
	control medium	Freund's
IV	control medium	none
V	DomB medium	Freund's
VI	DomB medium	none

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Primary immunizations consisted of 50 µg of the immunogen diluted in phosphate buffered saline, with or without complete Freund's adjuvant as indicated above. Three boosts of 25 µg each were delivered at two-week intervals with no adjuvant (Groups II, IV, and VI) or Freund's incomplete adjuvant (Groups I, III, and V). Test bleeds were obtained one week after the first and second boosts, and the mice were bled out one week after the third boost.

- 15 Western blot and ELISA analysis of the first test bleeds confirmed a strong immune response to DomB among the Group VI (DomB, no adjuvant) mice. After the second bleed, this response was found to be titratable to greater than 1:6,400 by ELISA. Group VI was the only group that
- 20 developed high titer DomB antibodies. The mice immunized with DomB in Freund's adjuvant (Group V) mounted a weak immune response; based on reactivity to the immunogen in Western blot format and DomB-ELISA antibody titers in the 100-400 range, Group V mice appeared to be immunosuppressed.
- 25 The control mice (Groups I-IV) were all negative for antibody to DomB in both the Western and ELISA screens. In spite of high DomB antibody titers in Group VI mice, plaque reduction neutralization tests (PRNT) revealed that none of the mice immunized with the crude DomB produced neutralizing 30 antibody (Table 1).

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Table 1								
Plaque Reduction Neutralization Tests of Sera from Mice								
Immunized with DomB in Concentrated, Total Secreted Yeast Proteins								
Group	Trial			Dilu	tion			
				Plaque	Counts	,,, _,, ,, _,, ,, ,, ,, ,, ,, , _,, ,, ,, , _, ,, ,, ,, ,, ,, , _, ,, ,, ,, ,, ,, , _, ,		
PBS +		1:100	1:200	1:400	1:4	1:8	1:16	
Freund's Adjuvants								
	1	109, 105	~100	~100				
	2				34, 33	38, 30	44, 44	
PBS		1:10	1:20	1:40	1:4	1:8	1:16	
	1	100, 106	~ 100	~100				
	2				29, 33	33, 40	39, 37	
Negative secreted proteins +		1:10	1:20	1:40	1:4	1:8	1:16	
Freund's								
	1	106, 95	~ 100	~ 100				
	2				40, 35	37, 39	39, 47	
Negative secreted proteins		1:10	1:20	1:40	1:4	1:8	1:16	
	1	109, 94	~ 100	~100				
	2				38, 42	44, 40	45, 42	
Crude DomB+ Freund's		1:10	1:20	1:40	1:4	1:8	1:16	
	1	100, 92	~100	~100				
	2				37±5°	35 ± 6	39±3°	
Crude DomB		1:10	1:20	1:40	1:4	1:8	1:16	
	1	105, 91	~ 100	~ 100				
	2				39±3°	41±3	39±3°	
DEN-2 HMAF		1:100	1:200	1:400	1:800	1:1600	1:3200	
	1	0, 0	0, 0	0, 0	11, 8	31, 19	81, 75	
					1:250	1:1000	1:4000	
	2		·		3,6	12, 15	21, 21	

PRNT assays performed on VERO cells (Trial 1) or BHK-21 C15 cells (Trial 2) with DEN-2 NGC strain. All plaque counts indicate the number of plaques obtained with the sera from five animals were pooled and assayed in duplicate, <u>except</u> those indicated (*) where each serum sample was individually assayed in duplicate and the number of plaques averaged (mean \pm SD).

B. <u>Pure DomB Immunogenicity in Mice</u>: In contrast to mice immunized with crude DomB, outbred ICR mice (Charles River) immunized with purified DomB demonstrated high titer DEN-2 virus neutralizing antibodies. Purified DomB, at a concentration of 3.5 mg/ml, was used to immunize mice (three per group) with 175 μg of purified DomB mixed 1:1 with Freund's adjuvant, with alum, or without adjuvant (PBS). Test bleeds taken after three inoculations were assayed by PRNT (Table 2).

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Table 2 Plaque Reduction Neutralization Tests of Sera									
from Mice Immunized with Purified DomB									
Group	Tr ial		Dilution						
				Plaque	Counts				
Negative Ascites Fluid		1:10	1:20	1:40					
	1	36,40	35,34	40,38					
	2	39,38	47,45	41,44					
DEN-2 HMAF		1:100	1:200	1:400	1:800	1:1600	1:3200		
	1	0,0	0,0	1,0	2,1	8,9	13,14	1500	
	2	0,1	1,0	1,2	1,1	7,6	16,17	>1600	
DomB no adjuvant		1:10	1:20	1:40	1:80	1:160	1:320		
	1	36,29	28,36	29,27	36,30	40,25	28,31	>10	
	2	9,17	20,15	22,26	33,26	41,36	42,47	~10	
DomB alum		1:10	1:20	1:40	1:80	1:160	1:30		
	1	8,6	10,9	10,12	21,20	32,26	30,28	10	
	2	1,0	5,4	8,7	12,15	20,25	27,35	40	
DomB Freund's		1:40	1:80	1:160	1:320	1:640	1:1200		
	1	0,0	1,0	1,4	9,2	ND	ND	>320	
	2 0,0 3,4 5,5 5,20 13,9 23,20 ~640								

PRNT assays performed on VERO cells with DEN-2 NGC strain. ND = Not Determined.

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Mice immunized with purified DomB in the absence of adjuvant lacked neutralizing antibodies. DomB administered in combination with alum elicited low titer (80T PRNT ~1:10) neutralizing antibodies, and mice receiving purified DomB in combination with Freund's adjuvants had a high PRNT titer (80T PRNT >1:320).

Three groups of 3 mice each, 5-6 week-old outbred ICR strain (Charles River) were used. Inoculation was intramuscularly in the rump at one site using 10 µg of antigen in 0.1 ml administered solution. Three inoculations were given to each group on days 1, 20, and 43. In one group, inoculation on day 1

incomplete Freund's adjuvant, and that at day 43, no 15 adjuvant. In a separate group, no adjuvant was supplied and in a third group, alum was supplied with all three inoculations. The sera were withdrawn on day 57 and the sera from each group were pooled, heat-inactivated at 56°C for 30 minutes and tested for their ability to reduce 20 plagues formed from VERO cells.

incorporated complete Freund's adjuvants, on day 20

C. <u>DomB Protection from Virus Challenge</u>: Suckling mice were immunized with purified DomB in Freund's, Alum, Hunter's TiterMax (Vaxcel), or no adjuvant for protection against an intracerebral injection of DEN-2 New Guinea C (NGC) strain. DomB administered in all adjuvants conferred comparable moderate survival against dengue virus challenges although survival was statistically significant (P< 0.5 G test) only for mice immunized with DomB and Hunter's TiterMax. The results are shown in Figure 10.</p>

D. <u>KLH-DomB Immunogenicity in Mice</u>: Two series of mouse immunizations were initiated to determine the 50% effective immunizing dose (EID₅₀) of unconjugated and KLHconjugated DomB. Effects of alum and Freund's adjuvants to a no-adjuvant control are compared.

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DomB was conjugated to KLH via EDC (1-ethyl-3-(3-dimethylaminopropyl) carbodiimide HCl) using the carbodiimide coupling method at a 1.5:1 DomB-to-KLH mass ratio. The amount of unconjugated and conjugated DomB used for these immunizations was normalized, relative to the amount of unconjugated DomB. This normalization was based upon the specific immunoreactivity of each preparation as assayed by indirect ELISA.

Mice were immunized with the priming does of 10 174, 52, or 5.2 µg (total protein) of the KLH-conjugated DomB in Freund's, alum, or no adjuvant. Additional mice were immunized with 87, 26, or 2.6 µg (total protein) of unconjugated DomB in Freund's adjuvant to allow direct comparison of conjugated and unconjugated material.

Boosts consisting of one-half the priming dose are being given at two-week intervals. Test bleeds are assayed for the induction of anti-DomB antibodies by ELISA and Western blot. Final bleeds are tested for induction of a neutralizing immune response by PRNT assay as well as for the production of binding antibodies by ELISA and Western blot. The results are summarized in Table 3.

The response to unconjugated DomB with Freund's adjuvants was low compared to the results in Table 2, which had shown that unconjugated DomB induced a strong

- 25 virus neutralizing response in outbred ICR mice when administered with Freund's adjuvants. This apparent difference may be grounded in testing pooled sera for the data in Table 2. Pooling sera may mask individual variability. The variability in Table 3 may be
- 30 attributed to the limited epitopes in DomB and to differences in the MHC genes for the outbred Swiss mice used in Table 3.

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	Table 3 Mouse Immune Response to Unconjugated and KLH-Conjugated Purified Domain B								
mouse	antigen	adjuvant	Final Titer Western ^e	Final Titer ELISA ^c	Final Titer				
1			N/T ^a	<1:100	N/T				
2	saline	none	N/T	<1:100	<1:10				
3			N/T	<1:100	<1:10				
4			N/T	<1:100	<1:10				
5	87 μg B	none	N/T	<1:100	N/T				
6			N/T	<1:100	<1:10				
7			>1:100,000 Dblt ^b	>1:102,400	1:40				
8			N/T	<1:100	<1:10				
9	87 µg B	Freund's	1:1,000,000 Dblt	>1:409,600	<1:10				
10			N/T	<1:100	<1:10				
11			1:100,000 Dblt	1:25,600	<1:10				
12			N/T	<100	<1:10				
13			1:100,000 Dblt	1:102,400	<1:10				
14	26 µg B	Freund's	1:10,000 Dblt	1:102,400	N/T				
15			1:10,000 Dblt	>1:25,600	<1:20 ^c				
16			1:10,000 Dblt	1:409,600	<1:10				
17			1:1,000 Dblt	1:25,600	<1:10				
18			N/T	<1:100	<1:46				
19	2.6 µg B	Freund's	N/T	<1:100	<1:48				
20			N/T	<1:100	<1:24				
21			N/T	<1:100	<1:10				
22			N/T	1:100	<1:10				
23			<1:100	1:400	<1:24				
24	"87 μg" KLH-B	none	N/T	<1:100	<1:24				
25			N/T	<1:100	<1:24				
26			N/T	<1:100	<1:20				
27			N/T	<1:100	<1:34				
28			N/T	<1:100	<1:70				
29	"26 μg" KLH-B	none	N/T	<1:100	<1:34				
30			N/T	<1:100	N/T				
31			N/T	<1:100	<1:10				

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Mouse Immune Response to Unconjugated and KLH-Conjugated Purified Domain B							
mouse	antigen	adjuvant	Final Titer Western ^c	Final Titer ELISA ^c	Final Titer PRNT ⁶		
32	"2.6 Tg" KLH-B	none	N/T	<1:100	N/T		
33	1		N/T	<1:100	<1:10		
34	1		N/T	<1:100	<1:10		
35	1		N/T	<1:100	<1:10		
36	1		N/T	<1:100	<1:10		
37	"87 Tg" KLH-B	Freund's	N/T	>1:1,600	1:20		
38	1		N/T	>1:100	<1:10		
39	1		N/T	<1:100	<1:10		
40	1		1:1,000	>1:1,600	1:20		
41	1		N/T	<1:100	<1:10		
42	"26 Tg" KLH-B	Freund's	1:10,000 Dblt	>1:25,600	1:10		
43	1		1:10,000 Dblt	>1:25,600	1:2560		
44	1		1:10,000 Dblt	>1:25,600	1:5120		
45] [N/T	<1:100	<1:10		
46			N/T	1:100	<1:10		
47	"2.6 Tg" KLH-B	Freund's	N/T	<1:100	<1:10		
48	1		N/T	<1:100	<1:10		
49]		<1:50	1:6,400	<1:10		
50			N/T	<1:100	<1:10		
51			<1:50	1:25,600	<1:10		
52	"87 Tg" KLH-B	Alum	<1:100	<1:400	1:60		
53			N/T	<1:100	<1:10		
54			<1:100	>1:1,600	1:320		
55			<1:100	>1:1,600	1:80		
56			1:1000	>1:6,400	1:640		
57	"26 Tg" KLH-B	Alum	N/T	>1:100	<1:10		
58]		N/T	<1:100	<1:10		
59]		N/T	<1:100	<1:10		
60] · [N/T	<1:100	<1:10		
61			N/Ţ	<1:100	<1:10		
62	"26 Tg" KLH-B	Alum	<1:100	1:1,600	1:120		
63]		1:1000	>1:1,600	1:120		
64]		N/T	<1:100	<1:10		
65			1:100	>1:1,600	1:80		
66			N/T	<1:100	<1:10		

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^{*}N/T - not tested ^{*}Dblt - a doublet at approximately 12 kD ^{*}If serum was insufficient for testing at 1:10 dilution then higher initial dilutions were used and are indicated.

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DomB Immunizations for Hybridoma Generation: Ε. Six BALB/c mice were immunized with 87 μ g of unconjugated DomB or 174 µg KLH-conjugated DomB in Freund's adjuvant. Upon demonstration of a strong anti-DomB response, the mice are sacrificed and their spleen cells fused to hybridoma cells for monoclonal antibody production.

Example 9

Production of Domain B in Drosophila

10 The cloning vector p29GEB2.4PS, containing the domain B-encoding nucleotide sequence was digested with PvuII and the short fragment ligated into pMttSma, which is derived from pMttbns by digesting with BglII and inserting an adapter oligonucleotide containing an Sma The duplex linker adapter inserted has the site. sequence pGATCCCGG. pMttsma then contains a unique SmaI site at the 3' end of sequences encoding the tPA leader. The parent vector pMttbns SmithKline contains the Drosophila metalothionein gene (mtn), the human tissue 20 plasminogen activator secretion signal (tPAL) and the SV40 early polyadenylation signal. The resulting expression vector, pMttDomB is thus obtained as shown in Figure 11. The sequence of the fusion is shown in Figure

25 The resulting pMttDomB vector was transfected into Schneider S2 cells which were maintained in Schneider medium (Life Technologies, Grand Island, NY) supplemented with 10% heat-inactivated FBS at 25°C. The cells were plated the day prior to transfection at 5 x 10^5 cells in 60 mm tissue culture dishes in a total 30 volume of 4 ml Schneider medium plus 10% FBS. The Schneider cells were then cotransfected with pMttDomB and pCOHygro (a hygromycin B phosphotransferase-based selection vector obtained from SmithKline Beecham) using 35 standard calcium phosphate transfection. 48 hours after transfection, the medium was supplemented with hygromycin

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at 300 µg/ml; hygromycin-resistant cells grew in 2-3 weeks. Varying levels of the cotransformed plasmids were used in the transfection procedure ranging from expression plasmid: PCOHygro, 1:1 to 100:1.

Cells cotransfected at ratios of 1:1, 5:1 and 20:1 were induced with 200 μ M copper sulfate and the media and cells were harvested at days 1, 4 and 7. Western blots showed secretion of domain B into the medium.

Example 10

Production of 60%E and 80%E in Drosophila

Using the adapted expression vector of Example 9, containing the Drosophila metalothionein gene, the 15 human tissue plasminogen activator signal and the SV-40 early polyadenylation signal, the nucleotide sequences encoding 80%E, prM 80%E, 60%E and prM 60%E are inserted and the resulting vectors used to transform Schneider cells as described in Example 9. The mature truncated 20 forms of the envelope protein are secreted into the medium, or properly processed, and are conformationally correct with respect to the corresponding native portions of the envelope protein.

Furthermore, crude media from these 25 transformants employed in the protocol set forth in paragraphs A and B of Example 8 produce antibodies which are neutralizing against dengue virus.

Example 11 Expression of 80%E in Saccharomyces cerevisiae

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An expression vector (pLS6-80%E) was constructed for secretion of the N-terminal 80% (codons 1-395) of the DEN-2 PR-159 S1 envelope glycoprotein (80%E) from S. cerevisiae. The 80%E DNA sequences were obtained from plasmid p29D280E, described in example 1, by restriction endonuclease digestion with both BglII and SalI. The released fragment was isolated by agarose gel

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electrophoresis and subcloned between the *BglII* and *SalI* sites of pLS6, a yeast expression vector described in example 2. The resulting recombinant plasmid, pLS6-80%E, contains truncated E as a translationally in-frame fusion to the leader region of mating-factor α (MF α), a secreted

- 5 to the leader region of mating-factor α (MF α), a secreted yeast protein. The MF α leader (MF α_L) contains a 19 amino acid secretion signal peptide followed by a 66 amino acid propeptide. Cleavage of the signal peptide is expected to occur concomitantly with translocation across the
- 10 endoplasmic reticulum membrane. Maturation of MF α normally involves removal of the propeptide by Kex2p, a golgi protease, and subsequent trimming of N-terminal (Glu/Asp)Ala dipeptides by dipeptidyl aminopeptidase (DPAP). The herein described MF α_L -80%E fusion was made
- 15 such that processing of the MFα_L propeptide and trimming of a GluAla dipeptide results in 80%E with eight additional N-terminal amino acids derived from sequences present in the multiple cloning site of the pLS6 vector or in the PCR primer-adapter used to synthesize the 80%E 20 cDNA (see below).

