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(54) Title: DERIVATIVES OF PNEUMOCOCCAL CHOLINE BINDING PROTEINS FOR VACCINES

(57) Abstract

The present invention provides bacterial immunogenic agents for adminstration to humans and non-human animals to stimulate an immune response. It particularly relates to the vaccination of mammalian species with pneumococcal derived polypeptides that include an alpha helix but exclude a choline binding region as a mechanism for stimulating production of antibodies that protect the vaccine recipient against infection by pathogenic bacterial species. In another aspect the invention provides antibodies against such proteins and protein complexes that may be used as diagnostics and/or as protective/treatment agents for pathogenic bacterial species.

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DERIVATIVES OF PNEUMOCOCCAL CHOLINE BINDING PROTEINS FOR VACCINES

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This application claims the benefit of U.S. Prov. Appl'n Serial No. 60/085,743, filed May 15, 1998 and U.S. Prov. Appl'n Serial NO 60/080,878, filed April 7, 1998.

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This invention relates generally to the field of bacterial antigens and their use, for example, as immunogenic agents in humans and animals to stimulate an More specifically, it relates to the immune response. 15 vaccination of mammalian species with a polypeptide comprising an alpha helix-forming polypeptide obtained from a choline binding polypeptide as a mechanism for stimulating production of antibodies that protect the recipient against infection by vaccine pathogenic bacterial species. Further, the invention relates to 20 antibodies and antagonists against such polypeptides useful in diagnosis and passive immune therapy with respect to diagnosing and treating such pneumococcal infections.

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particular aspect, the present invention In a relates to the prevention and treatment of pneumonococcal infections such of the middle infections as ear, nasopharynx, lung and bronchial areas, blood, CSF, and the 30 like, that are caused by pneumonococcal bacteria. In this regard, certain types of Streptococcus pneumoniae are of particular interest.

S. pneumoniae is a gram positive bacteria which is a 35 major causative agent in invasive infections in animals and humans, such as sepsis, meningitis, otitis media and lobar pneumonia (Tuomanen, et al. NEJM 322:1280-1284 (1995)). As part of the infective process, pneumococci 5

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readily bind to non-inflamed human epithelial cells of the upper and lower respiratory tract by binding to eukaryotic carbohydrates in a lectin-like manner (Cundell et al., Micro. Path. 17:361-374 (1994)). Conversion to invasive pneumococcal infections for bound bacteria may involve the generation of inflammatory factors local which may activate the epithelial cells to change the number and type of receptors on their surface (Cundell, et al., Nature. 377:435-438 (1995)).Apparently, one such receptor, platelet activating factor (PAF) is engaged by the pneumococcal bacteria and within a very short period of time (minutes) from the appearance of PAF, pneumococci strongly enhanced adherence and invasion exhibit of tissue. Certain soluble receptor analogs have been shown to prevent the progression of pneumococcal infections (Idanpaan-Heikkila et al., J. Inf. Dis., 176:704-712 (1997)).

A family of choline binding proteins (CBPs), which are non-covalently bound to phosphorylcholine, are present on the surface of pneumococci and have a non-covalent association with teichoic acid or lipoteichoic acid. An example of such family is choline binding protein A (CbpA), an approximately 75kD weight type of CBP which includes a unique N-terminal domain, a proline rich region, and a C-terminal domain comprised of multiple 20 amino acid repeats responsible for binding to choline. A segment of the N-terminal portion of CbpA protein forms an alpha helix as part of its three-dimensional structure.

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Accordingly, it is an object of the present invention to provide a polypeptide having broad protection against pneumococcal infections.

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Definitions

In order to facilitate understanding of the description below and the examples which follow certain

frequently occurring methods and/or terms will be described.

"Plasmids" are designated by a lower case p preceded 5 and/or followed by capital letters and/or numbers. The starting plasmids herein are either commercially available, publicly available on an unrestricted basis, or can be constructed from available plasmids in accord with published procedures. In addition, equivalent plasmids to 10 those described are known in the art and will be apparent to the ordinarily skilled artisan.

"Digestion" of DNA refers to catalytic cleavage of the DNA with a restriction enzyme that acts only at 15 certain sequences in the DNA. The various restriction enzymes used herein are commercially available and their reaction conditions, cofactors and other requirements were used as would be known to the ordinarily skilled artisan. For analytical purposes, typically 1 µg of plasmid or DNA 20 fragment is used with about 2 units of enzyme in about 20 μ l of buffer solution. For the purpose of isolating DNA fragments for plasmid construction, typically 5 to 50 µg of DNA are digested with 20 to 250 units of enzyme in a larger volume. Appropriate buffers and substrate amounts 25 for particular restriction enzymes are specified by the manufacturer. Incubation times of about 1 hour at 37°C are ordinarily used, but may vary in accordance with the supplier's instructions. After digestion the reaction is electrophoresed directly on a polyacrylamide gel to

30 isolate the desired fragment.

Size separation of the cleaved fragments is performed using 8 percent polyacrylamide gel described by Goeddel, D. *et al.*, Nucleic Acids Res., 8:4057 (1980).

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"Oligonucleotides" refers to either а single stranded polydeoxynucleotide or two complementary polydeoxynucleotide strands which may be chemically

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Such synthetic oligonucleotides have no 5' synthesized. phosphate and thus will not ligate to another oligonucleotide without adding a phosphate with an ATP in the presence of a kinase. A synthetic oligonucleotide will ligate to а fragment that has not been dephosphorylated.

"Ligation" refers to the process of forming phosphodiester bonds between two double stranded nucleic 10 acid fragments (Maniatis, T., et al., Id., p. 146). Unless otherwise provided, ligation may be accomplished using known buffers and conditions with 10 units to T4 DNA ligase ("ligase") per 0.5 µg of approximately equimolar amounts of the DNA fragments to be ligated.

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"HPS portion" as used herein refers to an amino acid sequence as set forth in SEQ ID NO:2 for a choline binding protein ("CBP") of a pneumococcal bacteria that may be located amino terminal with respect to the proline rich 20 portion of the overall amino acid sequence for such CBP.