ATGGCT......GAG GCC TTT AGA TCT CGA GTA CCC GGG ACC ATG ...GAA TAA TAG Met-18aa Ala-65aa Glu Ala Phe Arg Ser Arg Val Pro Gly Thr Met1...Gly395End 25 End A A A 80%E Signalase Kex2p DPAP

30 Transcription of the gene fusion is driven by the S. cerevisiae copper-inducible copper metallothionein (CUP1) promoter.

After confirming the DNA sequence of the ligated junctions of expression vector pLS6-80%E, the recombinant DNA was transformed into *S. cerevisiae* strain GL43 (*MATa* $trpl\Delta 1$ ura3-52 pep4::URA3; SmithKline Beecham) according to standard protocols (Gutherie & Fink, eds. 1991; Rose et al, 1990). Transformants were selected by their

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ability to grow on minimal medium (SD medium: Guthrie and Fink, 1991) without tryptophan supplementation.

In order to test for expression and secretion of 80%E, several transformants were grown as small-scale cultures (5 ml medium in 17x150 mm tubes). Single colonies were used to inoculate SD medium and the inoculated cultures were grown to saturation (overnight, 30° C, 220 rpm). 1x10⁸ cells from the overnight culture were used to inoculate 5 ml of minimal SD medium supplemented with Casamino acids (2 g/l; Difco) and CuSO₄ (200 μ M). This expression culture was fed with glucose (4 g/l, final concentration) and sodium phosphate (pH 6.7, 20 mM, final concentration) at 24 hours post inoculation and was harvested by centrifugation at 48

15 hours of growth (30°C, 220 rpm, in 17x150 mm tubes). Cell-free spent medium was buffer-exchanged with TEEN+PIC (50 mM Tris, 10 mM EDTA, 10 mM EGTA, 150 mM NaCl, pH 8.0 with 1 µg/ml each of pepstatin and leupeptin and 1 mM phenylmethylsulfonylfluoride) and concentrated 250-fold

20 using Centricon-30 (Amicon) ultrafiltration. An extract of cellular protein was prepared by lysing the yeast cells with vigorous agitation in the presence of glass beads (425-600 μm) and TEEN+PIC using a Mini Beadbeater apparatus (BioSpec Products, Bartlesville, OK). Samples

25 were endoglycosidase H_f digested according to the manufacturer's (New England Biolabs, Beverly, MA) protocol prior to SDS-PAGE analysis. Protein gels were Coomassie-stained directly as well as Western blotted and immunoprobed using mouse polyclonal serum raised against

30 recombinant domain B (described in example 15). Negative control yeast carrying the expression vector without a Dengue gene insert secreted no proteins recognized by the anti-domain B serum, while the major immunoreactive band from pLS6-80%E medium had a relative mobility matching 35 that of other recombinant 80%E proteins (see Example 17).

The pLS6-80%E medium also contained a minor

immunoreactive species with an apparent molecular weight 6-8 kD higher than 80%E; this is likely to be unprocessed MFα propeptide-80%E. The pLS6-80%E cellular protein extract contained many immunoreactive polypeptides not observed in negative control cells, two of which match the secreted products discussed above. A Coomassiestained protein corresponding to recombinant 80%E could not be identified in either the pLS6-80%E secreted sample or the total cellular protein sample. This indicates the relatively low levels of recombinant protein expression in these preparations.

Example 12

<u>Construction of expression vector pPIC9-80%E and</u> <u>secretion of 80%E by P. pastoris expressing MFa_L-80%E</u>

The expression vector constructed to secrete 80%E from *P. pastoris* was engineered to express amino acids 1-395 of the DEN-2 PR-159 S1 envelope glycoprotein as a fusion to the MF α_L . The DNA sequences encoding 80%E were obtained from the clone p29D280E, described in example 1, by digestion with the restriction enzymes *SmaI* and *SaII*. The isolated fragment was treated with the Klenow DNA polymerase I fragment enzyme to make the *SaII* end blunt. This 80%E fragment was then cloned into the

25 Pichia expression vector pPIC9 (Invitrogen, San Diego, CA) which contains the MF α secretion leader (MF α_L) sequence, SnaBI, EcoRI, and NotI cloning sites, and uses the HIS4 gene as a selectable marker. The described 80%E fragment was ligated with pPIC9 plasmid DNA that was

30 previously digested with the restriction enzyme SnaBI. The orientation and genetic integrity of the resulting gene fusion expression vector, pPIC9-80%E, was confirmed by restriction digestion and DNA sequence analysis. The organization, partial nucleotide and predicted amino acid 35 sequences of the MFα_L-80%E fusion gene are shown below:

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(SEQ ID NO:19 and SEQ ID NO:20) ATG GAG GCC TTTAGATCTCGAGTACCCGGGACC ATG ...GGA TAA Met-18aa-65aa Glu Ala PheArgSerArgValProGlyThr Met1...Gly395 END

5 Signalase Kex2p DPAP

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The location of the signalase and Kex2 cleavage sites which remove the pre and pro portions of the MF α leader peptide, respectively, are indicated. The dengue sequences are indicated in bold type. The Met₁ residue is the N-terminal amino acid of the E glycoprotein and Gly₃₉₅ is residue 395 from the amino terminal end of the envelope glycoprotein. The expression of a recombinant product in *Pichia* from the pPIC9 vector is driven by the methanol inducible promoter derived from the *Pichia AOX1*

(alcohol oxidase 1) gene.

The pPIC9-80%E expression vector was transformed into spheroplasts of P. pastoris strain GS115 (his4) and transformants were selected for their ability to grow on minimal medium without histidine supplementation. Strain GS115 and the protocol used for transformation were obtained from Invitrogen (San Diego, CA). Transformants were tested for their ability to express and secrete 80%E by growing selected clones in small cultures (5 to 50 ml). The transformants were first grown to saturation (24 to 36 hrs.) in BMGY medium (1% yeast extract, 2% peptone, 100 mM potassium phosphate, pH 6.0, 1.34% yeast nitrogen base without amino acids, 4×10^{-5} % biotin, 1% glycerol). The cells

- 30 were collected by centrifugation and suspended in one half the original culture volume with BMMY (identical to BMGY except the glycerol component of BMGY is replaced in BMMY with 0.5% methanol) medium and cultured for 48 hrs. Proteins secreted into the culture medium as well as
- 35 cellular proteins were treated with endoglycosidase H_f , (EndoH_f, New England Biolabs, Beverly, MA) and separated by SDS-PAGE. Protein gels were analyzed by both Coomassie staining and immunoprobing of Western blots. When nonEndoH treated samples are compared to EndoH



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treated samples on Coomassie stained gels of proteins prepared from the culture medium, a unique staining band of approximately 50 kD is present in the EndoH treated lanes of all the pPIC9-80%E transformants.

5 Immunoprobing with anti-domain B serum (see example 6) detected a smear ranging from 50 to 90 kD in the nonEndoH treated samples and a unique band of approximately 50 kD in the EndoH treated samples on Western blots of proteins prepared from the culture medium. No bands were detected

10 by the anti-domain B serum in negative control lanes. Variable amounts of the corresponding immunoreactive band were detected in EndoH treated samples of cellular protein samples. The approximately 50 kD 80%E product produced by the MFa_L-80%E construct is consistent with

- 15 the approximate molecular weight as determined by SDS-PAGE of other recombinant 80%E proteins (see Example 17). The amount of secreted 80%E in the culture medium is about 1% of the total secreted protein as estimated by the intensity of the Coomassie staining band detected.
- 20 In one liter cultures, the amount of 80%E secreted into the culture medium was determined to be 500 ng/ml by use of a sandwich ELISA method.

Example 13

25 <u>Construction of Expression vector pMttbns-80%E and</u> <u>secretion of 80%E by Drosophila melanogaster Schneider</u> <u>cells expressing tPA_L-80%E</u>

The expression vector constructed to secrete 80%E from Drosophila melanogaster tissue culture cells 30 included the sequences encoding the DEN-2 PR159/S1 envelope glycoprotein amino acids 1-395. The DNA sequences for 80%E were obtained from the clone p29D280E (described in Example 1) by digestion with the restriction enzymes BglII and SalI. The released 80%E 35 fragment was cloned into the BglII plus XhoI-digested D.

melanogaster expression vector pMtt Δ Xho. The expression vector pMtt Δ Xho is a derivative of the vector pMttbns

below:

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(SmithKline Beecham) that contains a *D. melanogaster* metallothionein gene promoter (*Mtn*), the human tissue plasminogen activator leader sequence (tPA_L), and the SV40 early polyadenylation signal. A 15 bp *Bam*HI DNA

- 5 fragment containing a XhoI site was deleted from pMttbns to make pMtt∆Xho, in which the BglII and another XhoI restriction endonuclease sites are unique. This construction resulted in the 80%E fragment being fused to the tPAL sequence. During normal maturation of tissue
- 10 plasminogen activator the 20 amino acid prepeptide region of the leader sequence is removed by signalase in the endoplasmic reticulum and the 11 amino acid propeptide region is enzymatically removed in the Golgi. The genetic integrity of the gene fusion expression vector, 15 pMtt80%E, was confirmed by restriction digestion and DNA sequence analysis. The nucleotide and predicted amino acid sequences of the tPAL-80E fusion gene are shown
- 25 The tPA pre- and propeptide regions are delineated by preand pro-tPA, respectively, and the dengue sequences are indicated in bold type. The Met₁ residue is the Nterminal amino acid of the envelope glycoprotein and Gly₃₉₅ is residue 395 from the amino terminal end of the anvelope glycoprotein.

The selection plasmid, pCOHygro (SmithKline Beecham), carries the *E. coli* hygromycin B phosphotransferase gene under the transcriptional control of a *D. melanogaster copia* transposable element long

35 terminal repeat and confers resistance to hygromycin B. Others have demonstrated that Schneider cells that have been cotransfected with both the pMtt expression vector and the pCOHygro plasmid and are then selected for

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hygromycin B resistance produce stable cell populations that contain multiple copies of the cotransfected gene of interest. Drosophila melanogaster Schneider cells (ATCC, Rockville, MD) were cotransfected with the pMtt80%E and pCOHygro plasmids at a ratio of 20:1 using the calcium 5 phosphate coprecipitation method (Gibco BRL). Transformants were selected by outgrowth in Schneider medium (Gibco BRL) supplemented with 10% fetal bovine serum (Hyclone) and 300 µg/ml hygromycin (Boerhinger Mannheim). Transformants were split to a cell density of 10 2 X 10⁶ cells/ml in serum free Excell medium (JRH Biosciences) and induced with 200 μ M CuSO₄. The medium and cells from induced cells were separately harvested after 6-7 days of induction. Proteins secreted into the culture medium were separated by SDS-PAGE, and proteins 15 were analyzed by both Coomassie staining and immunoprobing of Western blots. The Coomassie bluestained SDS-PAGE gels shows that the approximately 50 kD secreted 80%E product is one of the predominant proteins 20 in the unconcentrated medium, comprising as much as 20% of the total protein. Immunoblots probed with anti-DEN2 hyperimmune mouse ascites fluid (HMAF; from Robert Putnak, Walter Reed Army Institute of Research) and polyclonal anti-domain B antisera (Example 6) revealed a single immunoreactive polypeptide of approximately 50 kD 25 present in unconcentrated medium. In addition, immunoblots revealed that the 80%E produced by the tPAL-80%E construct was slightly larger than that obtained upon expression of a tPAL-prM90%E construct (described in detail in Example 17). This additional mass may owe to 30 the nine adapter amino acids at the amino terminus of 80%E (GARSRVPGT-80%E) (SEQ ID NO:23) when expressed from pMtt80%E versus 80%E expressed from pMttprM80%E (Example 17). The tPA propeptide, if not proteolytically removed, may also contribute to the additional molecular weight of 35

80%E expressed from pMtt80%E.





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<u>Example 14</u>

Subcloning of dengue prM100%E and prM80%E cDNAs and mutagenesis of E secretion signal-encoding sequence (mutSS)

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A cDNA clone of DEN-2 PR159/S1 designed to encode the preMembrane, Membrane, and Envelope genes (prM100%E) was constructed by PCR amplification essentially as described in Example 1 for the subcloning of 80%E. This cDNA clone includes nucleotides 439 to 2421 of the DEN-2 genome. The dengue cDNA fragment was generated using synthetic oligonucleotide primers D2pM439p and D2E2421m (see example 1 for nomenclature) and plasmid pC8 (Hahn et al, 1988, *supra*) as template. In addition to DEN-2 specific sequences, the primers contained the identical adapter sequences described in Example 1, except that a methionine codon (ATG) was

included immediately preceding the first codon of the preMembrane sequence (phenylalanine). The primers are:

20 <u>D2prM439p</u>(SEQ ID NO:24)

<u>Bg111</u>

5'-at<u>tctaga</u>t<u>ctcgag</u>ta<u>cccggg</u>acc atg TTT CAT CTG ACC ACA CGC-3' XbaI XhoI SmaI

25 <u>D2E2421m</u>(SEQ ID NO:25) <u>Sall</u> 5'-tc<u>tctaga</u>gtcga cta tta GGC CTG CAC CAT AAC TCC Xbal END END

30 The PCR-generated prM100%E cDNA fragment was digested with the restriction endonuclease XbaI and ligated into the XbaI site of pBluescript SK⁺ (Stratagene, San Diego, CA) to obtain the plasmid p29prME13. DNA sequence analysis of the PCR-generated

35 cDNA clone identified three PCR-introduced nucleotide differences between pC8 and p29prME13. A T→C transition at nucleotide 1255 resulted in a silent mutation, while the A→G transition at nucleotide 1117 resulted in the conservative amino acid substitution of a valine for an WO 96/37221

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isoleucine. The final transition observed at nucleotide 1750 replaced a methionine, adjacent to a cysteine involved in a disulfide bond in domain A, with a valine. The effect of this substitution on the ability of the disulfide to form or its stability is not known.

To generate a cDNA subclone representing prM80%E, a 794 bp BamHI-SalI fragment from p29prME13 representing the envelope carboxy terminal-encoding fragment was removed. This fragment was replaced with a

- 10 431 basepair BamHI-SalI fragment from p29D280E (described in Example 1) that encodes a 20% carboxy end truncation of the envelope glycoprotein. The BamHI site is a naturally occurring site within the envelope cDNA, and the SalI site is an engineered site that immediately
- 15 follows the stop codons encoded by the PCR primers. The resulting truncated cDNA clone, p48BSprM80E, was confirmed by restriction digestion and DNA sequence analysis to encode amino acids 1 through 395 of the envelope glycoprotein following prM.

20 Expression of the prM80%E cDNA in S. cerevisiae (Example 15) demonstrated absence of proteolytic processing between the prM and 80%E proteins in this yeast. To improve processing of E from prM, oligonucleotide-directed mutagenesis was performed to

- 25 alter the naturally occurring signalase cleavage site between the prM and E proteins. Based on the algorithm of Von Heijne (1986, Nucl. Acids Res. 14:4683-4690), the natural DEN-2 E secretion signal peptide receives a poor predictive score for its function as a secretion signal.
- 30 The algorithm of von Heijne is based on N-terminal amino acid sequences found in secreted and nonsecreted proteins. Scores for functional secretion signal peptides range from 3.5 to 17, with a mean of ~10. The score for the secretion signal peptide of E of DEN-235 PR159/S1 is 5.2, near the lower end of the range for signal peptides. In the mutagenesis, the sequence encoding the four amino acids immediately preceding and

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the amino acid immediately following the signalase cleavage site were altered to change these five amino acids. The modified signal sequence has a score of 12.4 based on von Heijne's algorithm. The original and altered nucleotide sequences of the mutagenized region and the encoded amino acid sequences relative to the signalase site are (SEQ ID NO:26 through SEQ ID NO:29):

Original ATC GCT CCT TCA ATG ACA ATG CGC TGC Ile Ala Pro Ser Met Thr Met Arg Cys Mutagenized ATC GCT GGC GCT CAA GCT CAA CGC TGC Ile Ala Gly Ala Gln Ala Gln Arg Cys Membrane | Envelope

Signalase Cleavage

The mutagenized sequence and the resulting amino acid changes are indicated in bold.

To perform the mutagenesis, a 1,122 bp SmaI-HindIII fragment spanning the prM-E signalase cleavage site from the p29prME13 cDNA clone was subcloned into pAlter1 (Promega, Madison, WI) to yield the plasmid pAltSmaH3prME. The 1,122 bp SmaI-HindIII fragment contains all of prM and 611 bp of the E sequence. The HindIII site is a naturally occurring site within the E sequence that is located at nucleotide 1547 of the genomic sequence. The mutagenized clone, pAltSmaH3prME(mutSS), was verified by DNA sequence analysis.