"identity", "% identity" or The terms "percent identity" as utilized in this application refer to a calculation of differences between two contiguous sequences which have been aligned for "best 25 fit" (to provide the largest number of aligned identical corresponding sequence elements, wherein elements are either nucleotides or amino acids) and all individual differences are considered as individual difference with respect to the identity. In this respect, all individual 30 element gaps (caused by insertions and deletions with respect to an initial sequence ("reference sequence")) over the length of the reference sequence and individual substitutions of different elements (for individual 35 elements of the reference sequence) are considered as individual differences in calculating the total number of differences between two sequences. Individual differences may be compared between two sequences where an initial

sequences (reference sequence) has been varied to obtain a variant sequence (comparative sequence) or where a new sequence (comparative sequence) is simply aligned and compared to such a reference sequence. When two aligned sequences are compared all of the individual gaps in BOTH 5 sequences that are caused by the "best fit" alignment over length of the reference sequence are considered the individual differences for the purposes of identity. If an alignment exists which satisfies the stated minimum 10 identity, then a sequence has the stated minimum identity to the reference sequence. For example, the following is a comparison of two hypothetical sequences having 100 elements each that are aligned for best fit wherein one sequence is regarded as the "reference sequence" and the other as the comparative sequence. All of the individual 15 alignment gaps in both sequences are counted over the length of the reference sequence and added to the number individual element substitution of changes (aligned elements that are different) of the comparative sequence 20 for the total number of element differences. The total number of differences (for example 7 qaps and 3 substitutions) is divided by the total number of elements in the length of the reference sequence (100 elements) for "percentage difference" (10/100). the The resulting 25 percentage difference (10%) subtracted from is 100% identity to provide a "% identity" of 90% identity. For the identity calculation all individual differences in both sequences are considered in the above manner over a discrete comparison length (the length of the reference 30 sequence) of two best fit aligned sequences to determine identity. Thus, no algorithm is necessary for such an identity calculation.

"Isolated" in the context of the present invention 35 with respect to polypeptides and/or polynucleotides means that the material is removed from its original environment (e.q., the natural environment if it is naturally occurring). For example, naturally-occurring а polynucleotide or polypeptide present in a living organism

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is not isolated, but the same polynucleotide or polypeptide, separated from some or all of the co-existing materials in the natural system, is isolated. Such polynucleotides could be part of a vector and/or such polynucleotides or polypeptides could be part of а composition, and still be isolated in that such vector or composition is not part of its natural environment. The polypeptides and polynucleotides of the present invention are preferably provided in isolated an form. and preferably are purified to homogeneity.

Summary of the Invention

In one aspect the present invention relates to a vaccine for treating or preventing pneumococcal bacterial 15 infections which utilizes as an immunogen at least one polypeptide truncate of a pneumococcal surface-binding protein, analog, or variant having a highly conserved immunogenic alpha-helical portion (corresponding generally to a "consensus" amino acid sequence as set forth in SEQ 20 ID NO:1) with respect to different types of pneumococcal bacteria, which polypeptide does not include a choline-Preferably, the C-terminal cholinebinding portion. binding portion is absent from such polypeptides. More preferred are such polypeptides wherein the HPS amino acid 25 sequence is also absent. Even further preferred are polypeptides wherein the highly conserved immunogenic portion corresponding alpha-helical generally to а "consensus" amino acid sequence as set forth in SEQ ID NO:1 also corresponds generally to the amino acid sequence 30 as set forth in SEQ ID NO:19 (amino acids 1 to 103 of SEQ ID NO:19 are identical to amino acids 1 to 103 of SEQ ID Also preferred as vaccines are recombinantly-NO:1). produced, isolated polypeptides that are missing both an HPS portion and the choline-binding portion.

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More preferred as vaccines are one or more polypeptide truncates of pneumococcal surface-binding proteins, analogs or variants including a single highly conserved alpha-helix immunogenic portion with respect to

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different types of pneumococci, which polypeptides do not include a C-terminal choline-binding portion. Further preferred are isolated recombinantly produced polypeptides having such structure. Also preferred are such polypeptides that do not include either a C-terminal choline-binding portion or a HPS portion.

The present invention further provides a vaccine comprising a polypeptide including an immunogenic portion 10 that is capable of forming an alpha helix, which polypeptide includes a sequence that has at least 85% identity and preferably at least 87% identity to the amino SEQ ID NO:1, wherein the isolated acid sequence of polypeptide does not include a C-terminal choline-binding 15 portion. Further preferred are such polypeptides that comprise a polypeptide sequence that has at least 85% identity and preferably at least 87% identity to an amino acid sequence according SEQ ID NO:19. Preferably, the sequence of the isolated polypeptide includes neither an 20 HPS portion (SEQ ID NO:2) nor a C-terminal choline-binding Further preferred are isolated recombinantly portion. produced polypeptides having such structure. In particular, such polypeptides corresponding to alpha helical structures of different types of S. pneumoniae 25 bacteria are contemplated. Particularly preferred are the serotypes 1-5, 6A, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15B, 17F, 18C, 19F, 19A, 20, 22F, 23F and 33F of such S. pneumoniae bacteria. Examples of such serotypes of bacteria readily available from are standard ATCC 30 catalogs.

In additional aspect, the present an invention further provides a vaccine against S. pneumoniae comprising synthetic polypeptide а or recombinant comprising a plurality of alpha-helical portions, each derived from different naturally occurring S. pneumoniae choline-binding polypeptides wherein such alpha-helical portions have at least 85% identity to the amino acid SEQ ID NO:1, and wherein sequence of the isolated

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polypeptide does not include a choline-binding portion. Further preferred are those wherein the amino acid sequence for the alpha-helix areas is at least 85% identical to the amino acid sequence of SEQ ID NO:19.

- 5 Preferably, such synthetic polypeptide includes neither a HPS portion nor a choline-binding portion. Analogs and variants of such chain structure polypeptides wherein such alpha helical portions may be synthetic variant amino acid sequences (or may be a mixture of naturally occurring and 10 variant sequences) are also contemplated and embraced by
- the present invention. In a preferred aspect, chain vaccines polypeptides having at least ten different alpha helical structures corresponding to S. pneumoniae serotypes 1-5, 6A, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15 15B, 17F, 18C, 19F, 19A, 20, 22F, 23F and 33F are
- Further preferred are polypeptides including at provided. least fifteen of such alpha-helical structures, more preferred are polypeptides including at least 20 such alpha-helical structures and more preferred are 20 polypeptides including at least one alpha-helical structure corresponding to each of the S. pneumoniae serotypes 1-5, 6A, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14,
- 15B, 17F, 18C, 19F, 19A, 20, 22F, 23F and 33F. Another preferred polypeptide comprises each of the alpha helical 25 structures from the amino acid sequences of SEQ ID NOS:3-18 which correspond to SEQ ID NO:1.