Example 15

<u>Construction of expression vectors pLS6-prM80%E and pLS6-prM(mutSS)80%E, expression of MFa_L-prM80%E and MFa_L-prM(mutSS)80%E in Saccharomyces cerevisiae, and secretion of 80%E by Saccharomyces cerevisiae expressing MFa_L-prM(mutSS)80%E</u>

For expression of DEN-2 PR159/S1 preMembrane protein amino acids 1-166 and Envelope glycoprotein amino

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acids 1-395 as a single continuous open reading frame in S. cerevisiae, DNA sequences encoding these proteins were obtained by digestion of plasmid p48BSprM80E (see Example 14) with restriction endonucleases BglII and SalI. This fragment was cloned into the pLS6 yeast expression vector (see Example 2) that had been digested with BglII and SalI. The structure of the resulting gene fusion expression vector, pLS6-prM80%E, was confirmed by restriction digestion and DNA sequence analysis. The nucleotide and predicted amino acid sequences of the MF α_L -prM80%E fusion gene are shown below (SEQ ID NO:30

and SEQ ID NO:31):

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BglII Small

- The location of the signalase, Kex2p, and DPAP cleavage sites involved in the processing of the MFα leader peptide are indicated (See Example 1 of this application for a detailed explanation). The dengue sequences are indicated in bold. The Phe₁ and Thr₁₆₆ residues are the N-terminal and C-terminal amino acid residues of prM, respectively. The Met₁ residue is the N-terminal amino acid of the envelope glycoprotein and Gly₃₉₅ is residue 395 from the amino terminal end of the envelope glycoprotein.

30 The pLS6prM80%E plasmid was transformed into Saccharomyces cerevisiae strain GL43 (MATa ura3-52 trp1 Δ 1 pep4:URA3) and screened for 80%E expression as described in Example 1. Proteins secreted into the culture medium as well as total cellular proteins were treated with 35 Endoglycosidase H_f (EndoH_f, New England Biolabs, Beverly, MA) prior to analysis by SDS-PAGE followed by Coomassie Blue staining or Western blot immunoprobing. A novel secreted protein could not be identified either on



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Coomassie Blue stained SDS-PAGE gels or Western blots probed with anti-DEN2 <u>hyperimmune mouse ascitic fluid</u> (HMAF; from R. Putnak, Walter Reed Army Institute for Research) or anti-DomB antiserum (see Example 6).

Western blot analysis of total cellular protein revealed a HMAF immunoreactive band of approximately 90 kD suggesting that the recombinant product had not been processed to prM and E. Probing of companion Western blots with polyclonal antisera that recognized the MFa

10 leader peptide (from J. Rothblatt, Dartmouth University) confirmed that the product recognized by the anti-DEN2 HMAF was identical to that recognized by anti-MF α serum, demonstrating that the MF α_L -prM80%E fusion protein was not processed into its individual components (MF α_L , prM, 15 and 80%E).

The unsuccessful processing of E from prM in the MF α_{L-} prM80%E fusion protein may be an obstacle to the proper folding and secretion of E. To assess whether the optimized dengue signal sequence (see Example 16) facilitated the processing of the envelope protein at the prM-E junction, the altered E signal sequence from

pLS6prM(mutSS)100E-TGA was introduced into pLS6prM80%E to create plasmid pLS6prM(mutSS)prM80%E. This procedure replaced the native E signal sequence (Pro-Ser-Met-Thr₋₁-25 Met₊₁) (SEQ ID NO:32) with the optimized E signal sequence (Gly-Ala-Gln-Ala₋₁-Gln₊₁) (SEQ ID NO:33).

Plasmid pLS6prM(mutSS)100E-TGA was obtained by homologous replacement of a SmaI-SacI fragment between plasmids pAlterSmaH3prME(mutSS) (see Example 14) and pLS6prM100E. DNA sequencing of pLS6prM(mutSS)100E-TGA identified an unintended TGA stop codon within E downstream of the mutated secretion signal. To transfer the altered secretion signal encoding sequence to pLS6prM80%E and to separate the cDNA fragment containing the altered secretion signal of E from the TGA stop

codon, a BglII-EcoNI fragment from pLS6prM(mutSS)100E-

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TGA, encompassing prM and the first 430 nucleotides of E and lacking the TGA stop codon, was transferred to plasmid pLS6prM80%E which had been similarly digested to yield the expression plasmid pLS6prM(mutSS)80%E. The sequence of the expression plasmid was confirmed by restriction digestion and DNA sequence analysis. The nucleotide and predicted amino acid sequences of the MFa_{L} -prM junction are identical to the sequences listed above.

As described for the nonmutagenized plasmid, the DNA was transformed into the S. cerevisiae GL43 strain and transformants were selected based upon their ability to grow on unsupplemented minimal medium (see Example 11). Transformants were cultured, induced, and

- evaluated as described above for the nonmutated $MF\alpha_{L}$ prM80%E transformants. Proteins secreted into the culture medium as well as total cellular proteins were treated with EndoH_f prior to separation by SDS-PAGE. Protein gels were analyzed by Coomassie staining and
- 20 immunoprobing of Western blots. SDS-PAGE analysis of concentrated culture medium failed to identify a novel Coomassie staining band. Immunoprobing with anti-DEN2 HMAF and anti-DomB antiserum, however, revealed a small amount of processed immunoreactive E protein in the
- 25 medium. The size of the immunoreactive protein (approximately 50 kD) was similar to the secreted protein from pLS6-80%E expression vector. Evaluation of intracellular expression of the fusion protein containing the optimized secretion signal by SDS-PAGE and Western
- 30 blot demonstrated that the transformed cells produce immunoreactive product recognized by anti-DEN2 HMAF and anti-DomB antiserum. Unlike the immunoreactive product seen in pLS6prM80%E transformants, the immunoreactive band found in pLS6 (mutSS) prM80%E transformants was not recognized by $MF\alpha_L$ anti-serum suggesting that processing 35 had occurred at the prM-E junction. Thus, the mutagenesis of the signalase cleavage site resulted in

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greatly enhanced processing of the $MF\alpha_L$ -prM80%E product at the prM-E junction.

Example 16

5 <u>Construction of expression vector pPIC9-prM(mutss)80%E</u> and secretion of 80%E by *P. pastoris* expressing MFα_LprM(mutss)80%E

The expression vector constructed to express preMembrane-mutated secretion signal-80%Envelope 10 (pPICprM(mutSS)80%E) in P. pastoris from a single continuous open reading frame utilized the DEN-2 PR159/S1 prM and E gene sequences described above (Example 14). The plasmid, pPIC9-prM(mutSS)80%E, was constructed by transferring a prM(mutSS)80%E fragment from the S. cerevisiae expression plasmid pLS6prM(mutSS)80%E into 15 pPIC9. The P. pastoris expression vector pPIC9 (Example 4) and the S. cerevisiae expression vector pLS6 (Example 2) both use the MFa prepropeptide leader (MFa_L) sequence to direct the secretion of expressed proteins, and the 20 two MFa₁ sequences share an XhoI restriction endonuclease site, encoding amino acids leucine and glutamic acid, just upstream of the Kex2 protease site. The transfer of the dengue cDNA fragment made use of this XhoI site.

Prior to transferring the prM(mutSS)80%E cDNA 25 fragment, sequences encoding extraneous amino acids and an extraneous XhoI site at the MF α_L -prM fusion were first removed. This was accomplished by digesting pLS6prM(mutSS)80%E with restriction endonuclease XhoI and XmaI into which was ligated a synthetic oligonucleotide

30 duplex {5'-TCGAGAAGAGAGAGAG-3'(SEQ ID NO:36) and 5'-CCGGCTTCTCTCTCTC-3'(SEQ ID NO:35)}. This manipulation deleted the extraneous XhoI site, the XmaI site, and six extraneous codons and preserved the MFa_L proteolytic processing sites and the XhoI site within MFa_L required

35 for the cDNA fragment transfer from pLS6 to pPIC9. The nucleotide and predicted amino acid

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sequence at the MF α_L -prM fusion junction from pLS6prM(mutSS)80%E and pLS6 Δ -prM(mutSS)80%E are:

MFα_L-prM junction of pLS6-prM(mutSS)80%E 5 (SEO ID NO:36 and SEO ID NO:37)

<u>XhoI</u> <u>XmaI</u> ATG ...CTC GAG AAA AGG GAG GCC TTTAGATCTCGAGTACCCGGGACCATG TTT .. Met ...Leu Glu Lys Arg Glu Ala PheArgSerArgValProGlyThrMet Phei..

Kex2p DPAP

MF α_L -prM junction of pLS6 Δ -prM(mutSS)80%E (SEQ ID NO:38 and SEQ ID NO:39)

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ATGCTC GAG AAA AGG GAG GCC GGGACCATG TTT .. MetLeu Glu Lys Arg Glu Ala GlyThrMet Phe1..

>) 🔺 Kex2p DPAP

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To construct the clone pPIC9-prM(mutSS)80%E, a XhoI-SalI fragment encoding prM(mutSS)80%E sequences was obtained from pLS6 Δ -prM(mutSS)80%E and was inserted into the pPIC9 vector that had been digested with XhoI. The genetic integrity of the expression plasmid, pPIC9prM(mutSS)80%E, was confirmed by restriction digestion.

The pPIC9-prM(mutSS)80%E expression vector was transformed into spheroplasts of *P. pastoris* strain GS115 30 (*his4*), and His⁺ transformants were selected for their ability to grow on minimal medium without histidine supplementation. The transformants were screened for expression and secretion of 80%E as described in Example 12. No unique Coomassie staining bands were detected in

- 35 the culture medium of either nonEndoH_f or EndoH_f treated samples (similar to that observed for culture medium from pPIC9-80%E transformants - see Example 12). Western immunoblots of proteins from the culture medium probed with anti-domain B serum detected multiple bands with a
- 40 diffuse area of reactivity ranging from 50 to 90 kD in the nonEndoH treated samples and a unique band of approximately 50 kD in the EndoH treated samples. No immunoreactive polypeptides were detected by the anti-



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domain B serum in negative control samples. Various amounts of the corresponding immunoreactive protein were detected in EndoH treated samples of cellular proteins. We estimate that the amount of secreted 80%E in the culture medium is less than 1% of the total amount of secreted protein based on the observation that no

- polypeptide of the appropriate molecular weight was detected by Coomassie staining of EndoH_f treated samples.
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<u>Example 17</u>

Construction of pMttbns-prM80%E and Secretion of 80%E by Drosophila melanogaster Schneider cells Expressing tPALprM80%E

- For expression of DEN-2 PR159/S1 preMembrane 15 protein amino acids 1-166 and Envelope glycoprotein amino acids 1-395 as a single continuous open reading frame in Drosophila melanogaster Schneider 2 tissue culture cells, DNA sequences encoding these proteins were obtained by digestion of the p48BSprM80%E clone (described in Example
- 20 14) with the restriction enzymes BgIII and SaII. This fragment was cloned into the unique BgIII and XhoIrestriction sites of plasmid pMtt ΔXho (described in Example 13). The cloning junctions of the resulting gene fusion expression vector, pMttprM80E, were confirmed by
- 25 restriction endonuclease digestion and DNA sequence analysis. The partial nucleotide and predicted amino acid sequences of the tPA_L-prM80E fusion gene are (SEQ ID NO:40 and SEQ ID NO:41):
- 35 The tPA pre- and propeptide regions are delineated by pre^A and pro-tPA^A, respectively, and the dengue sequences are indicated in bold type. The Phe₁ and Thr₁₆₆ residues are the N-terminal and C-terminal amino acid residues of prM, respectively. The Met₁ residue is the N-terminal

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amino acid of envelope glycoprotein and Gly₃₉₅ is residue 395 from the amino terminal end of the envelope glycoprotein.

As described previously in Example 13, 5 Schneider 2 cells were cotransfected with pMttprM80%E DNA at ratios of 1:1, 5:1, and 20:1 relative to pCOHygro DNA. Transformants were induced with 200µM CuSO₄ and expression of prM80%E was examined at various times after induction. Proteins secreted into the culture medium as well as 10 cellular proteins were separated by SDS-PAGE. Protein gels were analyzed by both Coomassie Blue staining and

immunoprobing of corresponding Western blots. Analysis of Coomassie Blue-stained SDS-PAGE gels identified a novel band of approximately 50 kilodalton in all

15 transfectants. This novel band was also recognized by anti-DEN-2 HMAF in Western blot analysis. This ~50 kD immunoreactive band is roughly the same size as the secreted EndoH-treated product from pLS6-80%E transformed yeast cells (Example 11) and slightly smaller than the

secreted 80%E from pMttbns80%E-transformed D. melanogaster Schneider cells (Example 13), suggesting the Envelope protein had been processed away from the preMembrane protein. (The size discrepancy between 80%E secreted by pMtt80%E and pMttprM80%E Schneider cells is

25 discussed in Example 13.) Polyclonal antisera to the pr portion of prM (from Peter Wright, Monash University, Australia) did not recognize the ~50 kD protein, confirming that the 80%E produced in the transfected cells was processed from prM. In fact, no evidence of a

30 higher molecular weight band that might correspond to unprocessed prM80%E was detected in any sample, suggesting that the proteolytic processing of prM from E is extremely efficient in Schneider cells. The fate of the prM portion of the fusion remains unresolved as no 35 distinct immunoreactive band was detected by probing with the anti-pr antisera.

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The secreted 80%E glycoprotein was partially purified (judged by the presence of a single major band on a sliver stained SDS-PAGE gel) and its N-terminal amino acid sequence was determined. To purify the secreted glycoprotein, culture medium was concentrated and buffer exchanged against 20 mM succinate pH 5.7. The buffer exchanged material was loaded onto a CM-BioGel column and eluted in 150 mM NaCl. The 150 mM NaCl eluant was separated on an SDS-PAGE gel and electro-transferred to Immobilon-P membrane (Millipore). The 80%E band was excised, and the N-terminal amino acids were determined by Edman sequencing. Two amino acid sequences were obtained. One, Met-Xxx-Xxx-Ile-Gly-Ile (SEQ ID NO:42), had an individual residue yield of 7.9 -10.0 picomoles, while the other, Val-Xxx-Val-Gly-Ala-Val (SEQ ID NO:43), had a 3.2- 4.2 picomole yield. Incomplete reduction of the Cys at position three may account for lack of its detection. the first sequence is consistent with the

expected sequence, Met-Arg-Cys-Ile-Gly-Ile (SEQ ID 20 NO:44), supporting the interpretation that the ~50 kD secreted immunoreactive glycoprotein is correctly processed 80%E of DEN-2.

Sensitivity of the secreted 80%E to endoglycosidases was evaluated by molecular weight shift of the protein in SDS-PAGE and Western immunoblots following endoglycosidase treatment. Resistance of the secreted 80%E to Endoglycosidase H_f (Endo H_f; New England Biolabs) and sensitivity to N-glycosidase F (PNGase F; New England Biolabs) digestion indicated that the secreted product contains N-linked glycosylation, and that the glycosylation is probably complex and is neither

The secreted protein is one of the predominant proteins in the unconcentrated medium, comprising as much 35 as 20% of the total secreted protein. Estimates of the concentration of the 80%E product in unconcentrated medium based upon sandwich ELISA assays (described in detail in Example 7) and Coomassie blue staining range

high mannose nor hybrid in composition.



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from 3-16 µg/ml depending on the preparation. Immunoblots probed with polyclonal anti-dengue 2 hyperimmune mouse ascites fluid (DEN-2 HMAF; from R. Putnak, WRAIR) demonstrated that the amount of secreted 80%E produced by the transfectants increased over time from day 1 post induction to 7 days post induction. The amount of 80%E detected intracellularly in the transfectants correlated with the cotransfection ratio, but the increase in intracellular 80%E with time was not as dramatic as for secreted 80%E, suggesting efficient secretion of 80%E and accumulation in the medium.

Example 18

Induction of anti-Dengue 2 antibodies in mice by Pichia pastoris-secreted 80%E

P. pastoris cells transformed with pPIC-80%E (described in Example 12) were induced with 0.5% methanol and the medium was collected after 40 hours of induction (for additional details on culture conditions see Example 12). The medium was filtered through a 0.5 μ m low protein binding filter (Opticap, Millipore), then buffer exchanged with phosphate buffered saline (10 mM sodium phosphate, 2 mM potassium phosphate, 0.15 M sodium chloride, and 27 mM potassium chloride, pH 7.5) and concentrated approximately 40 fold using a combination of tangential flow (Minitan; Millipore) and centrifugal ultrafiltration (Centriprep 30; Amicon). The concentrated medium was analyzed by Western blot and assayed by sandwich ELISA (described in detail in Example 7) prior to injection into mice. Five week-old Swiss Webster outbred mice (Simonsen) were immunized by intraperitoneal (I.P.) injection with 100 µg total protein of the crude concentrated 80%E medium with or

without complete Freund's adjuvant. Controls for this experiment included a negative control medium prepared from a nonrecombinant *P. pastoris* culture as described above for the 80%E medium. Protein precipitation was

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observed during the concentration of the negative control medium, consequently the final protein concentration of the concentrated medium was lower than that from the 80%E medium. (For this reason, 12.5 μ g of total protein in Freund's complete adjuvant was used for immunization with

- 5 Freund's complete adjuvant was used for immunization with the negative control medium.) Additional controls included saline and KLH-domain B, a recombinant dengue product previously shown to induce neutralizing antibodies (Example 8), both were administered with
- 10 Freund's complete adjuvant. The mice received three I.P. boosts consisting of one half the priming dose with or without Freund's incomplete adjuvant, and consistent with the specific priming immunization. The boosts were administered at two week intervals.
- 15 Following the second boost, the animals were test bled (tail vein) and the immune response was monitored using an indirect ELISA. In the ELISA assay, plates were coated with a bovine serum albumin (BSA) domain B conjugate, blocked with BSA, and serial
- 20 dilutions of the mouse sera were then incubated with the coating antigen. Alkaline phosphatase-labeled goat anti-mouse IgG was used as the secondary detecting antibody, and the color development upon addition of an alkaline phosphatase chromogenic substrate was monitored.
- 25 The ELISA titer is that which corresponded to the highest dilution of serum resulting in an optical density twofold above background (reactivity of the serum against BSA only). Following the third boost, the animals were bled-out and the serum was tested for anti-DEN2
- 30 responsiveness by indirect ELISA and the plaque reduction neutralization assay (PRNT). In the PRNT assay, the mouse sera were serially diluted in Eagles minimal essential medium (EMEM; BioWhittaker) supplemented with 10% fetal bovine serum (FBS; Hyclone) and mixed with 100 35 plaque forming units (pfu) of DEN2 virus (New Guinea C strain, from Robert Putnak, WRAIR). After allowing one hour for binding and neutralization of the virus, the

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serum-virus mixtures were plated onto susceptible monkey kidney monolayers (Vero cells, from Robert Putnak, WRAIR) plated in EMEM (BioWhittaker) supplemented with 10% FBS (Hyclone) in 6 well tissue culture dishes (Corning). The cells were overlaid with 0.9% agarose (Seakem; FMC) in 5 EMEM supplemented with 5% FBS and viral cytopathic effect was allowed to develop for 6-7 days. The resulting viral plaques were stained with 0.012% neutral red (Sigma) in 1% agarose (Seakem; FMC). The number of plaques in each cluster were counted and compared to a no-serum viral 10 control. The $PRNT_{80}$ was the highest dilution of serum that resulted in an 80% reduction in the number of plaques from a given viral inoculum. Results from the ELISA and PRNT assays are summarized in Table 4. The P. pastoris expressed 80%E induces a potent anti-DEN2 response in mice, with ELISA titers of up to 1:102,400. The titers obtained in the presence of adjuvant exceeded those obtained without adjuvant. The lack of a strong virus neutralizing response ($PRNT_{80}$ titers, Table 4) may simply reflect the crude nature of the immunogen, as

20 similar results were obtained when crude domain B was used as an immunogen (see Example 8).

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Table 4 Induction of Anti-DEN2 Immune Response in Mice Immunized with <i>P. pastoris</i> - expressed 80%E							
mouse	antigen	adjuvant	30 titer ELISA	40 titer ELISA	40 titer PRNT ₈₀		
30-1	Saline	Freund's	< 1:50	< 1:100			
30-2			< 1:50	< 1:100			
30-3			< 1:50	< 1:100			
30-4			< 1:50	< 1:100			
30-5			< 1:50	< 1:100			
32-1	KLH- DomB "26"Tg	Freund's	>1:6400	1:25600			
32-2			> 1:1600	1:102,400			
32-3			<1:100	1:100			
32-4			>1:6400	1:102,400			
32-5			<1:100	< 1:100			
34-1	12.5 Tg Pichia negative control medium	Freund's	< 1:100	< 1:100			
34-2			<1:100	1:100			
34-3			< 1:100	< 1:100			
34-4			< 1:100	< 1:100			
34-5			< 1:100	1:100			
36-1	100 Tg Pichia 80%E total medium	Freund's	>1:6400	1:25,600	<1:10		
36-2			1:6400	1:25,600	< 1:10		
36-3			>1:6400	1:25,600	<1:10		
36-4			1:100	1:100	<1:10		
36-5			1:6400	1:102,400	<1:10		
37-1	100 Tg Pichia 80%E total medium	None	< 1:100	1:100			
37-2	1		1:1600	1:6400			
37-3	1.		1:100	1:400			
37-4	1		1:400	1:6400			
37-5			1:100	<1:100			
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Example 19

Induction of dengue virus-neutralizing antibodies by immunizing mice with 80%E secreted by Drosophila melanogaster Schneider cells expressing tPAL-prM80%E and <u>tPA_I-80%E</u>

Schneider cells, transformed with pMtt-prM80%E and pMtt-80%E expressing the tissue plasminogen activator leader fusion proteins tPAL-prM80%E and tPAL-80%E,

- 10 respectively (Described in detail in Examples 17 and 13, respectively), were cultured in serum-free medium (Excell; JRH Biosciences) and induced by addition of CuSO₄ to a final concentration in the culture medium of 0.2 mM (see examples 13 and 17 for more detail on culture conditions). The cells were maintained in inducing 15 medium for seven days prior to harvesting. The cells were removed by centrifugation at 1000 X G in a Beckman TJ-6 refrigerated centrifuge and the media were filtered through a 0.2 µm cellulose acetate filter (Nalgene). The
- 20 media containing the recombinant 80%E were concentrated 20-fold using centrifugal concentrators (Centriprep 30; Amicon) and assayed by ELISA (described in detail in Example 7) and Western immunoblots prior to their use as an immunogen in mice. Negative control medium, derived
- from Schneider cells transformed with pCOHygro only (See 25 Example 13), was produced as described above. In two series of immunizations, outbred Swiss Webster mice (Simonsen) were immunized intraperitoneally (I.P.) with 100 µg total protein in Freund's complete adjuvant.
- 30 Control animals were immunized with 100 μ g of total protein from the concentrated negative control medium, 80 µg of purified Saccharomyces-expressed domain B (see Example 8), or saline only, each in Freund's complete adjuvant. In the first series of immunizations, the mice received three I.P. boosts, consisting of one half 35 the priming dose in Freund's incomplete adjuvant, at two

week intervals. In the second series of I.P. immunizations, the mice received two boosts, each at one month intervals.

Following the first and second boosts, the animals were test bled (tail bleed) and the immune response was monitored using an indirect ELISA as described in example 18. Following the final boost, the animals were bled out and the sera were tested for anti-DEN2 responsiveness by indirect ELISA and PRNT as

- 10 described in Example 18. Results for the ELISA and PRNT assays are summarized in Tables 5 and 6. In both series of immunizations, the mice immunized with the crude media containing 80%E, expressed cotranslationally with prM or independently without prM, developed high titer, virus-15 neutralizing antibodies. These titers are higher than any previously reported titers for any immunogen produced
 - from any flavivirus, suggesting the utility of these immunogens as efficacious vaccine candidates.

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Table 5							
Immune Response of Mice Immunized with Crude <i>Drosophila</i> Media Containing Dengue 2 Virus 80%E Expressed as a prM80%E Fusion							
mouse	antigen	adjuvant	20 titer ^a ELISA	30 titer⁵ ELISA	Final titer ^e ELISA	PRNTso	
						titer ^c	
20-1	Saline	Freund's	< 1:50	< 1:50	< 1:100	<1:10	
20-2			< 1:50	< 1:50	<1:100	<1:10	
20-3			< 1:50	< 1:50	< 1:100	<1:10	
20-4			< 1:50	< 1:50	<1:100	<1:10	
20-5			< 1:50	<1:50	<1:100	<1:10	
21-1	pCoHygro	Freund's	< 1:50	< 1:50	< 1:100	<1:10	
21-2			< 1:50	< 1:50	<1:100	<1:10	
21-3			< 1:50	< 1:50	< 1:100	<1:10	
21-4			< 1:50	< 1:50	< 1:100	<1:10	
21-5			< 1:50	< 1:50	< 1:100	<1:10	
22-1	prM80%E	Freund's	1:3200	>1:25,600	1:102,400	1:2560	
22-2			>1:800	> 1:6,400	1:25,600	1:2560	
22-3			>1:200	>1:1,600	1:25,600	1:2560	
22-4			< 1:50	>1:102,400	>1:102,400	1:2560	
22-5			1:3200	>1:25,600	>1:25,600	1:640	

^aDetermined following the 2nd injection. ^bDetermined following the 3rd injection. ^CDetermined following the 4th and final injection.

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Table 6						
Immune Response of Mice Immunized with Crude <i>Drosophila</i> Media Containing Dengue 2 Virus 80%E Expressed with or without prM						
mouse	antigen	adjuvant	20 titer*	3/ titer ^b	Final titer ^e	
			ELISA	ELISA	PRNT ₈₀	
25-1	Saline	Freund's	< 1:50	>1:50	< 1:10	
25-2			< 1:50	>1:50	< 1:10	
25-3			< 1:50	>1:50	< 1:10	
25-4			< 1:50	>1:50	<1:10	
25-5			< 1:50	>1:50	< 1:10	
27-1 [¢]	80Tg	Freund's	1:1600	1:102,400	< 1:100	
	Purified					
	Sacchro					
	DomB					
27-2			1:100	1:6400	1:10	
27-3			>1:6400	1:102,400	< 1:100	
27-4			>1:6400	1:409,600	1:40	
27-5			>1:6400	1:409,600	< 1:500	
28-1	100Tg	Freund's	DEAD	NT	· NT	
	Drosophila					
	prM80%E					
	total medium					
28-2			>1:6400	1:102,400	1:8000	
28-3			>1:6400	1:409,600	1:8000	
28-4			>1:6400	1:102,400	1:4000	
28-5			>1:6400	1:102,400	1:1000	
29-1	100Tg	Freund's	1:6400	1:6400	1:500	
	Drosophila					
	80%E					
	total medium					
29-2			1:6400	1:102,400	1:4000	
29-3			<1:100	1:25,600	1:1000	
29-4			>1:6400	1:25,600	1:8000	
29-5			>1:6400	1:102,400	1:4000	

^aDetermined following the 2nd injection. ^bDetermined following the 3rd injection. ^CDetermined following the 4th and final injection. ^dmouse 27-1 did not receive second boost.

NT - not tested.

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Example 20

Protection from dengue virus challenge by immunizing mice with 80%E secreted by Drosophila melanogaster Schneider cells expressing tPA_L-prM80%E or tPA_L-80%E.

Schneider cells, transformed with pMtt-prM80%E and pMtt-80%E expressing the tissue plasminogen activator leader fusion proteins tPAL-prM80%E and tPAL-80%E respectively (described in detail in Examples 17 and 13, respectively), were cultured in serum-free medium 10 (Excell; JRH Biosciences) and induced by addition of $CuSO_4$ to the culture medium at final concentration of 0.2 mM (see examples 13 and 17 for additional details on culture conditions). The cells were maintained in inducing medium for seven days prior to collecting the

15 medium. The cells were removed by centrifugation at 1000 X G in a Beckman TJ-6 refrigerated centrifuge and the media were filtered through a 0.2 µm cellulose acetate filter (Nalgene). The media were concentrated 20-fold using centrifugal ultrafiltration (Centriprep 30, Amicon)

20 and assayed by ELISA and Western immunoblots prior to use as an immunogen in mice. Negative control medium, derived from Schneider cells transformed with pCOHygro only (see Example 13), was produced as described above. These immunogens were used to immunize 10-13 day old

25 Balb/c mice (Jackson Labs). A total of 0.1 ml of each preparation (corresponding to 70 μ g total protein from negative control medium, 230 µg total protein from tPAL-80%E medium, and 150 µg total protein from tPAL-prM80%E medium) were used to subcutaneously inoculate groups of

30 10 mice each, using Alum as adjuvant. An identical second dose was administered 14 days later as a booster. One week after the second dose the mice were challenged with an intracranial injection of 100 LD₅₀ units of DEN2 New Guinea C strain (10⁵ pfu/injection).

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The morbidity and mortality of the mice were monitored for 17 days post challenge. Six days after "WO 96/37221

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challenge the mice immunized with the negative control medium began exhibiting symptoms of infection including ruffled fur, hindlimb paralysis, wasting, and death. Eight of the 10 negative control mice were dead by day 12 post-challenge and the remaining two mice recovered by

- day 15 post-challenge. In contrast, of the 10 mice immunized with the tPA_{T} -80%E, only four exhibited any symptoms of infection, and eight of 10 survived the challenge. Similarly, nine of 10 mice immunized with
- 10 tPA_L-prM80%E survived the challenge, although seven of these mice exhibited mild symptoms of infection during the monitoring period. These survival data are illustrated in Figure 13, and show that both Drosophilaexpressed 80%E antigens efficiently protected mice from viral challenge. These results emphasize the utility of 15 the Drosophila cell expressed 80%E dengue immunogens as vaccine candidates.

Example 21

Construction of DEN-2 N-terminal 60%E and prM60%E cDNA Fragments

A subclone encoding the N-terminal 60% of E was constructed using p29GEB24PS and p29D280E. Example 1 of the parent application --- describes construction of

- 25 p29GEB24PS. Plasmid p29GEB24PS holds a BamHI fragment insert containing, in part, DEN-2 E sequences (nucleotides 1696-2121) starting at a DEN-2 genomic BamHI site and ending with the Gly395 codon, followed immediately by two stop codons and a Sall site.
- 30 Oligonucleotide mutagenesis had deleted codons for Leu292 through Lys295 (nucleotides 1810-1821) and had inserted two stop codons and a SalI site immediately following the Lys291 codon (see Figure 4 and Example 1).

The N-terminal 80%E insert in p29D280E was then converted to a 60%E insert by replacing a restriction fragment encoding the 3' end of 80%E with a restriction fragment from p29GEB24PS encoding the 3' end of 60%E. То

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accomplish this, DNA of p29GEB24PS was digested with BamHI, the ~590 bp BamHI fragment was isolated by agarose gel electrophoresis and then digested with Sall, and finally the 119 bp BamHI-SalI fragment released from the ~590 bp BamHI fragment and containing dengue nucleotides 1696-1809 was isolated by agarose gel electrophoresis and ligated into p29D280E prepared as follows. Plasmid p29D280E was digested with BamHI, which cuts the BamHI site (dengue nucleotides 1696-1701) within 80%E, and with Sall, which cuts immediately 3' of 80%E and also within the vector, pBR322, 422 base pairs distal to the 3' end of the 80%E fragment. Following ligation, the desired product, a plasmid containing the cDNA encoding the N-terminal 60% of E in pBR322 (p29D260E), was recovered by transformation of E. coli with the ligation mixture and screening transformant colonies for plasmids of the appropriate size and restriction digestion pattern. Proper ligation of the BamHI-Sall fragment in p29D260E was confirmed by DNA sequence determination.

To construct a cDNA encoding prM and the amino terminal 60% of E (prM60%E), we used a strategy identical to that used to construct prM80%E (Example 14). The prM100%E plasmid, p29prME13, was digested with BamHI and SalI to release the 794 bp 3' end fragment of E, which was then replaced with the 119 bp BamHI-SalI fragment encoding a 40% carboxy-end truncation of E from p29D260E. The resulting truncated cDNA clone, p48BSprM60E, encodes a prM-60%E fusion ending with Lys291 of E and was

confirmed by restriction digestion and DNA sequence 30 analysis.

Example 22

<u>Construction of expression vector pLS6-60%E and secretion</u> <u>of 60%E by Saccharomyces cerevisiae expressing MFa-60%E</u> An expression vector (pLS6-60%E) was constructed for secretion of the N-terminal 60% (codons

1-291, 60%E) of the DEN-2 PR-159 S1 envelope glycoprotein

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from S. cerevisiae. The 60%E DNA sequences were obtained from plasmid p29D260E, described in example 21, by restriction endonuclease digestion with both BglII and SalI. The released fragment was isolated by agarose gel electrophoresis and subcloned into the BglII and SalI sites of pLS6, a yeast expression vector described in example 2. The MF α_L -60%E fusion was made such that processing of the MF α propeptide and trimming of a Glu-Ala dipeptide results in 60%E with eight additional Nterminal amino acids encoded by sequences present in the multiple cloning site of the pLS6 vector and the E gene PCR primer adapter (see below) (SEQ ID NO:45 and SEQ ID

NO:46). 15 ATGGCT......GAG GCC TTT AGA TCT CGA GTA CCC GGG ACC ATG ...AAA TAA TAG Not 1900 Ale Clea Cle Ale Che Are See Are Val Dee Cle The Meter Lyaw End Fed

Met-18aa Ala-65aa Glu Ala Phe Arg Ser Arg Val Pro Gly Thr Met...LyszaiEnd End 60%E Signalase Kex2p DPAP

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Transcription of the gene fusion is driven by the S. cerevisiae copper-inducible copper metallothionein (CUP1) promoter.

After confirming the DNA sequence of the 25 ligated junctions of expression vector pLS6-60%E, the recombinant DNA was transformed into *S. cerevisiae* strain GL43 (*MATa trp1A1 ura3-52 pep4::URA3*; SmithKline Beecham) according to standard protocols (Gutherie & Fink, eds. 1991; Rose et al, 1990). Transformants were selected by 30 their ability to grow on minimal medium (SD medium:

Guthrie and Fink, 1991) without tryptophan supplementation.