In another aspect, the invention relates to passive immunity vaccines formulated from antibodies against a polypeptide 30 including a highly conserved immunogenic portion with respect to different types of pneumococcal bacteria which portion is capable of forming an alphahelix having the hereinbefore described identity to the amino acid sequence of SEQ ID NO:1, which polypeptide does 35 not include a C-terminal choline-binding portion, wherein said antibodies will bind to at least one S. pneumoniae species. Preferably, if such polypeptide is a truncate of a native pneumococcal surface-binding protein both its HPS portion (where applicable) and its choline-binding portion 5

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are absent from such polypeptide. Such passive immunity utilized to prevent vaccines can be and/or treat infections in immunocompromised patients, pneumococcal patients having an immature immune system (such as young children) or patients who already have an ongoing In this manner, according to a further aspect infection. of the invention, a vaccine can be produced from a recombinant polypeptide wherein the synthetic or polypeptide includes the conserved alpha helical portions of two or more different choline binding polypeptides of S. pneumoniae.

This invention also relates generally to the use of isolated polypeptide having a highly conserved an immunogenic portion with respect to different types of 15 pneumococcal bacteria which portion is capable of forming an alpha-helix (corresponding generally to SEQ ID NO:1 or to SEQ ID NO:19) wherein the isolated polypeptide does not include a choline-binding portion, to raise antibodies in non-human mammalian species useful, for 20 example, as diagnostic reagents and vaccines.

In yet another aspect, the present invention relates to the production of a polypeptide including a highly 25 conserved immunogenic portion with respect to different types of pneumococcal bacteria which portion is capable of alpha-helix forming an whose sequence corresponds generally to the amino acid sequence of SEQ ID NO:1 or SEQ NO:19, wherein the isolated polypeptide does not ID include a choline-binding portion. 30 Preferably, such recombinant production is of truncated native а pneumococcal surface-binding polypeptide wherein both the HPS portion (where applicable) and the choline-binding portion are absent.

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In still another aspect, the present invention provides an isolated choline-binding polypeptide, wherein the non-choline binding region of such polypeptide has at least 90% identity to the corresponding amino acid

sequence portion of a naturally occurring pneumococcal surface-binding protein which is a member selected from the group consisting of SEQ ID NOS:3-18. The invention relates to fragments of such polypeptides which include at least the conserved alpha-helical portion corresponding generally to SEQ ID NO:1, and which has at least 85% identity thereto, wherein the isolated polypeptide preferably is free of a choline binding region.

In another aspect the present invention provides an isolated polypeptide comprising an amino acid sequence which has at least 90% identity to one of the amino acid sequences selected from the group consisting of SEQ ID NO:3-18. Preferably, such isolated polypeptide comprises an amino acid sequence which has at least 95% identity, and more preferably 97% identity, to one of the amino acid sequences selected from the group consisting of SEQ ID NO:3-18. The invention further relates to fragments of such polypeptides.

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aspect, the present vet further invention In а provides a S. pneumoniae CBP polypeptide encoded by a polynucleotide that will hybridize under highly stringent conditions to the complement of a polynucleotide encoding 25 a polypeptide having an amino acid selected from the group consisting of SEQ ID NOS:1 and 3-18. Particularly preferred are polypeptides comprising an amino acid sequence segment that is at least 90% identical to the amino acid sequence of SEQ ID NO:1. Further preferred are 30 such polypeptides comprising a contiguous amino acid sequence that has at least 95% identity with respect to the amino acid sequence of SEQ ID NO:1. And, even more preferred are polypeptides comprising an amino acid sequence that has at least 97% identity with respect to 35 the amino acid sequence of SEQ ID NO:1.

In another aspect the present invention provides polynucleotides which encode the hereinabove described polypeptides of the invention. The polynucleotide of the

present invention may be in the form of RNA or in the form of DNA. which DNA includes cDNA, genomic DNA, and synthetic DNA. The DNA may be double-stranded or singlestranded, and if single stranded may be the coding strand or non-coding (anti-sense) strand. The polynucleotides which encode polypeptides including the amino acid sequences least one of of at SEO ID NOS:3-18 (or polypeptides that have at least 90% identity to the amino acid sequences of such polypeptides) may be one of the coding sequences shown in SEQ ID NOS:20-35 or may be of a different coding sequence which coding sequence, as a result of the redundancy or degeneracy of the genetic code, encodes the same polypeptides as the DNA of SEQ ID NOS:20-35.

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The polynucleotides which encode the polypeptides of SEQ ID NOS:3-18 may include: only the coding sequence for the polypeptide; the coding sequence for the polypeptide (and optionally additional coding sequence) and non-coding 20 sequence, such as introns or non-coding sequence 5' and/or 3' of the coding sequence for the polypeptide. The polypeptides encoded may comprise just a single alphahelical portion or multiple alpha-helical portion and may independently or collectively include N-terminal sequences 25 5' such helical of alpha areas and/or sequences corresponding to the "X" structures or proline rich areas (as set forth in Figure 1, for example).

The invention further relates to a polynucleotide 30 comprising a polynucleotide sequence that has at least 95% identity and preferably at least 97% identity to a polynucleotide encoding one of the polypeptides comprising SEO NO:3-18. The invention further relates to ID fragments of such polynucleotides which include at least 35 the portion of the polynucleotide encoding the polypeptide sequence corresponding to SEQ ID NO:1.

Thus, the term "polynucleotide encoding a polypeptide" encompasses a polynucleotide which includes

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only coding sequence for the polypeptide as well as a polynucleotide which includes additional coding and/or non-coding sequence. In particular, the polypeptides may include any or all of the types of structures set forth schematically in Figure 1.

The present invention further relates to variants of the hereinabove described polynucleotides which encode for fragments, analogs and derivatives of the polypeptides including the amino acid sequences of SEQ ID NOS:3-18. The variants of the polynucleotides may be a naturally occurring allelic variant of the polynucleotides or a nonnaturally occurring variant of the polynucleotides. Complements to such coding polynucleotides may be utilized to isolate polynucleotides encoding the same or similar polypeptides. In particular, such procedures are useful to obtain alpha helical coding segments from different serotypes of *S. pneumoniae*, which is especially useful in

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polypeptide.

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Thus, the present invention includes polynucleotides encoding polypeptides including the same polypeptides as shown in the Sequence Listing as SEQ ID NOS:3-18 as well as variants of such polynucleotides which variants encode for a fragment, derivative or analog of the polypeptides of SEQ ID NOS:3-18. Such nucleotide variants include deletion variants, substitution variants and addition or insertion variants.

the production of "chain" polypeptide vaccines containing

multiple alpha helical segments.

30 As hereinabove indicated, the polynucleotides may have a coding sequence which is a naturally occurring allelic variant of the coding sequence shown in the Sequence Listing as SEQ ID NOS:20-35. As known in the art, an allelic variant is an alternate form of a 35 polynucleotide sequence which may have a substitution, deletion or addition of one or more nucleotides, which does not substantially alter the function of the encoded

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The polynucleotides of the present invention may also have the coding sequence fused in frame to a marker sequence which allows for purification of the polypeptides of the present invention. The marker sequence may be, for example, a hexa-histidine tag supplied by a pQE-9 vector to provide for purification of the mature polypeptides fused to the marker in the case of a bacterial host, or, for example, the marker sequence may be a hemagglutinin (HA) tag when a mammalian host, e.g. COS-7 cells, is used. The HA tag corresponds to an epitope derived from the influenza hemagglutinin protein (Wilson, I., et al., Cell, 37:767 (1984)).