In order to test for expression and secretion of 60%E, several transformants were grown in small-scale 35 cultures (5 ml medium in 17x150 mm tubes). Single colonies were used to inoculate SD medium and the inoculated cultures were grown to saturation (overnight, 30°C, 220 rpm). 1x10⁸ cells from the overnight culture were used to inoculate 5 ml of minimal SD medium

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supplemented with Casamino acids (2 g/l; Difco) and CuSO₄ (200 μ M). This expression culture was fed with glucose (4 g/l, final concentration) and sodium phosphate (pH 6.7, 20 mM, final concentration) at 24 hours post inoculation and, after 48 hours of growth (30°C, 220 rpm, in 17x150 mm tubes), was harvested by centrifugation. Cell-free spent medium was buffer-exchanged with TEEN (50 mM Tris, 10 mM EDTA, 10 mM EGTA, 150 mM NaCl, pH 8.0) and concentrated 200-fold using Centricon-30 (Amicon) ultrafiltration. An extract of cellular protein was prepared by lysing the yeast cells with vigorous agitation in the presence of glass beads (425-600 μ m) and TEEN+PIC (TEEN with 1 μ g/ml each of pepstatin and

leupeptin and 1 mM phenylmethylsulfonylfluoride) using a Mini Beadbeater apparatus (BioSpec Products, Bartlesville, OK). Samples were endoglycosidase H_f digested according to the manufacturer's (New England Biolabs, Beverly, MA) protocol prior to SDS-PAGE analysis. Protein gels were Coomassie-stained directly as well as Western blotted and immunoprobed using anti-DEN2 HMAF.

Negative control yeast carrying the expression vector without a Dengue gene insert secreted no proteins recognized by anti-DEN2 HMAF, while pLS6-60%E medium 25 contained several immunoreactive species. The major band presumably represents full-length 60%E since its apparent molecular weight is approximately 10 kD less than that of recombinant 80%E. A protein band comigrating with this immunoreactive material was visible in Coomassie-stained

30 gels of proteins secreted by pLS6-60%E transformants, but this band was absent in medium of negative control transformants. The immunoblot of proteins secreted by pLS6-60%E transformants evidenced a minor band of apparent molecular weight 6-8 kD larger than 60%E; this likely represents unprocessed MFa propeptide-60%E. The

cellular protein extract of pLS6-60%E transformants

contained many immunoreactive polypeptides not observed in negative control cells; two of these match the secreted products discussed above.

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Example 23

<u>Construction of expression vectors pLS6-prM60%E and</u> <u>pLS6-prM(mutSS)60%E, expression of MFa_L-prM60%E and</u> <u>MFa_L-prM(mutSS)60%E in Saccharomyces cerevisiae, and</u> <u>secretion of 60%E by Saccharomyces cerevisiae expressing</u>

MFa_L-prM(mutSS)60%E

For expression of DEN-2 PR159/S1 premembrane protein amino acids 1-166 and envelope glycoprotein amino acids 1-291 as a single contiguous open reading frame in S. cerevisiae, DNA sequences encoding these proteins were 15 obtained by digestion of plasmid p48BSprM60E (see Example 21) with the restriction enzymes BqlII and Sall. This fragment was cloned into the pLS6 S. cerevisiae expression vector (see Example 2) that had been digested with BglII and SalI. The structure of the resulting gene fusion expression vector, pLS6-prM60%E, was confirmed by 20 restriction digestion and DNA sequence analysis. The Nterminal MFa_L-prM fusion amino acid sequence of the MFa_LprM60%E fusion protein are identical to those described for MF α_{L} -prM(mutSS)80%E fusions (Example 15), while the

25 C-terminal amino acid of the fusion protein is Lys₂₉₁ of the dengue envelope glycoprotein.

The pLS6-prM60%E plasmid was transformed into S. cerevisiae strain GL43 (MATa ura3-52 trp1 Δ 1 pep4::URA3) and screened for 60%E expression as described in Example 11. Proteins secreted into the culture medium as well as cellular proteins were treated with endoglycosidase H_f (EndoH, New England Biolabs, Beverly, MA) and separated by SDS-PAGE. Proteins were analyzed by both Coomassie staining of polyacrylamide gels and immunoprobing of Western blots. Similar to the expression of secreted 80%E from pLS6-prM80%E (Example

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15), secreted 60%E was not detected on Coomassie stained polyacrylamide gels nor Western blots. An immunoblot of intracellular protein confirmed that the construct (MF α_L prM60%E) was expressed, but the fusion product had not been processed to prM and E (evaluation was performed as described in Example 15).

Because the dengue E signal sequence itself may limit processing of the prME fusion proteins, expression of MF α_{L} -prM(mutSS)60%E was evaluated. The BallI-EcoNI fragment from pLS6-prM(mutSS)100%E-TGA encoding the altered secretion signal peptidase cleavage site (see Example 15) was used to replace the homologous fragment from the pLS6-prM60%E to produce pLS6-prM(mutSS)60%E.

The sequence of the expression plasmid was confirmed by 15 restriction digestion and DNA sequence analysis. Plasmid pLS6prM(mutSS)60%E was transformed into the S. cerevisiae GL43 strain and transformants were selected as described in Example 11.

Transformants were cultured, induced, and 20 evaluated as described in Examples 11 and 15. In contrast to the expression of MFa_{I} -prM60%E, Western blot analysis of total intracellular proteins from pLS6prM(mutSS)60%E transformants demonstrated that the transformed cells produce an approximately 40 kilodalton

25 product recognized by anti-DEN2 HMAF. For analysis of secreted proteins, media from induced cultures were concentrated, treated with endoglycosidase H_f , and analyzed on Western blots for E antigen. A small amount of processed 60%E could be identified in the culture

30 medium upon immunoprobing with anti-DEN2 HMAF. Thus, the mutagenesis of the signalase cleavage site resulted in greatly enhanced processing of the MFa_L -prM(mutSS)60%E product at the prM-E junction which produced secretion of a processed 60%E from the $MF\alpha_L$ -prM(mutSS)60%E in 35 S. cerevisiae.

Example 24

Construction of expression vector pPIC9-60%E and secretion of 60%E by *P. pastoris* expressing MFa_L-60%E The expression vector constructed to secrete 5 60%E from *P. pastoris*, pPIC9-60%E, included the DEN-2 PR-159 S1 envelope glycoprotein amino acids 1-291. As a precursor to this 60%E expression vector, a modified pLS6-60%E plasmid (pLS6-alt60%E), encoding a fusion between MFa_L and the amino terminal 60% of the dengue 10 envelope, was constructed that encodes fewer nondengue amino acids between the MFa_L and E segments. The 60%E cDNA fragment from pLS6-alt60%E was then transferred to pPIC9.

The sequences in pLS6-alt60%E encoding the 15 dengue E protein amino terminus were derived from pLS6-2x80E, which encodes a tandemly arrayed dimer of 80%E that are linked by a synthetic linker peptide. To convert pLS6-2x80E to pLS6-alt60%E, pLS6-2x80E was digested with *Eco*NI, which cuts within the first member

20 of the 80%E dimer, and SalI, which cuts immediately downstream of the dimer, thereby removing the 3' portion of the first member of the 80E dimer and the second member of the 80E dimer entirely. In its place was ligated an EcoNI-SalI fragment from pLS6-prM(mutSS)60E

25 (see Example 23), encoding the 3' portion of 60%E, to complete construction of pLS6-alt60%E.

The minimizing of nondengue codons between $MF\alpha_L$ and E, found in pLS6-2x80E, were preserved in pLS6alt60%E, and, subsequently, in pPIC9-60%E. The codons

30 between MFα_L and E were minimized, owing to use of the XmaI site in p29D280E (XmaI is the first adapter restriction enzyme site upstream of E; see Example 1) and the StuI site in pLS6. To ligate the XmaI 5' end of E to the StuI site in the MFα_L, the XmaI site was treated with 35 Klenow polymerase to make the end blunt during the construction of pLS6-2x80E.

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To transfer 60%E from pLS6-alt60%E to pPIC9, pLS6-alt60E was digested by XhoI plus SalI, which released a fragment that includes a portion of the MF α leader and the entire 60%E coding region. This fragment was ligated with the Pichia expression vector pPIC9 (Invitrogen, San Diego, CA; described in Example 12) that was previously digested with the restriction enzyme XhoI. This ligation restored the complete MF α leader sequence including the Kex2 cleavage site. The genetic integrity of the resulting gene fusion expression vector, pPIC9-60%E, was confirmed by restriction digestion and DNA sequence analysis. The partial nucleotide and predicted amino acid sequences of the MF α_L -60%E fusion gene are shown below (SEQ ID NO:47 and SEQ ID NO:48):

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The location of the signalase and Kex2 cleavage sites which remove the pre and pro portions of the MFα leader peptide, respectively, are indicated. The dengue 25 sequences are indicated in bold type. The Met₁ residue is the N-terminal amino acid of the E glycoprotein and Lys₂₉₁ is residue 291 from the amino terminal end of the envelope glycoprotein. The expression of a recombinant product in Pichia from the pPIC9 vector is driven by the 30 methanol inducible promoter derived from the Pichia pastoris aox1 (alcohol oxidase 1) gene.

The pPIC9-60%E expression vector was transformed into spheroplasts of *P. pastoris* strain GS115 (*his4*) and transformants were selected for their

35 ability to grow on minimal medium without histidine supplementation. Strain GS115 and the protocol used for transformation were obtained from Invitrogen (San Diego, CA). Transformants were tested for their ability to



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express and secrete 60%E by growing selected clones in small cultures (5 ml). The transformants were grown to saturation (24 to 36 hrs.) in BMGY medium (1% yeast extract, 2% peptone, 100 mM potassium phosphate, pH 6.0, 1.34% yeast nitrogen base without amino acids, 4 x 10^{-5} % The cells were collected by biotin, 1% glycerol). centrifugation and suspended in one half the original culture volume with BMMY (identical to BMGY except the glycerol component of BMGY is replaced in BMMY with 0.5% methanol) medium and cultured for 48 hrs. Proteins secreted into the culture medium, as well as cellular proteins, were treated with endoglycosidase H_f (EndoH, New England Biolabs, Beverly, MA) and separated by SDS-PAGE. Western immunoblots probed with DEN-2 HMAF

15 indicated that the recombinants expressed significant levels of 60%E. Protein gels analyzed by Coomassie staining also showed strong levels of 60%E expression and secretion.

Example 25

<u>Construction of expression vector pPIC9-prM(mutSS)60%E</u> and secretion of 60%E by *P. pastoris* expressing MFa_LprM(mutSS)60%E

To construct clone pPIC9-prM(mutSS)60%E, a 25 strategy identical to that describe in Example 16 was used. Clone pLS6-prM(mutSS)60%E (described in Example 23) was digested with restriction endonuclease XhoI and XmaI and sequences within the $MF\alpha_{I}$ and dengue cloning adapter were replaced by oligonucleotides to remove an 30 extraneous XhoI site, to preserve a critical XhoI site, and to regenerate the Kex2 protease process site. The XhoI-SalI prM(mutSS)60%E fragment from pLS6 Δ prM(mutSS)60%E was ligated into the unique XhoI site of pPIC9. The nucleotide and amino acid sequences at the Nterminus of the fusion protein are identical to that 35 shown in Example 16 for $pLS6\Delta$ -prM(mutSS)80%E. The

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structure of the *Pichia* expression vector pPIC9prM(mutSS)60%E was confirmed by restriction digestion and DNA sequence analysis.

The pPIC9-prM(mutSS)60%E expression vector was 5 transformed into spheroplasts of P. pastoris strain GS115 (his4), and transformants were selected and evaluated as described in Example 12. After EndoH_f treatment of secreted proteins, a novel Coomassie Blue-staining band could be detected by SDS-PAGE. Immunoprobing with anti-10 domain B serum of Western blots of proteins prepared from the culture medium detected a diffuse area of reactivity ranging from 40 to 90 kD in the nonEndoH treated samples and a unique band of approximately 40 kD in the EndoH_f treated samples that corresponds to the band observed following Coomassie Blue staining. 15

Example 26

Construction of pMttbns-prM100%E, Expression of tPaprM100%E by Drosophila melanogaster Schneider cells, and Induction of a Virus Neutralizing Response by Immunizing Mice with tPa-prM100%E Membrane Preparations

For expression of DEN-2 PR159/S1 preMembrane protein amino acids 1-166 and Envelope glycoprotein amino acids 1-495 as a single continuous open reading frame in *Drosophila melanogaster* Schneider 2 tissue culture cells, DNA sequences encoding these proteins were obtained by digestion of the p48BSprME13 clone (described in Example 14) with the restriction enzymes Bg1II and SalI. This fragment was cloned into the unique Bg1II and XhoI restriction sites of plasmid pMtt Δ Xho (described in Example 13).

The cloning junctions of the resulting gene fusion expression vector, pMttprM100E, were confirmed by restriction endonuclease digestion and DNA sequence analysis. The partial nucleotide and predicted amino acid sequences of the tPA₁-prM100E fusion gene are:

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The TPA pre- and propeptide regions are delineated by pre' and pro-tPA', respectively, and the dengue sequences are indicated in bold type. The Phe₁ and Thr₁₆₆ residues are the N-terminal and C-terminal amino acid residues of prM, respectively. The Met₁ residue is the N-terminal amino acid of envelope glycoprotein and Ala₄₉₅ is residue 495 from the amino terminal end of the envelope glycoprotein (the carboxy-terminal residue).

As described previously in Example 13, Schneider 2 cells were cotransfected with pMttprM100%E DNA at ratios of 1:1, and 5:1, relative to pCOHygro DNA and selected for growth in medium containing 300 µg/ml hygromycin. Transfectants were induced with 200µM CuSO₄

20 and expression of prM100%E was examined at various times after induction. Proteins secreted into the culture medium as well as cellular proteins were separated by SDS-PAGE. Protein gels were analyzed by both Coomassie Blue staining and immunoprobing of corresponding Western

- 25 blots. Analysis of Coomassie Blue-stained SDS-PAGE gels failed to reveal a unique protein band either intracellularly or in the culture medium. However, a novel band was recognized intracellularly by polyclonal anti-dengue 2 hyperimmune mouse ascites fluid (anti-DEN-2
- 30 HMAF, gift of R. Putnak, WRAIR) in Western blot analysis. This ~60 kD immunoreactive band comigrated with viral Envelope derived from dengue-2 infected mosquito C6/36 cells suggesting the recombinant 100%E protein had been processed away from the preMembrane protein. Polyclonal
- 35 antisera to the pr portion of prM (from Peter Wright, Monash University, Australia) recognized a ~20 kD protein, which comigrated with viral prM, confirming that the 100%E produced in the transfected cells was processed



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from prM. In fact, no evidence of a higher molecular weight band that might correspond to unprocessed prM100%E was detected in any sample, suggesting that the proteolytic processing of prM from E is extremely efficient in Schneider cells. No 100%E was detected in the culture medium indicating that 100%E remains anchored in cell-associated membranes.

Immunoblots probed with anti-DEN-2 HMAF demonstrated that the amount of intracellular 100%E produced by the transfectants increased over time from day 1 post induction to 7 days post induction. The amount of 100%E detected intracellularly in the transfectants correlated with the cotransfection ratio. Sensitivity of the intracellular 100%E to

15 endoglycosidases was evaluated by molecular weight shift of the protein in SDS-PAGE and Western immunoblots following endoglycosidase treatment. Partial resistance of the recombinant 100%E to Endoglycosidase H_f (Endo H_f; New England Biolabs) digestion indicated that the product 20 contains N-linked glycosylation, and that the composition of the recombinant products probably represents a mixture of complex, high mannose, and/or hybrid glycosylation.

Schneider cells, transformed with pMtt-prM100%E expressing the tissue plasminogen activator leader fusion 25 protein tPA_L-prM100%E were cultured in serum-free medium (Excell; JRH Biosciences) and induced by addition of CuSO₄ to a final concentration in the culture medium of 0.2 nM (see Examples 13 and 17 for more detail on culture conditions). The cells were maintained in inducing 30 medium for seven days prior to harvesting. The cells

were harvested by centrifugation at 1000 X G in a Beckman TJ-6 refrigerated centrifuge. The cells were washed one time with phosphate buffer saline (PBS), resuspended in PBS and disrupted with a tissue homogenizer. Undisrupted
35 cells were removed by centrifugation at 1000 X G in an Eppendorf microcentrifuge. Cellular membranes were pelleted at 10,000 X G in an Eppendorf microcentrifuge

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and 100,000 X G in a Beckman L8-80M Ultracentrifuge. The 10,000 X G and 100,000 X G pellets were resuspended in PBS and pooled for mouse immunization. Negative control cells transformed with pCOHygro only (see Example 13) were cultured, induced, and harvested as described for the prM100%E-expressing cells. The immunogens were assayed by Western blot prior to immunization of mice.

Outbred Swiss Webster mice (Simonsen) were immunized intraperitoneally (I.P.) with 75 µg total protein of the prM100%E membrane preparation in Freund's complete adjuvant. Control animals were immunized with either 75 µg of total protein from the negative control membrane preparation or with saline only, each in Freund's complete adjuvant. The mice received three I.P. boosts, consisting of one half the priming dose in Freund's incomplete adjuvant, at two week intervals.

Following the first and second boosts, the animals were test bled (tail bleed) and the immune response was monitored using an indirect ELISA as

- 20 described in Example 18. One week after the final boost, the animals were bled out and the sera were tested for anti-DEN2 responsiveness by indirect ELISA and PRNT as described in Example 18. Results for the ELISA and PRNT assays are summarized in Table 7. The mice immunized
 25 with the membrane preparation containing 100%E developed
 - virus-neutralizing antibodies despite the crude nature of the immunogen, suggesting the utility of these immunogens as efficacious vaccine candidates.

Table 7							
Immune Response of Mice Immunized with Recombinant Dengue prM100%E							
mouse	antigen	adjuvant	20 titer	30 titer [®]	final titer	PRNT80	
			ELISA	ELISA	ELISA	titer®	
20-1	Saline	Freund's	< 1:50	<1:50	<1:100	<1:10	
20-2			<1:50	<1:50	<1:100	<1:10	
20-3			<1:50	< 1:50	< 1:100	<1:10	
20-4			< 1:50	< 1:50	< 1:100	< 1:10	
20-5			< 1:50	< 1:50	< 1:100	< 1:10	
23-1	pCoHygro	Freund's	<1:50	<1:50	<1:100	<1:10	
	membrane						
23-2			< 1:50	< 1:50	< 1:100	<1:10	
23-3			< 1:50	< 1:50	< 1:100	<1:10	
23-4			< 1:50	< 1:50	< 1:100	<1:10	
23-5			< 1:50	<1:50	< 1:100	< 1:10	
24-1	prM100%E	Freund's	1:200	1:1,600	>1:25600	1:320	
	membrane						
24-2			< 1:50	>1:1600	>1:1600	1:320	
24-3]		<1:50	>1:6000	>1:1600	1:80	
24-4			< 1:50	>1:6000	>1:25600	1:640	
24-5]		< 1:50	< 1:50	1:400	1:20	

^aIndirect ELISA titer using recombinant domain B as coating antigen. ^b80% reduction of plaque titer of DEN-2 NGC virus on Vero cells.

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Example 27

Immunoaffinity Purification of E and Subunits Preparation of immunoaffinity columns:

10 Materials:

Cyanogen bromide (CNBr) activated sepharose 4B, lot 17-0430-01 (Pharmacia Biotech, Upsala, Sweden). (HiTrap) Bead size 45-165 μm.

1 mM HCl

15 Binding buffer (100 mM NaHCO3, 500 mM NaCl pH 8.3) Low pH washing buffer (100 mM glycine, pH 2.5) Equilibration buffer (100 mM phosphate, pH 7.2) Monoclonal antibody 9D12, 4G2 or other (dengue envelope specific/reactive)

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Procedure: 0.6g (~2 ml) of freeze-dried HiTrap was suspended in 5 ml of 1 mM HCl for 15 min. The gel was poured onto a sintered glass filter and washed with 120 ml of 1 mM HCl, followed by 10 ml of binding buffer. Monoclonal antibody 9D12 was buffer exchanged into binding buffer using a Centricon 30 (Amicon). The concentration was estimated from its absorbance at 280 nm $(A_{280} \text{ of } 1 \text{ mg/ml} = 1.4)$. Approximately 20 mg of antibody was added to washed matrix to a final volume of about 4 The coupling reaction was allowed to proceed at 4°C ml. with end-over-end rotation. The antibody modified gel was then poured into a polypropylene column (Biorad) and washed. Washing involved sequential elution with

equilibration buffer - washing buffer - equilibration 10 buffer - binding buffer, for three cycles. The amount of unbound antibody was estimated by measuring absorbance of Typically, about 18 mg of the eluants at 280 nm. antibody bound per 2 ml of gel. The column was stored at 4 °C until used. Prior to running media containing 15 protein, the column was washed with washing buffer followed by equilibration buffer.

Purification of proteins by immunoaffinity chromatography. Chromatography was performed using low Media or buffer was supplied to the antibody-20 pressure. sepharose column by a peristaltic pump (Pharmacia or Gilson). The effluent from the column was monitored for absorbance at 280 nm using a Gilson 112 UV/vis detector equipped with a flow through cell. Media containing

25 classical domain B, domain B+T, MF080E and 80%E from Drosophila cells was circulated through the immunoaffinity column for antigen binding and washed with phosphate buffer pH 7.0. The bound protein is eluted with buffer at pH 2.5.

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Comparison of 9D12, 4G2 and 3H5 MAbs as affinity supports. N-hydroxysuccinimide (NHS) activated HiTrap columns (1 ml) were reacted with 5 mg of dengue envelope-specific MAb 9D12, 4G2 or 3H5 according to methods provided by the manufacturer. The columns were then tested for their ability to purify DEN-2 80%E from Drosophila cell culture medium, using the purification procedure described above. The 4G2 column had a greater

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capacity than the 9D12 column, while the 3H5 column did not bind significant amounts of DEN-2 80%E. Up to 150 μ g of 80%E could be bound by the 4G2 column, while the capacity of the 9D12 column was about 70 μ g.

General protocol for purification of DEN-2 80%E by 4G2 and 9D12 HiTrap columns. The columns (1 ml or 5 ml bed volume) are perfused at a rate of 1 ml per minute with media containing 80%E adjusted to pH 7.2. Following washing with 10 mM phosphate buffer, pH 7.2, the antigen is eluted with 100 mM glycine pH 2.5. The eluted product is neutralized with 1 M Tris, pH 7.5 (final concentration 0.2 M), and NaCl is added to a final concentration of 150 mM. The sample is then concentrated by membrane ultrafiltration in a Centricon 30 (Amicon).

Concentration and purity determination. The concentration of purified 80%E was determined by ELISA, using purified domain B as a standard. The purity of the antigen was estimated by SDS-PAGE followed by Coomassie staining or Western transfer, showing the immunoaffinity purified 80%E antigens to be greater than 98% pure.

Purification of $(MFa_L) 80\%E$ from Pichia media. (MFa_L)80%E was purified by immunoaffinity chromatography using a MAb 9D12 column as described. For each run, 50 ml of yeast culture medium [~1 µg/ml (MFa_L)80%E] was cycled for 3 hr through a 2 ml Sepharose column containing 20 mg of MAb 9D12. The column was washed with

5 mM phosphate buffer, pH 7.0, until a baseline absorbance at 280 nm was re-established. The bound protein was eluted with 5 mM phosphate buffer, pH 2.5,

30 then concentrated 20-fold by evaporation using a SpeedVac (Savant), and further concentrated by membrane ultrafiltration using a Centricon 30 (Amicon). The concentration of purified protein was estimated using a MAb 9D12/rabbit anti-domain B sandwich ELISA. This assay 35 incorporates a purified *Pichia*-secreted domain B standard, quantified by amino acid analysis (Protein and Nucleic Acid Facility, Stanford CA). The molar amount of

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 $(MFa_L) 80\%$ eluted from the columns was determined by comparison to a domain B standard curve. The concentration of $(MFa_L) 80\%$, was estimated by multiplying the concentration of domain B equivalents by five, to compensate for the mass/size difference between the domain B standard and $(MFa_L) 80\%$. Purity of the $(MFa_L) 80\%$ product was assayed by SDS-PAGE analysis on a 12% gel. Prior to SDS-PAGE analysis, the protein was treated with 5,000 units of Endo Hf (New England Biolabs) for two hours to remove heterogeneous carbohydrates added post-translationally to the molecule by the yeast. The treated protein migrates as a single band of molecular weight approximately 50,000 daltons.

Approximately 0.6 mg of (MFa_L)80%E was 15 recovered by this method (~50% yield). The purified subunit protein was stored in PBS at 4 °C, until used to immunize mice.

Example 28

Ion-Exchange and Size Exclusion Chromatography Purification of Envelope Subunit

Sterile-filtered, frozen stocks of supernatant from Drosophila cell cultures grown in a bioreactor were used as starting material for the purification procedure. The sample was thawed then concentrated and diafiltered using a tangential flow ultrafiltration device (Millipore, Bedford, MA) containing membranes with molecular weight cut-off of 10 kilodaltons. The sample was first concentrated to approximately 200 ml then diafiltered against 10mM Tris-succinate buffer (TSB), pH 8.1. The volume of TSB, pH 8.1, that was used for the diafiltration was at least twice the starting sample volume. Under the described conditions, 80%E was retained by the membranes along with the majority of the Drosophila proteins.

The retentate from the diafiltration was loaded directly onto a Fractogel DEAE (E-M Sciences, Gibbstown,

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NJ) column (1.4cm id. x 5.5cm ht.) that was equilibrated against TSB, pH 8.1. The sample was loaded at a flowrate of 1 ml per minute. Under these loading conditions, all of the 80%E in the sample was bound to the anionexchange matrix. After the sample was loaded, the column was washed with TSB, pH 8.1, until the absorbance at 280nm was zero. The column was then treated with TSB, pH 8.1, containing 100mM NaCl; this buffer eluted all bound proteins including 80%E. The product from this step contained large amounts of Drosophila contaminants; however, a significant contaminating Drosophila protein component as well as nonprotein components of the culture media were removed.

The peak fractions from the 100mM NaCl elution 15 profile were then adjusted to pH 5.7 and loaded onto a Fractogel-COO⁻ (E-M Sciences) column (1.4cm id. x 1.6cm ht.) that was equilibrated against TSB, pH 5.7, containing 100mM NaCl. The sample was loaded at a flowrate of 1 ml per minute. Most of the 80%E was bound to 20 the cation-exchange column, while many of the Drosophila proteins were not. Once the sample was loaded, the column was washed with TSB, pH 5.7 (no NaCl), followed by TSB, pH 8.1 (no NaCl). Bound 80%E was eluted with TSB, pH 8.1, containing 50 mM NaCl. The 80%E fraction from this step was approximately 80% pure. 25

The Fractogel-COO⁻ column product was further purified by gel filtration chromatography on a BioSep 3000 HPLC column (600x7.8mm, Phenomenex, Torrance, CA). Phosphate buffered saline (25 mM phosphate, 100mM NaCl), pH 6.8, was used as the running buffer at a flow rate of 0.5 ml per minute. Some of the 80%E was eluted in an early peak fraction along with various Drosophila contaminants; however, most of the material was eluted in a later peak. The 80%E in the later fraction was >90% pure.

Based on sandwich ELISA and SDS-PAGE analyses, the recombinant Dengue protein retains complete

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reactivity with MAb 9D12 that recognizes a conformationally sensitive envelope epitope. Reactivity with MAb 9D12 was retained during each step of the purification process.

Example 29

Expression and secretion of DEN-2 NS1 from Drosophila cells

A fully sequenced DEN-2 PR159 S1 NS1 clone was used as the starting construct. The plasmid clone, 10 p29D2NS1.2, contains a 1,055 bp fragment that corresponds to nucleotides 2422 and 3477 of the parent DEN-2 genomic map. The 5' end is flanked by a BglII site and the 3' end is flanked by two stop codons and a Sall site. This 15 BglII-SalI fragment was cloned into the Drosophila expression plasmid pMTT digested with the same two Two of the resulting recombinants were selected enzymes. for further characterization.

Two clones encoding the DEN2 NS1 protein, 20 pMttNS1.1 and pMttNS1.2, were transfected into S2 cells and transfectants selected for hygromycin resistance. The expression of the NS1 protein by the mixed populations was evaluated by Western blot analysis. Immunoreactive product was detected both intracellularly

25 and in the medium, with the vast majority of the product being secreted into the medium. A corresponding Coomassie blue staining band was seen on SDS-PAGE gels. The secreted 80%E product appeared to accumulate in the medium out to at least day 7 post induction. By Western

30 blot analysis, the NS1.2 transfectant population appears to express at slightly higher levels than the NS1.1 transfectant population. The immunoreactive product is recognized by anti-DEN2 HMAF and anti-NS1 MAb 7E11. Digestion of the secreted product with Endoglycosidase H (EndoH) and Peptide:N-glycosidase F (PNGase F)

demonstrated that the secreted product is sensitive to both EndoH and PNGase F digestion and therefore is

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glycosylated. Interestingly, the mobility shift on EndoH digestion is intermediate between the undigested and the PNGase F digested product, although all products are discrete bands on a Western blot. This may suggest that the two glycosylation sites are not identically modified by the secretory pathway of the S2 cells. Estimates of the expression level based upon Coomassie stained gel and Western blot is ~10 μ g/ml.

S2 DEN-2 NS1 transfectants were plated at 10 limiting dilutions in supplemented serum-free medium (IPL-41; Life Technologies Inc.). The subcultures were induced with CuSO₄ and the expression levels screened by immuno-dot blot, using MAb 7E11 (NS1 specific) for detection. Several of these cultures expressed and 15 secreted into the medium significant levels of NS1. The highest level expressors will be expanded and a clonal cell population secreting DEN-2 NS1 will be produced.

A bioreactor culture of S2 DEN-2 NS1 cells was grown using the *Drosophila* cell culture methods described 20 above. Based on Western dot blot analysis, we estimate that approximately 3-4-fold more NS1 was produced per unit volume in the bioreactor medium (~10 μ g/ml), compared to S2 DEN-2 NS1 cultures grown in tissue culture flasks. Analysis by SDS-PAGE of unpurified S2 DEN-2 NS1

25 culture medium, heated at 37 °C for 15 minutes in nonreducing buffer, showed a distinct band migrating at ~80-85 kDa, possibly corresponding to NS1 dimer. Western blots employing the anti-NS1 specific MAb, 7E11, confirmed that >90% of the NS1 secreted into culture

30 medium exists as a homodimer. SDS-PAGE/Western blot analysis also indicated that S2 DEN-2 NS1 cells secreted ~3-4-fold higher levels of NS1 dimer when cultured in the bioreactor, compared to culture in flasks.

For nondisruptive HPLC purification preserving 35 the dimer form a Hewlett-Packard Ti-series 1050 HPLC system was employed, with UV detection at 280 nm. Preliminary attempts with hydrophobic interaction

chromatography (TOSOHAAS phenyl-5PW) failed to resolve any of the medium components. Thus, an ion exchange method based on Cellufine sulfate (Amicon), was employed. S2 DEN-2 NS1 containing medium from a bioreactor culture

- 5 was exchanged into 20 mM succinic acid/sodium succinate, pH 6.0, and concentrated using a Minitan tangential flow ultrafiltration unit (Millipore, Bedford, MA) equipped with 10K molecular weight cut-off membranes. The concentrate was applied to a Cellufine sulfate column
- 10 equilibrated with a running buffer of 20 mM succinic acid/sodium succinate, pH 6.0, and the NS1 protein was eluted with a linear 0--3M NaCl gradient developed over 5 column volumes. NS1 eluted in the 1--1.5 M NaCl fraction and gave nearly complete recovery of 250% pure NS1 dimer, 15 as judged by SDS-PAGE analysis.

Further purification of the NS1 obtained in the previous paragraph employed Fractogel DEAE (EM Sciences, Gibbstown, NJ) which efficiently removes a major Drosophila cell contaminant present in the Cellufine sulfate fractions. Since the calculated isoelectric point of NS1 is similar to that of 80%E, we initially employed Fractogel DEAE column conditions that had been developed for 80%. NS1 medium was exchanged into running buffer, 20 mM Tris succinate, pH 8.1, containing 50 mM NaCl, applied to Fractogel DEAE, then NS1 was eluted with a NaCl step-gradient in pH 8.1 buffer. This protocol resulted in ~50-75% pure NS1 dimer, as determined by SDS-PAGE and Western blot.

Throughout this specification and the claims which follow, unless the context requires otherwise, the word "comprise", and variations such as "comprises" and "comprising", will be understood to imply the inclusion of a stated integer or step or group of integers or steps but not the exclusion of any other integer or step or group of integers or steps.





THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

1. An immunogenic composition which generates protective, neutralizing antibody responses to a *Flavivirus* in a murine host which responses confer protection against intracerebral challenge by the homologous *Flavivirus*, said strain of *Flavivirus* selected from the group consisting of a strain of dengue, a strain of Japanese encephalitis virus (JEV), a strain of yellow fever virus (YF), and a strain of tick-borne encephalitis virus (TBE)

which composition contains an adjuvant; and

a portion of the envelope protein (E) of the *Flavivirus* strain against which said responses are sought, which portion is 80%E, wherein said 80%E represents that portion of the envelope protein that constitutes 80% of its length starting from amino acid 1 at its N-terminus and which portion has been secreted as a recombinantly produced protein from *Drosophila* cells.

2. The immunogenic composition of claim 1 wherein the 80% E is encoded in a DNA construct operably linked downstream from human tissue plasminogen activator prepropeptide secretion leader (tPA_1).

3. The immunogenic composition of claim 1 wherein said *Flavivirus* is a dengue virus.

4. The immunogenic composition of claim 1 wherein said *Drosophila* cells are *D.melanogaster* Schneider cells.

5. The immunogenic composition of claim 4 wherein the *Flavivirus* is a dengue virus.

6. The immunogenic composition of claim 1 wherein said adjuvant is an alum adjuvant.

7. A method to protect a subject against a *Flavivirus* strain, said strain selected from the group consisting of a strain of dengue, a strain of YF, and a strain of JEV and a strain of TBE, which method comprises administering to a subject in need of such protection in an effective amount of the immunogenic composition of claim 1.