The present invention further relates to polynucleotides (hybridization target sequences) 15 which hybridize to the complements of the hereinabove-described sequences if there is at least 70% and preferably 80% identity between the target sequence and the complement of the sequence to which the target sequence hybridizes, preferably at least 85% identity. More preferred are such 20 sequences having at least 90% identity, preferably at least 95% and more preferably at least 97% identity between the target sequence and the sequence of complement of the polynucleotide to which it hybridizes. The 25 invention further relates to the complements to both the target sequence and to the polynucleotide sequence that encodes an amino acid sequence selected from the group consisting of SEQ ID NOS:3 to 18. The present invention particularly relates to polynucleotides which hybridize under stringent conditions to the complements of the 30 hereinabove-described polynucleotides as well as to those herein used, complements. As the term "stringent conditions" means hybridization will occur with the complement of polynucleotide and corresponding a а sequence only if there is at least 95% and preferably at 35 least 97% identity between the target sequence and the sequence of complement of the polynucleotide to which it The polynucleotides which hybridize to the hybridizes. complements of the hereinabove described polynucleotides

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in a preferred embodiment encode polypeptides which retain an immunogenic portion that will cross-react with an antibody to at least one of the polypeptides having a sequence according to SEQ ID NOS:3-18, or to a polypeptide that includes an amino acid sequence which has at least 85% identity to that of SEQ ID NO:1.

In a still further aspect, the present invention provides for the production of such polypeptides and vaccines as set forth above having a histidine label (or other suitable label) such that the full-length proteins, truncates, analogs or variant discussed above can be isolated due to their label.

In another aspect the present invention relates to a method of prophylaxis and/or treatment of diseases that are mediated by pneumococcal bacteria that have surface-binding CBP proteins. In particular, the invention relates to a method for the prophylaxis and/or treatment of infectious diseases that are mediated by *S. pneumoniae* that have a CBP surface-binding protein that forms an alpha helix (comprising a sequence that has at least an 85% identity to the amino acid sequence of SEQ ID NO:1). In a still further preferred aspect, the invention relates

25 to a method for the prophylaxis and/or treatment of such infections in humans.

still another aspect the present In invention relates to a method of using one or more antibodies 30 (monoclonal, polyclonal or sera) to the polypeptides of the invention as described above for the prophylaxis and/or treatment of diseases that are mediated bv pneumococcal bacteria that have CBP surface-binding In particular, the invention relates to a proteins. 35 method for the prophylaxis and/or treatment of infectious diseases that are mediated by S. pneumoniae CBP proteins which include an alpha helical portion having the hereinbefore described identity to the consensus sequence of SEQ ID NO:1. In a still further preferred aspect, the 5

invention relates to a method for the prophylaxis and/or treatment of otitis media, nasopharyngeal, bronchial infections, and the like in humans by utilizing antibodies to the alpha-helix containing immunogenic polypeptides of the invention as described above.

Brief Description of Drawings

Figure 1 is a diagram of a pneumococcal CBP protein to the C-terminal, which shows from the N-terminal 5 respectively, (a) a N-terminal sequence, (b) one of a potential alpha-helical forming area conserved segment (R1) that may not be present in some CBP polypeptides, (c) an optional small bridging sequence of amino acids that may bridge two conserved alpha-helical segments (X), (d) a 10 second of a potential alpha-helical forming area consensus sequence (R2) related to the first consensus sequence (which corresponds to SEQ ID NO:1), (e) a proline rich area sequence, (f) a choline binding repeats area, and (e) a C-terminal tail sequence. Where relevant, an 15 optional HPS sequence may naturally occur 5' of the proline rich sequence and 3' of the R1, X, and/or R2 areas.

Figure 2 reports the results for passive immunity 20 protection against 1600 cfu virulent serotype 6B S. pneumoniae SP317 (in mice) that was provided by day 31 rabbit antisera to pneumococcal CBP а truncate polypeptide, NR1XR2 (truncate missing both the proline and the choline binding areas, but including two conserved alpha-helical areas R1 and R2). 25 Eighty percent of the mice immunized with the truncate antisera prior to challenge survived the 14 day observation period. By contrast, all mice immunized with a control sera (preimmune rabbit sera) were dead by day 7.

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Figure 3 reports the results for passive immunity protection aqainst virulent 3450 cfu serotype 6B S. pneumoniae SP317 (in mice) that was provided by day 52 rabbit antisera to a pneumococcal CBP truncate polypeptide, NR1XR2 (truncate missing both the proline and the choline binding areas, but including two conserved alpha-helical areas R1 and R2). One hundred percent of the mice immunized with the truncate antisera prior to challenge survived the 10 day observation period. By

contrast, ninety percent of the mice immunized with a control sera (pre-immune rabbit sera) were dead at day 10.

Figure 4 reports the results for passive immunity protection against 580 cfu virulent 5 serotype 6B S. pneumoniae SPSJ2 (in mice) that was provided by day 31 to pneumococcal rabbit antisera a CBP truncate polypeptide, NR1XR2 (truncate missing both the proline and the choline binding areas, but including two conserved alpha-helical areas R1 and R2). Fifty percent of the mice 10 immunized with the truncate antisera prior to challenge survived the 10 day observation period. By contrast, all mice immunized with a control sera (pre-immune rabbit sera) were dead by day 8.

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Figure 5 reports the results for active immunity protection against 560 cfu virulent serotype 6B S. pneumoniae SPSJ2 (in mice) that was provided by immunization with a pneumococcal CBP truncate polypeptide, 20 NR1X (truncate missing the second conserved alpha-helical area R2, as well as both the proline and the choline binding areas). Eighty percent of the mice actively immunized with the NR1X CBP truncate prior to challenge survived the 14 day observation period. By contrast, all 25 mice immunized with a control (sham mice) of PBS and adjuvant were dead by day 8.

Figure 6 reports the results for active immunity protection 680 cfu virulent against serotype 6B 30 S. pneumoniae SPSJ2 (in mice) that was provided by immunization with a pneumococcal CBP truncate polypeptide, NR1XR2 (truncate missing both the proline and the choline binding areas, but including two conserved alpha-helical areas R1 and R2). Fifty percent of the mice actively 35 immunized with the NR1XR2 CBP truncate prior to challenge survived the 14 day observation period. By contrast, all mice immunized with a control (SP90) protein and adjuvant were dead by day 9.