8. The method of claim 7 wherein said *Flavivirus* is a dengue virus.

9. A method to generate protective, neutralizing antibody responses in a subject against a *Flavivirus* strain, said strain selected from the group consisting of a strain of dengue, a strain of YF, a strain of JEV and a strain of TBE, which method comprises administering to a subject in need of such a protective, neutralizing antibody response an effective amount of the immunogenic composition of claim 1.

10. The method of claim 9 wherein said *Flavivirus* is a dengue virus.

11. Use of an immunogenic composition according to claim 1 in the manufacture of a medicament to protect a subject against a *Flavivirus* strain wherein said strain is selected from the group consisting of dengue, a strain of YF, a strain of JEV and a strain of TBE.

12. Use according to claim 11 wherein said *Flavivirus* is a dengue virus.

13. Use of an immunogenic composition according to claim 1 in the manufacture of a medicament to generate protective, neutralizing antibody responses in a subject against a *Flavivirus* strain wherein said strain is selected from the group consisting of dengue, a strain of YF, a strain of JEV and a strain of TBE.

14. Use according to claim 13 wherein said *Flavivirus* is a dengue virus.

15. An immunogenic composition which generates a neutralizing antibody response to a *Flavivirus* in a murine host against the homologous *Flavivirus*, said strain of *Flavivirus* selected from the group consisting of a strain of dengue, a strain of Japanese encephalitis virus (JEV), a strain of yellow fever virus (YF), and a strain of tick-borne encephalitis virus (TBE) which composition contains an adjuvant; and

a portion of the envelope protein (E) of the *Flavivirus* strain against which generation of said response is sought, which portion is 80% E, wherein said 80% E represents that portion of the envelope protein that constitutes 80% of its length starting from amino acid 1

at its N-terminus, and which portion has been secreted as a recombinantly produced protein from *Drosophila* cells.

16. The immunogenic composition of claim 15, wherein said *Drosophila* cells are Schneider cells.

17. The immunogenic composition of claim 15 wherein said *Flavivirus* is a dengue virus.

18. The immunogenic composition of claim 15 wherein the 80% E is encoded in a DNA construct operably linked downstream from human tissue plasminogen activator prepropeptide secretion leader (tPA_1).

19. The immunogenic composition of claim 15 wherein the adjuvant is an alum adjuvant.

20. A method to generate a neutralizing antibody response in a subject against a *Flavivirus* strain, said strain selected from the group consisting of a strain of dengue, a strain of YF, a strain of JEV and a strain of TBE, which method comprises administering to a subject in need of generating said response an effective amount of the immunogenic composition of claim 15.

21. The method of claim 20 wherein said *Flavivirus* is a dengue virus.

22. Use of an immunogenic composition according to claim 15 in the manufacture of a medicament to generate protective, neutralizing antibody responses in a subject against a *Flavivirus* strain wherein said strain is selected from the group consisting of dengue, a strain of YF, a strain of JEV and a strain of TBE.

23. Use according to claim 22 wherein said *Flavivirus* is a dengue virus.

24. An expression system for the recombinant production and secretion of a portion of an envelope (E) protein of a *Flavivirus* selected from the group consisting of dengue virus, Japanese encephalitis virus (JEV), tick-borne encephalitis virus (TBE) and yellow fever virus

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(YF), which expression system comprises *Drosophila* cells modified to contain a DNA molecule which comprises

(a) a first nucleotide encoding said portion of said E protein of the *Flavivirus* strain against which protection is sought, which portion is the N-terminal 80% of the protein from residue 1 to residue 395, and

(b) a second nucleotide sequence which encodes a secretory leader sequence or secretory signal sequence operably linked to said first nucleotide sequence and position so as to produce a fusion protein when said first and said second nucleotide sequences are expressed in a eucaryotic cell,

said encoding sequences operably linked to control sequences capable of effecting expression of said encoding nucleotide sequences in eucaryotic cells.

25. The expression system of claim 24, wherein said secretory leader sequence is human tissue plasminogen activator prepropeptide secretion leader (tPA_L) and optionally includes the premembrane leader of the E protein.

26. A method to produce a portion of an E protein of a *Flavivirus* selected from the group consisting of dengue virus, Japanese encephalitis virus (JEV), tick-borne encephalitis virus (TBE) and yellow fever virus (YF), which method comprises

(a) culturing the *Drosophila* cells of claim 24 in culture medium under conditions favourable for expression of the encoding nucleotide sequence so that the cells secrete said portion of the E protein of the *Flavivirus* strain against which protection is sought, which portion is the N-terminal 80% of the protein from residue 1 to residue 395 into the medium; and

(b) recovering the portion of the E protein from the culture medium.

27. A method to produce a portion of an E protein of a *Flavivirus* selected from the group consisting of dengue virus, Japanese encephalitis virus (JEV), tick-borne encephalitis virus (TBE) and yellow fever virus (YF), which method comprises

(a) culturing the *Drosophila* cells of claim 25 in culture medium under conditions favourable for expression of the encoding nucleotide sequence so that the cells secrete said

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portion of the E protein of the *Flavivirus* strain against which protection is sought, which portion is the N-terminal 80% of the protein from residue 1 to residue 395 into the medium; and

(b) recovering the portion of the E protein from the culture medium.

28. The expression system of claim 24 wherein the N-terminal 80% of the E protein from residue 1 to residue 395 is dengue virus E protein.

29. The method of claim 26 wherein the N-terminal 80% of the E protein from residue 1 to residue 395 is dengue virus E protein.

30. The method of claim 27 wherein the N-terminal 80% of the E protein from residue 1 to residue 395 is dengue virus E protein.

31. The expression system of claim 24, wherein the *Drosophila* cells are *Drosophila* Schneider cells.

32. The expression system of claim 25, wherein the *Drosophila* cells are *Drosophila* Schneider cells.

33. The method of claim 26, wherein the Drosophila cells are Drosophila Schneider cells.

34. The method of claim 27, wherein the *Drosophila* cells are *Drosophila* Schneider cells.

35. The expression system of claim 28, wherein the *Drosophila* cells are *Drosophila* Schneider cells.

36. The method of claim 29, wherein the *Drosophila* cells are *Drosophila* Schneider cells.



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37. The method of claim 30, wherein the *Drosophila* cells are *Drosophila* Schneider cells.

DATED this TWENTY SIXTH day of NOVEMBER, 1999 Hawaii Biotechnology Group, Inc. by DAVIES COLLISON CAVE Patent Attorneys for the Applicants WO 96/37221

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97 141 191 241 291 341	ÁTGA GCTGAAACGC GATTCTCACT GCCCTGGTGG ATTAAAAAGA GAGGCTTCAG	ATAACCAACG GAGAGAAACC TGGAATGCTG CATTCCTTCG TGGGGAACAA GAAAGAGATT	GAAAAAGGCG GCGTGTCAAC CAGGGACGAG TTTCCTAACA TTAAAAAATC GGAAGGATGC	AGAAACACGC TGTACAACAG GACCACTAAA ATCCCACCAA AAAGGCTATT TGAATATCTT	CTTTCAATAT TTGACAAAGA ATTGTTCATG CAGCAGGGAT AATGTTCTGA AAACAGGAGA preMembrane
391 441 541 591 641	CGTAGAACTG TCATCTGACC AAAAAGGGAA ACCCTCATGG TAAATGTCCC GCAACTCCAC	CAGGCATGAT ACACGCAACG AAGCCTTCTG CCATGGACCT TTTCTCAAGC GTCCACATGG	CATCATGCTG GAGAACCACA TTTAAGACAA TGGTGAGTTG AGAACGAACC GTAACTTATG Membrane	ATTCCAACAG CATGATCGTC AGGACGGCAC TGTGAAGACA AGAAGACATA GGACATGTAC	TGATGGCGTT AGTAGACAAG GAACATGTGT CAATCACGTA GATTGTTGGT CACCACAGGA
691 741 791 841	GAGCACAGAA GGGATTGGAG AACATGCCCA ATAATGGCCG	GAGAAAAAAG ACACGAACTG GAGAATTGAA CAATCCTGGC	ATCAGTGGCG AAACATGGAT ACTTGGATTC ATACACCATA	CTTGTTCCAC GTCATCAGAA TGAGACATCC GGAACGACGC	ACGTGGGAAT GGGGCCTGGA AGGCTTTACC ATTTCCAAAG Envelope
891 941 991 1041 1091 1141 1191 1241	AGTCCTGATA GCTGCATAGG AGTTGGGTTG AAAAAATAAA AACCCGCCAC ACAACAGACT GCAGGACAAA GAAATGGATG	TTCATCCTAC AATATCAAAT ACATAGTTTT CCAACACTGG CTTAAGGAAG CGCGCTGCCC AGGTTTGTCT TGGATTATTT	TGACAGCCAT AGGGACTTTG AGAACATGGA ACTTTGAACT TACTGTATAG AACACAAGGG GCAAACATTC GGAAAAGGAG	CGCTCCTTCA TGGAAGGAGT AGTTGTGTGA GATAAAAACA AGGCTAAACT GAACCCACCC CATGGTAGAC GCATCGTGAC	ATGACAATGC GTCAGGAGGG CGACGATGGC GAAGCCAAAC GACCAACACG TGAATGAAGA AGAGGATGGG CTGTGCCATG
1291	TTCACATGCA	AAAAGAACAT	GGAGGGAAAA	ATTGTGCAGC	CAGAAAACCT
1341 1391 1441 1541 1591 1641 1691 1741 1791	GGAATACACT GAAATGACAC TCCATCACAG CTCTCCAAGA AAGACAAAGC CCATGGCTGC GACACTGGTC TCTTAGGATC GAAATCCAGA CAGGCTGAGA	GTCGTŤATAA AGGAAAACAT AGGCGGAACT ACGGGCCTCG TTGGCTGGTG CCGGAGCAGA ACCTTCAAAA CCAAGAGGGG TGTCATCAGG ATGGACAAAT	CACCTCATTC GGTAAAGAAG GACAGGCTAT ACTTCAATGA CACAGACAAT CACAGACAAT CACACAAGGA ATCCCCATGC GCCATGCATA AAACCTGCTG TACAACTTAA	AGGGGAAGAA TCAAGATAAC GGCACTGTTA GATGGTGTTG GGTTCCTAGA TCAAATTGGA GAAAAAACAG CAGCACTCAC TTCACAGGAC AGGGATGTCA	CATGCAGTCG ACCACAGAGC CGATGGAGTG CTGCAAATGA CCTACCGTTG TACAGAAAGA GATGTTGTTG AGGGGCTACG ATCTTAAGTG TACTCCATGT

FIG. 2A

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1841	GCACAGGAAA	GTTTAAAGTT	GTGAAGGAAA	TAGCAGAAAC	ACAACATGGA
1891 1941	ACAATAGTCA CCCTTTTGAG	TTAGAGTACA ATAATGGATC	ATATGAAGGA Tggaaaaaag	GACGGCTCTC ACATGTTTTG	CATGCAAGAT GGCCGCCTGA
1991 2041 2091 2141 2191 2241 2291 2341	TCACAGTCAA GCAGAACCTC ACAATTGAAG TTGAGACAAC GCCTGGGATT TCTCCACCAG CATGGACTAT AACTCACGTA	CCCAATTGTA CATTCGGAGA CTGGACTGGT AATGAGGGGA TTGGATCTCT GTTTTTGGAG GAAGATCCTC GCACATCACT	ACAGAAAAGG CAGCTACATC TCAAGAAAGG GCGAAAAGAA GGGAGGAGTG CAATCTACGG ATAGGAGTTA GTCTGTGTCA	ACAGCCCAGT ATCATAGGAG AAGTTCCATC TGGCCATTTT TTCACATCAA GGCTGCTTTC TCATCACATG CTGGTATTAG NS1	CAACATAGAA TGGAACCAGG GGCCAAATGT GGGCGACACA TAGGAAAGGC AGTGGGGTCT GATAGGAATG TGGGAATCGT
2391 24491 2591 2691 2691 2691 2691 2791 2891 2991 2991 2991 3091 3191 3191 3191 3391	GACACTGTAC GCTGGAAGAA AACGTGCATA AAAACTGGCT TCCGCTCAGT GAATTGAATC AGGAGACATC AGGAGACATC ACCCACTGA CTCTCCACAG AGCAGAATGC ACTACGGCTT AAGCAGGATG CAACAGAGCC ATGGTAATT GACCAGGCTA GAGATGGAAAC TCATAACGGA AGAGGTGAGG	N TTGGGAGTTA CAAAGAACTA CATGGACAGA TCAGCCATCC AACAAGACTG ATATTCTATC AAAGGAATCA GTTGAGGTAT AACTCCATAA CCCAACACAA TGGAGTATTC CATTTTGTGA GTCCATGCTG GAAGATAGAG AGTCACACACA CCAAAGAATT TCACACACAA TTGATTTCTG AGAGGGCCCT ATGGTGTTGT ATGGATGCTG	TGGTGCAGGC AAATGTGGCA ACAATACAAG AGAAAGCTCA GAAAATCTTA AGAAAATCTTA AGAAAATGAA TGCAGGTAGG TCATGGAAAA TCAGACCTTC ACAGAGCTTG ACTACCAATA CTCAAAACTC ATATGGGTTA AAAGCTTCTT TCTATGGAGT TCGCTGGACC CGAAGGGACT CTTTAAGAAC CGATCTTGCA GTACGGGATG.	CGATAGTGGT GTGGAATATT TTCCAACCAG TGAAGAGGGC TGTGGAAACCA GTGAAACTGA AAAACGATCT CATGGGGTAA CTCATTGATG GAATTCACTA TATGGCTAAG ATGTCAGCGG TTGGATAGAA TCATTGAAGT AATGGAGTGC AGTGTCACAA CTTGGCATCT ACAGTGGTGGG AACCACTGCC CACTACCACC	TGCGTTGTGA CGTCACAGAT AATCCCCTTC ATCTGTGGAA AATAACATCA CCATCATGAC CTGCGGCCTC AGCGAAAATG GTCCCGAAAC GAAGTTGAGGG ATTGAGAGAA CCATAAAGGA AGCGCACTCA CAAAAGTTGC TAGAAAGCGA CATAATAACA AGGCAAGCTT TAACCGAGGA ACCAGGAAAAC ACTAAGATAC ACTAAGATAC

DEN-2 PR159/S1 cDNA coding strand sequence for Capsid, preMembrane, Envelope, and NS1 is given (Hahn etal. 1988). The start of genes are indicated. * indicates correction to published sequence Differences in the wild-type DEN-2 PR159 sequence are indicated above the PR159/S1 sequence.

FIG. 2B

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CTTTCAATAT GCTGAAACGC GAGAGAAACC GCGTGTCAAC TGTACAACAG TTGACAAAGA 131 ProPheAsnMet LeuLysArg GluArgAsn ArgValSerThr ValGlnGln LeuThrLys> GATTCTCACT TGGAATGCTG CAGGGACGAG GACCACTAAA ATTGTTCATG GCCCTGGTGG 191 ArgPheSerLeu GlyMetLeu GlnGlyArg GlyProLeuLys LeuPheMet AlaLeuVal> CATTCCTTCG TTTCCTAACA ATCCCACCAA CAGCAGGGAT ATTAAAAAGA TGGGGAACAA AlaPheLeuArg PheLeuThr lleProPro ThrAlaGlylle LeuLysArg TrpGlyThr> 251 TTAAAAAATC AAAGGCTATT AATGTTCTGA GAGGCTTCAG GAAAGAGATT GGAAGGATGC 311 IleLysLysSer LysAlalle AsnValLeu ArgGlyPheArg LysGlulle GlyArgMet> TGAATATCTT AAACAGGAGA CGTAGAACTG CAGGCATGAT CATCATGCTG ATTCCAACAG 371 LeuAsnIleLeu AsnArgArg ArgArgThr AlaGlyMetIle IleMetLeu IleProThr> TGATGGCGTT TCATCTGACC ACACGCAACG GAGAACCACA CATGATCGTC AGTAGACAAG ValMetAlaPhe HisLeuThr ThrArgAsn GlyGluProHis MetIleVal SerArgGin> 431 PreMembrane AAAAAGGGAA AAGCCTTCTG TTTAAGACAA AGGACGGCAC GAACATGTGT ACCCTCATGG 491 GluLysGlyLys SerLeuLeu PheLysThr LysAspGlyThr AsnMetCys ThrLeuMet> CCATGGACCT TGGTGAGTTG TGTGAAGACA CAATCACGTA TAAATGTCCC TTTCTCAAGC AlaMetAspLeu GlyGluLeu CysGluAsp ThrileThrTyr LysCysPro PheLeuLys 551 AGAACGAACC AGAAGACATA GATTGTTGGT GCAACTCCAC GTCCACATGG GTAACTTATG 611 GlnAsnGluPro GluAspIle AspCysTrp CysAsnSerThr SerThrTrp ValThrTyr> GGACATGTAC CACCACAGGA GAGCACAGAA GAGAAAAAAG ATCAGTGGCG CTTGTTCCAC 671 GlyThrCysThr ThrThrGly GluHisArg ArgGluLysArg SerValAla LeuValPro> Membrane ACGTGGGAAT GGGATTGGAG ACACGAACTG AAACATGGAT GTCATCAGAA GGGGCCTGGA 731 HisValGlyMet GlyLeuGlu ThrArgThr GluThrTrpMet SerSerGlu GlyAlaTrp> AACATGCCCA GAGAATTGAA ACTTGGATTC TGAGACATCC AGGCTTTACC ATAATGGCCG -791 LysHisAlaGin ArglieGiu ThrTrpIle LeuArgHisPro GlyPheThr IleMetAla> 851