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Figure 7 is an alignment report of the amino terminus of CBP polypeptides from various types of *S*. *pneumoniae* and a consensus sequence is reported at the top of each row (sets of lines) of the comparison. The consensus sequence for the comparison is listed as the "Majority" sequence (SEQ ID NO:36). One letter codes are utilized to represent the sequences which are aligned for a "best fit" comparison wherein dashes in a sequence indicate spacing gaps of the contiguous sequence.

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Figure 8 shows the sequence pair distances for the amino acid sequences as described for Figure 7 and set forth therein. A Clustal method with identity residue weight table is used. The percent similarity for such a comparison is reported for the amino acid sequences set forth in Figure 7.

Figure 9 is an alignment report for a first helical region in the amino acid sequences of CBP polypeptides 20 from various types of *S. pneumoniae* and a consensus sequence is reported at the top of each row (sets of lines) of the comparison. The consensus sequence for the comparison is listed as the "Majority" sequence (SEQ ID NO:38). One letter codes are utilized to represent the 25 sequences which are aligned for a "best fit" comparison wherein dashes in a sequence indicate spacing gaps of the contiguous sequence.

Figure 10 shows the sequence pair distances for the 30 amino acid sequences as described for Figure 9 and set forth therein. A Clustal method with identity residue weight table is used. The percent similarity for such a comparison is reported for the amino acid sequences set forth in Figure 9.

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Figure 11 is an alignment report for the region X in the amino acid sequences of CBP polypeptides from various types of *S. pneumoniae* and a consensus sequence is reported at the top of each row (sets of lines) of the

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comparison. The consensus sequence for the comparison is listed as the "Majority" sequence (SEQ ID NO:37). One letter codes are utilized to represent the sequences which are aligned for a "best fit" comparison wherein dashes in a sequence indicate spacing gaps of the contiguous sequence.

Figure 12 shows the sequence pair distances for the amino acid sequences as described for Figure 11 and set 10 forth therein. A Clustal method with identity residue weight table is used. The percent similarity for such a comparison is reported for the amino acid sequences set forth in Figure 11.

15 Figure 13 is an alignment report for the second helical region A in the amino acid sequences of CBP polypeptides from various types of S. pneumoniae and a consensus sequence is reported at the top of each row (sets of lines) of the comparison. The consensus sequence for the comparison is listed as the "Majority" sequence 20 (SEQ ID NO:1). One letter codes are utilized to represent sequences which are aligned the for a "best fit" comparison wherein dashes in a sequence indicate spacing gaps of the contiguous sequence.

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Figure 14 shows the sequence pair distances for the amino acid sequences as described for Figure 13 and set forth therein. A Clustal method with identity residue weight table is used. The percent similarity for such a comparison is reported for the amino acid sequences set forth in Figure 13.

Figure 15 is an alignment report for the second helical region B in the amino acid sequences of CBP 35 polypeptides from various types of *S. pneumoniae* and a consensus sequence is reported at the top of each row (sets of lines) of the comparison. The consensus sequence for the comparison is listed as the "Majority" sequence (SEQ ID NO:19). One letter codes are utilized to

represent the sequences which are aligned for a "best fit" comparison wherein dashes in a sequence indicate spacing gaps of the contiguous sequence.

5 Figure 16 shows the sequence pair distances for the amino acid sequences as described for Figure 15 and set forth therein. A Clustal method with identity residue weight table is used. The percent similarity for such a comparison is reported for the amino acid sequences set 10 forth in Figure 15.

Detailed Description of the Invention

- 15 In accordance with an aspect of the present invention there is provided a vaccine to produce a protective response against *S. pneumoniae* infections which employs a polypeptide which comprises a member selected from the group consisting of:
- 20 (a) an amino acid sequence which produces an alpha helical structure and which is at least 85% identical to the amino acid sequence of SEQ ID NO:1 and which is free of a choline binding region, and
- (b) an isolated truncate of a naturally occurring
 25 S. pneumoniae polypeptide that comprises an alpha helical portion that has at least 85% identity to the amino acid sequence of SEQ ID NO:1 and is free of a choline binding region,
- (C) an isolated truncate of a naturally occurring 30 S. pneumoniae polypeptide that comprises an alpha helical portion that has at least 90% identity to the amino acid sequence of SEQ ID NO:19 and is free of a choline binding region. In a preferred aspect, such isolated truncate polypeptide is a member selected from the group consisting 35 of SEQ ID NOS:3-18 and said isolated polypeptide is free of a choline binding region and, if relevant, a HPS region; or a fragment thereof which includes at least the alpha helical segment which corresponds to the consensus sequence of SEQ ID NO:1. Particularly preferred are

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vaccines which utilize such truncate polypeptides that include at least such alpha helical area or utilize a recombinant immunogen polypeptide comprising at least two of such alpha-helical segments. Such polypeptide may be recombinant polypeptide containing multiple alphaa areas from one or helical more trucates. Further preferred are recombinant immunogen polvpeptides comprising at least two alpha-helical areas corresponding to the alpha helical areas of two or more truncates from pneumococcal different types of bacteria. Such polypeptide may be a recombinant polypeptide containing multiple alpha-helical areas from one or more different types of pneumococcal bacteria.

- In accordance with the present invention, there is provided an isolated polypeptide comprising a truncated surface-binding polypeptide derived from S. pneumoniae, said isolated polypeptide containing an alpha-helical area whose amino acid sequence corresponds generally to the amino acid sequence of SEQ ID NO:1, but free of a choline binding area. Preferably, said isolated polypeptide also omits any naturally occurring repeats of the alpha-helical forming area and omits any HPS amino acid sequence that may be present.
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It is an object of the present invention to utilize as immunogenic composition for a vaccine (or to produce antibodies for use as a diagnostic or as a passive vaccine) comprising an immunogenic polypeptide comprising a pneumococcal surface-binding polypeptide with an alpha helical portion from which a choline binding region has been omitted. In one embodiment, such truncated proteins (naturally or recombiantly produced, as well as functional analogs) from *S. pneumoniae* bacteria are contemplated.

35 Even more particularly, *S. pneumoniae* polypeptides having a single alpha helical portion that omit any HPS areas that occur and choline binding areas of the native protein are contemplated.

A particularly preferred embodiment of such an immunogenic composition is for use as a vaccine (or as an immunogen for producing antibodies useful for diagnostics vaccines) wherein the active component of or the immunogenic composition is an isolated polypeptide comprising at least one member selected from the group consisting of:

(a) an amino acid sequence which is selected from SEQ ID NOS:3-19,

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(b) a polypeptide which has at least 90% identity to (a), preferably at least 95% identity to (a), and even more preferred at least 97% identity to (a), or

a fragment of (a) or (b) wherein such fragment (C) includes at least one alpha helical portion that corresponds to the consensus sequence which is SEQ ID NO:1 15 and said fragment does not comprise a choline binding region. Preferably, such vaccines utilize a polypeptide that contains neither a choline binding region nor an HPS region that occurs as part of the amino acid sequences in the native proteins. 20

In another preferred embodiment, there is provided a vaccine which includes at least one isolated polypeptide which includes an amino acid sequence which has at least 25 85% identity (preferably 87% identity and more preferably at least 90% identity) to SEQ ID NO:1, which isolated polypeptide is free of a choline binding portion and, where applicable, is also preferably free of an HPS The preferred polypeptide may also include one portion. 30 or more of the N-terminal sequences that are located 5' of the alpha helical areas in the polypeptides having an amino acid sequence selected from the group consisting of SEQ ID NOS:3-18, or the like. The polypeptide truncate may also include one or more of the proline regions (region "P" in Figure 1) and/or the spanning region 35 (region "X" in Figure 1).

In another aspect of the invention, such an immunogenic composition may be utilized to produce

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antibodies to diagnose pneumococcal infections, or to produce vaccines for prophylaxis and/or treatment of such pneumococcal infections as well as booster vaccines to maintain a high titer of antibodies against the immunogen(s) of the immunogenic composition.

antigens have been contemplated While other to produce antibodies for diagnosis and for the prophylaxis and/or treatment of pneumococcal infections, there is a 10 need for improved or more efficient vaccines. Such vaccines should have an improved or enhanced effect in preventing bacterial infections mediated pneumococci having surface-binding polypeptides. Further, to avoid unnecessary expense and provide broad protection against a 15 range of pneumococcal serotypes there is a need for polypeptides that comprise an immunogenic amino acid sequence corresponding to a portion of pneumococcal surface-binding polypeptides that is a highly conserved portion among various types of pneumococci. Preferably, such polypeptides avoid the inclusion of 20 amino acid sequences corresponding to other portions of the native polypeptides, such as the choline binding region and/or the HPS region.

25 There is a need for improved antigenic compositions comprising highly conserved portions of polypeptides that bind to the surface of pneumococcal bacteria for stimulating high-titer specific antisera to provide protection against infection by pathogenic pneumococcal bacteria and also for use as diagnostic reagents. 30

In such respect, truncated polypeptides, functional variant analogs, and recombinantly produced truncated polypeptides of the invention are useful as immunogens for 35 preparing vaccine compositions that stimulate the production of antibodies that can confer immunity against pathogenic species of bacteria. Further, preparation of vaccines containing purified proteins as antigenic ingredients are well known in the art.

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Generally, vaccines are prepared as injectables, in the form of aqueous solutions or suspensions. Vaccines in an oil base are also well known such as for inhaling. Solid forms which are dissolved or suspended prior to use 5 Pharmaceutical carriers are may also be formulated. generally added that are compatible with the active ingredients and acceptable for pharmaceutical use. Examples of such carriers include, but are not limited to, 10 water. saline solutions, dextrose, or glycerol. Combinations of carriers may also be used.

Vaccine compositions may further incorporate additional substances to stabilize pH, or to function as 15 adjuvants, wetting agents. or emulsifying agents, which can serve to improve the effectiveness of the vaccine.

Vaccines are generally formulated for parenteral administration and are injected either subcutaneously or 20 intramuscularly. Such vaccines can also be formulated as suppositories or for oral administration, using methods known in the art.

The amount of vaccine sufficient to confer immunity to pathogenic bacteria is determined by methods well known 25 to those skilled in the art. This quantity will be determined based upon the characteristics of the vaccine Typically, recipient and the level of immunity required. amount of vaccine to be administered will be the determined based upon the judgment of a skilled physician. 30 Where vaccines are administered by subcutaneous or intramuscular injection, a range of 50 to 500 μ g purified protein may be given.

35 The term "patient in need thereof" refers to a human that is infected with, or likely, to be infected with, pathogenic pneumococcal bacteria that produce CbpA, or the like, preferably *S. pneumoniae* bacteria (however a mouse

model can be utilized to simulate such a patient in some circumstances).

In addition to use as vaccines, the polypeptides of 5 the present invention can be used as immunogens to stimulate the production of antibodies for use in passive immunotherapy, for use as diagnostic reagents, and for use reagents in other processes such as affinity as chromatography.

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The polynucleotides encoding the immunogenic polypeptides described above may also have the coding sequence fused in frame to a marker sequence which allows for purification of the polypeptides of the present The marker sequence may be, for example, a 15 invention. hexa-histidine tag supplied by a pQE-9 vector to provide for purification of the mature polypeptides fused to the marker in the case of a bacterial host, or, for example, the marker sequence may be a hemagglutinin (HA) tag when a 20 mammalian host, e.g. COS-7 cells, is used. The HA tag corresponds to an epitope derived from the influenza hemagglutinin protein (Wilson, I., et al., Cell, 37:767 (1984)).

25 The identification of multiple coil structures of alpha helical amino acid segments in the S. pneumoniae polypeptides according to the invention may be determined by the location of proline rich areas with respect to one another. Further the "X" area optionally located between two or more alpha-helical structures can play a part in 30 the formation of a coil within a coil structure. Standard three-dimensional protein modeling may be utilized for determining the relative shape of such structures. An example of a computer program, the Paircoil Scoring Form 35 Program ("PairCoil program"), useful for such threedimensional protein modelling is described by Berger et al. in the Proc. Natl. Acad. of Sci. (USA), 92:8259-8263 (August 1995). The PairCoil program is described as a computer program that utilizes a mathematical algorithm to

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predict locations of coiled-coil regions in amino acid sequences. A further example of such a computer program is described by Wolf et al., Protein Science 6:1179-1189 (June 1997) which is called the Multicoil Scoring Form Program ("Multicoil program"). The MultiCoil program is based on the PairCoil algorithm and is useful for locating dimeric and trimeric coiled coils .

In a preferred aspect, the invention provides for 10 recombinant production of such polypeptides in a host bacterial cell other than a *S. pneumoniae* species host to avoid the inclusion of native surface-binding polypeptides that have a choline binding region. A preferred host is a species of such bacteria that can be cultured under 15 conditions such that the polypeptide of the invention is secreted from the cell.

The present invention also relates to vectors which include polynucleotides encoding one or more of the 20 polypeptides of the invention that include the highly conserved alpha-helical amino acid sequence in the absence of an area encoding a choline binding amino acid sequence, host cells which are genetically engineered with vectors of the invention and the production of such immunogenic 25 polypeptides by recombinant techniques in an isolated and substantially immunogenically pure form.