51 CAATCCTGGC ATACACCATA GGAACGACGC ATTTCCAAAG AGTCCTGATA TTCATCCTAC AlalleLeuAla TyrThrile GlyThrThr HisPheGlnArg ValLeuIle PhelleLeu>

FIG. 3A

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	911	TGACAGCCAT CGCTCCTTCA LeuThrAlaile AlaProSer	ATGACAATGC GCTGCATAGG AATATCAAAT MetThrMet ArgCysIleGly IleSerAsn	AGGGACTTTG ArgAspPhe>
	971	TGGAAGGAGT GTCAGGAGGG ValGluGlyVal SerGlyGly	AGTTGGGTTG ACATAGTTTT AGAACATGGA SerTrpVal AspileValLeu GluHisGly	AGTTGTGTGA SerCysVal>
	1031	CGACGATGGC AAAAAATAAA ThrThrMetAla LysAsnLys	CCAACACTGG ACTTTGAACT GATAAAAACA ProThrLeu AspPheGluLeu lleLysThr	GAAGCCAAAC GluAlaLys>
	1091	AACCCGCCAC CTTAAGGAAG GInProAlaThr LeuArgLys	TACTGTATAG AGGCTAAACT GACCAACACG TyrCysIle GluAlaLysLeu ThrAsnThr	ACAACAGACT ThrThrAsp>
	1151	CGCGCTGCCC AACACAAGGG SerArgCysPro ThrGlnGly	GAACCCACCC TGAATGAAGA GCAGGACAAA GluProThr LeuAsnGluGlu GlnAspLys	AGGTTTGTCT ArgPheVal>
	1211	GCAAACATTC CATGGTAGAC CysLysHisSer MetValAsp	AGAGGATGGG GAAATGGATG TGGATTATTT ArgGlyTrp GlyAsnGlyCys GlyLeuPhe	GGAAAAGGAG GlyLysGly>
WT	1271	GCATCGTGAC CTGTGCCATG GlylleValThr CysAlaMet	GAA(Glu) TTCACATGCA AAAAGAACAT GGAGGGAAAA PheThrCys LysLysAsnMet GluGlyLys	ATTGTGCAGC IleValGln>
WT	1331	CAGAAAAACCT GGAATACACT ProGluAsnLeu GluTyrThr	GTG(Val) GTCGTTATAA CACCTCATTC AGGGGAAGAA ValVallle ThrProHisSer GlyGluGlu	CATGCAGTCG HisAlaVal>
	1391	GAAATGACAC AGGAAAACAT GlyAsnAspThr GlyLysHis	GGTAAAGAAG TCAAGATAAC ACCACAGAGC GlyLysGlu ValLysIleThr ProGinSer	TCCATCACAG SerlleThr>
	1451	AGGCGGAACT GACAGGCTAT GluAlaGluLeu ThrGlyTyr	GGCACTGTTA CGATGGAGTG CTCTCCAAGA GlyThrVal ThrMetGluCys SerProArg	ACGGGGCCTCG ThrGlyLeu>
	1511	ACTTCAATGA GATGGTGTTG AspPheAsnGlu MetValLeu	CTGCAAATGA AAGACAAAGC TTGGCTGGTG LeuGlnMet LysAspLysAla TrpLeuVal	CACAGACAAT HisArgGln>
	1571	GGTTCCTAGA CCTACCGTTG TrpPheLeuAsp LeuProLeu	CCATGGCTGC CCGGAGCAGA CACACAAGGA ProTrpLeu ProGlyAlaAsp ThrGlnGly	TCAAATTGGA SerAsnTrp>
	1631	TACAGAAAGA GACACTGGTC IleginLysgiu ThrLeuVal	ACCTTCAAAA ATCCCCATGC GAAAAAACAG ThrPheLys AsnProHisAla LysLysGln	GATGTTGTTG AspValVal>
	1691	TCTTAGGATC CCAAGAGGGG ValLeuGlySer GlnGluGly	GCCATGCATA CAGCACTCAC AGGGGGCTACG AlaMetHis ThrAlaLeuThr GlyAlaThr	GAAATCCAGA GluIleGln>
	1751	TGTCATCAGG AAACCTGCTG MetSerSerGly AsnLeuLeu	TTCACAGĠAC ATCTTAAGTG CAGGCTGAGA PheThrGly HisLeuLysCys ArgLeuArg	ATGGACAAAT MetAspLys>

FIG. 3B

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FIG. 3C

1811	TACAACTTAA AGGGATGTCA LeuGlnLeuLys GlyMetSer	TACTCCATGT GCACAGGAAA GTTTAAAGTT GTGAAGGAAA TyrSerMet CysThrGlyLys PheLysVal ValLysGlu>
1871	TAGCAGAAAC ACAACATGGA IleAlaGluThr GlnHisGly	ACAATAGTCA TTAGAGTACA ATATGAAGGA GACGGCTCTC ThrileVal IleArgValGln TyrGluGly AspGlySer>
1931	CATGCAAGAT LCCTTTTGAG ProCysLysIle ProPheGlu	ATAATGGATC TGGAAAAAAG ACATGTTTTG GGCCGCCTGA IleMetAsp LeuGluLysArg HisValLeu GlyArgLeu>
1991	TCACAGTCAA CCCAATTGTA IleThrValAsn ProlleVal	AGT(Ser) ACAGAAAAGG ACAGCCCAGT CAACATAGAA GCAGAACCTC ThrGluLys AspSerProVal AsnileGlu AlaGluPro>
2051	CATTCGGAGA CAGCTACATC ProPheGly op Sertyrile	ATCATAGGAG TGGAACCAGG ACAATTGAAG CTGGACTGGT IlelleGly ValgluProgly GlnLeuLys LeuAspTrp>
2111	TCAAGAAAGG AAGTTCCATC PheLysLysGly SerSerIle	GGCCAAATGT TTGAGACAAC AATGAGGGGA GCGAAAAGAA GlyGlnMet PheGluThrThr MetArgGly AlaLysArg>
2171	TGGCCATTTT GGGCGACACA MetAlalleLeu GlyAspThr	GCCTGGGATT TTGGATCTCT GGGAGGAGTG TTCACATCAA AlaTrpAsp PheGlySerLeu GlyGlyVal PheThrSer>
2231	TAGGAAAGGC TCTCCACCAG IleGlyLysAla LeuHisGln	GTTTTTGGAG CAATCTACGG GGCTGCTTTC AGTGGGGTCT ValPheGly AlalleTyrGly AlaAlaPhe SerGlyVal>
2291	CATGGACTAT GAAGATCCTC SerTrpThrMet LysIleLeu	ATAGGAGTTA TCATCACATG GATAGGAATG AACTCACGTA IleGlyVal IleIleThrTrp IleGlyMet AsnSerArg>
2351	GCACATCACT GTCTGTGTCA SerThrSerLeu SerValSer	CTGGTATTAG TGGGAATCGT GACACTGTAC TTGGGAGTTA LeuValLeu ValGlyIleVal ThrLeuTyr LeuGlyVal>
2411	TGGTGCAGGC CGATAGTGGT MetValGlnAla AspSerGly	TGCGTTGTGA GCTGGAAGAA CAAAGAACTA AAATGTGGCA CysValVal SerTrpLysAsn LysGluLeu LysCysGly>
2471	GTGGAATATT CGTCACAGAT SerGlyIlePhe ValThrAsp	AACGTGCATA CATGGACAGA ACAATACAAG TTCCAACCAG AsnValHis ThrTrpThrGlu GlnTyrLys PheGlnPro>
2531	AATCCCCTTC AAAACTGGCT GluSerProSer LysLeuAla	TCAGCCATCC AGAAAGCTCA TGAAGAGGGC ATCTGTGGAA SerAlalle GlnLysAlaHis GluGluGly lleCysGly>
2591	TCCGCTCAGT AACAAGACTG IleArgSerVal ThrArgLeu	GAAAATCTTA TGTGGAAACA AATAACATCA GAATTGAATC GluAsnLeu MetTrpLysGln IleThrSer GluLeuAsn>
2651	ATATTCTATC AGAAAATGAA HisileLeuSer GluAsnGlu	GTGAAACTGA CCATCATGAC AGGAGACATC AAAGGAATCA ValLysLeu ThrIleMetThr GlyAspile LysGlyIle>
2711	TGCAGGTAGG AAAACGATCT MetGlnValGly LysArgSer	CTGCGGCCTC AACCCACTGA GTTGAGGTAT TCATGGAAAA LeuArgPro GinProThrGiu LeuArgTyr SerTrpLys>

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2771	CATGGGGTAA AGCGAAAATG	CTCTCCACAG AACTCCATAA TCAGACCTTC	CTCATTGATG
	ThrTrpGlyLys AlaLysMet	LeuSerThr GluLeuHisAsn GlnThrPhe	LeuileAsd>
2831	GTCCCGAAAC AGCAGAATGC	CCCAACACAA ACAGAGCTTG GAATTCACTA	GAAGTTGAGG
	GlyProGluThr AlaGluCys	ProAsnThr AsnArgAlaTrp AsnSerLeu	GluValGlu>
2891	ACTACGGCTT TGGAGTATTC	ACTACCAATA TATGGCTAAG ATTGAGAGAA	AAGCAGGATG
	AspTyrGlyPhe GlyValPhe	ThrThrAsn IleTrpLeuArg LeuArgGlu	LysGlnAsp>
2951	CATTTTGTGA CTCAAAACTC	ATGTCAGCGG CCATAAAGGA CAACAGAGCC	GTCCATGCTG
	AlaPheCysAsp SerLysLeu	MetSerAla AlaIleLysAsp AsnArgAla	ValHisAla>
3011	ATATGGGTTA TTGGATAGAA	AGCGCACTCA ATGATACATG GAAGATAGAG	AAAGCTTCTT
	AspMetGlyTyr TrpIleGlu	SerAlaLeu AsnAspThrTrp LysIleGlu	LysAlaSer>
3071	TCATTGAAGT CAAAAGTTGC	CACTGGCCAA AGTCACACAC TCTATGGAGT	AATGGAGTGC
	PheileGluVal LysSerCys	HisTrpPro LysSerHisThr LeuTrpSer	AsnGLyVal>
3131	TAGAAAGCGA GATGGTAATT	CCAAAGAATT TCGCTGGACC AGTGTCACAA	CATAATAACA
	LeuGluSerGlu MetVallle	ProLysAsn PheAlaGlyPro ValSerGln	HisAsnAsn>
3191	GACCAGGCTA TCACACACAA	ACAGCAGGAC CTTGGCATCT AGGCAAGCTT	GAGATGGACT
	ArgProGlyTyr HisThrGln	ThrAlaGly ProTrpHisLeu GlyLysLeu	GLuMetAsp>
3 251	TTGATTTCTG CGAAGGGACT	ACAGTGGTGG TAACCGAGGA CTGTGGAAAC	AGAGGGGCCCT
	PheAspPheCys GluGlyThr	ThrValVal ValThrGluAsp CysGlyAsn	ArgGlyPro>
3311	CTTTAAGAAC AACCACTGCC	TCAGGAAAAC TCATAACGGA ATGGTGTTGT	CGATCTTGCA
	SerLeuArgThr ThrThrAla	SerGlyLys LeulleThrGlu TrpCysCys	ArgSerCys>
3371	CACTACCACC ACTAAGATAC	AGAGGTGAGG ATGGATGCTG GTACGGGATG	GAAATCAGAC
	ThrLeuProPro LeuArgTyr	ArgGlyGlu AspGlyCysTrp TyrGlyMet	GluIleArg>
3431	CATTGAAAGA GAAAGAAGAA ProleulysGlu lysGluGlu	AATCTGGTCA GTTCTCTGGT CACAGCC AsnLeuVal SerSerLeuVal ThrAla	

FIG. 3D

Oligonucleotide Mutagenesis to Generate 60%E and Domain B



Shown are the nucleotide sequence and the corresponding translation of DEN-2 PR159/S1 E sequences between genomic nucleotides 1789 and 1848, the sequence of the oligonucleotide used in mutagenesis, and the resulting sequence and corresponding translations following mutagenesis. Dengue nucleotide sequence is uppercase, non-dengue nucleotides are lower case, inserted restriction endonuclease sites are indicated, and conserved cysteines are underlined.

FIG. 4

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5318	ATGAGATTTCCTTCAATTTTTACTGCAGTTTTATTCGCAGCATCCTCCGC MetArgPheProSerIlePheThrAlaValLeuPheAlaAlaSerSerAla> - MF& secretion signal peptide
5368	ATTAGCTGCTCCAGTCAACACTACAACAGAAGATGAAACGGCACAAATTC LeuAlaAlaProValAsnThrThrThrGluAspGluThrAlaGlnIle> ▲ Signalase cleavage H→MF∝ propeptide
5418	3 CGGCTGAAGCTGTCATCGGTTACTCAGATTTAGAAGGGGATTTCGATGTT ProAlaGluAlaVallleGlyTyrSerAspLeuGluGlyAspPheAspVal>
5468	3 GCTGTTTTGCCATTTTCCAACAGCACAAATAACGGGTTATTGTTTATAAA AlaValLeuProPheSerAsnSerThrAsnAsnGlyLeuLeuPheIleAsn>
5518	3 TACTACTATTGCCAGCATTGCTGCTAAAGAAGAAGGGGGTATCTCTCGAGA ThrThrIleAlaSerIleAlaAlaLysGlnGluGlyValSerLeuGlu>
5568	AAAGGGAGGCTGGGATGTCATACTCCATGTGCACAGGAAAGTTTAAAGTT LysArgGluAlaGlyMetSerTyrSerMetCysThrGlyLysPheLysVal>
5618	<pre>GTGAAGGAAATAGCAGAAACACAACATGGAACAATAGTCATTAGAGTACA ValLysGluIleAlaGluThrGlnHisGlyThrIleVallleArgValGln></pre>
5668	ATATGAAGGAGACGGCTCTCCATGCAAGATCCCTTTTGAGATAATGGATC TyrGluGlyAspGlySerProCysLysIleProPheGluIleMetAsp>
5718	3 TGGAAAAAAGACATGTTTTGGGCCGCCTGATCACAGTCAATCCAATTGTA LeuGluLysArgHisValLeuGlyArgLeuIleThrValAsnProIleVal>
5768	ACAGAAAAGGACAGCCCAGTCAACATAGAAGCAGAACCTCCATTCGGAGA ThrGluLysAspSerProValAsnIleGluAlaGluProProPheGlyAsp>
5818	3 CAGCTACATCATCATAGGAGTGGAACCAGGACAATTGAAGCTGGACTGGT SerTyrIleIleIleGlyValGluProGlyGlnLeuLysLeuAspTrp>
5868	3 TCAAGAAAGGATAA PheLysLysGlyEnd>

FIG. 7

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150 160 170 180 190 200 210 GTGAAGGAAA TAGCAGAAAC CATGGA ACAATAGTCA GTTTAAAATT ACA GAGTACA TATGAAGGA CAAATTTTAA CACTTCCTTT ATCGTCTTTG TGTTGTACCT TGTTATCAGT AATCTCATGT TATACTTCCT PheLysile ValLysGlu IleAlaGluThr GinHisGly ThrileVal IleArgValGln TyrGluGly> ſ C 23Q 220 240 250 260 270 280 TTTTGAG ATAATGGATC GACGGCTCTC CATGCAAGAT TGGAAAAAAG ACATGTTTTG GGCCGCCTGA CTGCCGAGAG GTACGTTCTA GGGAAAACTC TATTACCTAG ACCTTTTTC TGTACAAAAC CCGGCGGACT AspGlySer ProCysLysIle ProPheGlu IleMetAsp LeuGluLysArg HisValleu GlyArgLeu> TATTACCTAG ACCTTTTTTC TGTACAAAAC RGARSP-DOMB С С ſ C 29**0** 33Q 300 310 32Q 340 350 ACAGAAAAGG ACAGTCCAGT TCACAGTCAA CCCAATTGTA CAACATAGAA GCAGAACCTC CATTCGGAGA TGTCTTTTCC TGTCAGGTCA GTTGTATCTT AGTGTCAGTT GGGTTAACAT CGTCTTGGAG GTAAGCCTCT IleThrValAsn ProlleVal ThrGluLys AspSerProVal AsnileGlu AlaGluPro ProPheGlyAsp> C ______ RGARSP-DOMB______C____C____> 39**0** 360 37Q 38Q 40**0** 410 TGGAACCAGG ACAATTGAAG CAGCTACATC ATCATAGGAG CTGGACTGGT TCAAGAAAGG ATAATAG GTCGATGTAG TAGTATCCTC ACCTTGGTCC TGTTAACTTC GACCTGACCA AGTTCTTTCC TATTATC SerTyrIle IleIleGly ValGluProGly GlnLeuLys LeuAspTrp PheLysLysGly EndEnd> RGARSP-DOMB C ſ C C

FIG. 9B

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FIG. 12

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