Host cells are genetically engineered (transduced or transformed or transfected) with the vectors comprising a polynucleotide encoding a polypeptide comprising the 30 highly conserved alpha-helical region but not having a choline binding region, or the like of this invention which may be, for example, a cloning vector or an expression vector. The vector may be, for example, in the 35 form of a plasmid, a viral particle, a phage, etc. The engineered host cells can be cultured in conventional nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the polynucleotides which encode such polypeptides. The

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culture conditions, such as temperature, pH and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

include chromosomal, nonchromosomal and Vectors synthetic DNA sequences, e.g., derivatives of SV40; bacterial plasmids; phage DNA; baculovirus; yeast plasmids; vectors derived from combinations of plasmids and phage DNA, viral DNA such as vaccinia, adenovirus, 10 fowl pox virus, and pseudorabies. However, any other vector may be used as long as it is replicable and viable in the host.

15 The appropriate DNA sequence may be inserted into the vector by a variety of procedures. In general, the DNA sequence is inserted into an appropriate restriction endonuclease site(s) by procedures known in the art. Such procedures and others are deemed to be within the scope of 20 those skilled in the art.

sequence in the expression vector The DNA is operatively linked to an appropriate expression control (promoter) to direct mRNA synthesis. sequence(s) As 25 representative examples of such promoters, there may be mentioned: LTR or SV40 promoter, the E. coli. lac or trp, the phage lambda $P_{\rm\scriptscriptstyle L}$ promoter and other promoters known to control expression of genes in prokaryotic or eukaryotic cells or their viruses. The expression vector also ribosome binding site 30 contains а for translation initiation and a transcription terminator. The vector may also include appropriate sequences for amplifying expression.

35 In addition, the expression vectors preferably contain one or more selectable marker genes to provide a phenotypic trait for selection of transformed host cells such as dihydrofolate reductase or neomycin resistance for eukaryotic cell culture, or such as tetracycline or ampi-

cillin resistance in E. coli.

The vector containing the appropriate DNA sequence as hereinabove described, as well as an appropriate 5 promoter or control sequence, may be employed to transform an appropriate host to permit the host to express the proteins.

As representative examples of appropriate hosts, 10 there may be mentioned: bacterial cells, such as <u>E. coli</u>, <u>Streptomyces</u>, <u>Salmonella typhimurium</u>; fungal cells, such as yeast; insect cells such as <u>Drosophila S2</u> and <u>Spodoptera Sf9</u>; animal cells such as CHO, COS or Bowes melanoma; adenoviruses; plant cells, etc. The selection 15 of an appropriate host is deemed to be within the scope of those skilled in the art from the teachings herein.

More particularly, the present invention also includes recombinant constructs comprising one or more of 20 the sequences as broadly described above. The constructs comprise a vector, such as a plasmid or viral vector, into which a sequence of the invention has been inserted, in a forward or reverse orientation. In a preferred aspect of this embodiment. the further construct comprises 25 regulatory sequences, including, for example, a promoter, operably linked to the sequence. Large numbers of suitable vectors and promoters are known to those of skill in the art, and are commercially available. The following vectors are provided by way of example. Bacterial: pQE70, 30 pQE60, pQE-9 (Qiagen, Inc.), pbs, pD10, phagescript, psiX174, pbluescript SK, pbsks, pNH8A, pNH16a, pNH18A, pNH46A (Stratagene); ptrc99a, pKK223-3, pKK233-3, pDR540, pRIT5 (Pharmacia). Eukaryotic: pWLNEO, pSV2CAT, pOG44, (Stratagene) pXT1, pSG pSVK3, pBPV, pMSG, pSVL 35 (Pharmacia). However, any other plasmid or vector may be used as long as they are replicable and viable in the host.

Promoter regions can be selected from any desired

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gene using CAT (chloramphenicol transferase) vectors or other vectors with selectable markers. Two appropriate vectors are pKK232-8 and pCM7. Particular named bacterial promoters include lacI, lacZ, T3, T7, gpt, lambda P_R , P_L and TRP. Eukaryotic promoters include CMV immediate early, HSV thymidine kinase, early and late SV40, LTRs from retrovirus, and mouse metallothionein-I. Selection of the appropriate vector and promoter is well within the level of ordinary skill in the art.

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a further embodiment, the present invention In relates to host cells containing the above-described The host cell can be a higher eukaryotic constructs. cell, such as a mammalian cell, or a lower eukaryotic cell, such as a yeast cell, or the host cell can be a 15 prokaryotic cell, such as a bacterial cell. Introduction of the construct into the host cell can be effected by calcium phosphate transfection, DEAE-Dextran mediated transfection, or electroporation (Davis, L., Dibner, M., 20 Battey, I., Basic Methods in Molecular Biology, (1986)).

The constructs in host cells can be used in a conventional manner to produce the gene product encoded by the recombinant sequence. Alternatively, the polypeptides of the invention can be synthetically produced by conventional peptide synthesizers.

Mature proteins can be expressed in mammalian cells, yeast, bacteria, or other cells under the control of appropriate promoters. Cell-free translation systems can 30 also be employed to produce such proteins using RNAs derived from the DNA constructs of the present invention. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are described by Sambrook, et al., Molecular Cloning: A Laboratory Manual, 35 Second Edition, Cold Spring Harbor, N.Y., (1989), the disclosure of which is hereby incorporated by reference.

Transcription of the DNA encoding the polypeptides

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of the present invention by higher eukaryotes is increased by inserting an enhancer sequence into the vector. Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp that act on a promoter to increase its transcription. Examples including the SV40 enhancer on the late side of the replication origin bp 100 to 270, a cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers.

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Generally, recombinant expression vectors will include origins of replication and selectable markers permitting transformation of the host cell, e.g., the ampicillin resistance gene of E. coli and S. cerevisiae TRP1 gene, and a promoter derived from a highly-expressed gene to direct transcription of a downstream structural Such promoters can be derived from operons sequence. encoding glycolytic enzymes such as 3-phosphoglycerate kinase (PGK), α -factor, acid phosphatase, or heat shock 20 proteins, among others. The heterologous structural sequence is assembled in appropriate phase with translation initiation and termination sequences. Optionally, the heterologous sequence can encode a fusion protein including an N-terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product.

Useful expression vectors for bacterial use are constructed by inserting а structural DNA sequence 30 a desired together encoding protein with suitable translation initiation and termination signals in operable reading phase with a functional promoter. The vector will comprise one or more phenotypic selectable markers and an origin of replication to ensure maintenance of the vector 35 and to, if desirable, provide amplification within the host. Suitable prokaryotic hosts for transformation include E. coli, Bacillus subtilis, Salmonella typhimurium and various species within the genera Pseudomonas,

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Streptomyces, and Staphylococcus, although others may also be employed as a matter of choice.

As a representative but nonlimiting example, useful expression vectors for bacterial use can 5 comprise a selectable marker and bacterial origin of replication derived from commercially available plasmids comprising genetic elements of the well known cloning vector pBR322 (ATCC 37017). Such commercial vectors include. for 10 example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and GEM1 (Promega Biotec, Madison, WI, USA). These pBR322 "backbone" sections are combined with an appropriate promoter and the structural sequence to be expressed.

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Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period.

Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification.

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Microbial cells employed in expression of proteins can be disrupted by any convenient method, including freeze-thaw cycling, sonication, a french press, mechanical disruption, or use of cell lysing agents, such methods are well know to those skilled in the art. However, preferred are host cells which secrete the polypeptide of the invention and permit recovery of the

35 Various mammalian cell culture systems can also be employed to express recombinant protein. Examples of mammalian expression systems include the COS-7 lines of monkey kidney fibroblasts, described by Gluzman, Cell, 23:175 (1981), and other cell lines capable of expressing

polypeptide from the culture media.

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a compatible vector, for example, the C127, 3T3, CHO, HeLa and BHK cell lines. Mammalian expression vectors will comprise an origin of replication, a suitable promoter and enhancer, and also any necessary ribosome binding sites, polyadenylation site, splice donor and acceptor sites, transcriptional termination sequences, and 5' flanking nontranscribed sequences. DNA sequences derived from the SV40 splice, and polyadenylation sites may be used to provide the required nontranscribed genetic elements.

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The polypeptides can be recovered and/or purified from recombinant cell cultures by well-known protein recovery and purification methods. Such methodology may include ammonium sulfate or ethanol precipitation, acid chromatography, 15 extraction, anion or cation exchange phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxylapatite lectin chromatography. chromatography and Protein refolding steps can be used, as necessary, in completing 20 configuration of the mature protein. In this respect, chaperones may be used in such a refolding procedure. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps.

25 The polypeptides that are useful as immunogens in the present invention may be a naturally purified product, or a product of chemical synthetic procedures, or produced by recombinant techniques from a prokaryotic or eukaryotic host (for example, by bacterial, yeast, higher plant, 30 insect and mammalian cells in culture). Depending upon the host employed in a recombinant production procedure, polypeptides of the present invention the may be glycosylated or may be non-glycosylated. Particularly preferred immunogens are truncated pneumococcal 35 polypeptides that comprise a single highly conserved alpha helical area, but do not comprise a choline binding region or a HPS region. Therefore, antibodies against such polypeptides should bind to other pneumococcal bacterial species (in addition to the S. pneumoniae species from

which such polypeptides were derived) and vaccines against such *S. pneumoniae* should give protection against other pneumococcal bacterial infections.

5 Procedures for the isolation of the individually expressed alpha-helical containing polypeptides may be isolated by recombinant expression/isolation methods that are well-known in the art. Typical examples for such isolation may utilize an antibody to a conserved area of 10 the protein or to a His tag or cleavable leader or tail that is expressing as part of the protein structure.

The polypeptides, their fragments or other derivatives, or analogs thereof, or cells expressing them 15 can be used as an immunogen to produce antibodies thereto. These antibodies can be, for example, polyclonal or antibodies. The monoclonal present invention also includes chimeric, single chain, and humanized antibodies, as Fab fragments, or the product of as well an Fab 20 expression library. Various procedures known in the art may be used for the production of such antibodies and fragments.

Antibodies generated against the polypeptides 25 corresponding to a sequence of the present invention can be obtained by direct injection of the polypeptides into an animal or by administering the polypeptides to an animal, preferably a nonhuman. The antibody so obtained will then bind the polypeptides itself. In this manner, 30 sequence encoding only even а a fragment of the polypeptides can be used to generate antibodies binding the whole native polypeptides.

For preparation of monoclonal antibodies, any 35 technique which provides antibodies produced by continuous cell line cultures can be used. Examples include the hybridoma technique (Kohler and Milstein, 1975, Nature, 256:495-497), the trioma technique, the human B-cell hybridoma technique (Kozbor et al., 1983, Immunology Today

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4:72), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole, et al., 1985, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96).

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Techniques described for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce single chain antibodies to immunogenic polypeptide products of this invention. Also, transgenic mice may be used to express humanized antibodies to immunogenic polypeptide products of this invention.

In order to facilitate understanding of the above description and the examples which follow below, as well as the Figures included herewith, Table 1 below sets forth the bacterial source for the polypeptides of SEQ ID NOS:3-18 and the polynucleotides encoding them (SEQ ID NOS:20-35, respectively). The name and/or type of bacteria is specified and a credit or source is named. The sequences 20 from such types of bacteria are for illustrative purposes only since by utilizing probes and/or primers as described herein other sequences of similar type may be readily obtained by utilizing only routine skill in the art.

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TABLE 1

SEQ ID NO.	Type Of Pneumococcus	Source Credit or ATCC No.
3	1	ATCC 33400
4	2	SPATCC 11733
5	2	ATCC2 (catalog #6302)
6	4	ATCC4 (catalog #6304)
7	6B	ATCC 6B (catalog #6326)
8	18C	SPATCC 18C (ATCC catalog #10356)

SEQ ID NO.	Type Of Pneumococcus	Source Credit or ATCC No.
9	4	Norway type 4; Nat'l. Inst. of Public Health, Norway Ingeborg Aagerge
10	noncapsulated	R6X; Rockefeller Univ., Rob Masure (from D39, type 2)
11	6B	SPB 105; Boston Univ., Steve Pelton
12	23F	SPB 328; Boston Univ., Steve Pelton
13	14	SPB 331; Boston Univ., Steve Pelton
14	23F	SPB 365; Boston Univ., Steve Pelton
15	9V	SPR 332; Rockefeller Univ., Rob Masure
16	68	SPSJ 2p; St. Jude Children's Research Hospital, Pat Flynn (clinical isolate passaged 1x in mice for virulence)
17	14	SPSJ 9; St. Jude Children's Research Hospital, Pat Flynn (clinical isolate - nares, pneumonia)
18	19A	SPSJ 12; St. Jude Children's Research Hospital, Pat Flynn (clinical isolate)

TABLE 1 (Continued)

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The present invention will be further described with reference to the following examples; however, it is to be understood that the present invention is not limited to 10 such examples. All parts or amounts, unless otherwise specified, are by weight.

Example 1

Generation of CbpA Truncate Protein Vectors

A. Vector for Full Length CbpA (NR1XR2PC)

5 A virulent serotype 4 S. pneumoniae strain, Norway 4 (obtained from I. Aaberge, National Instute of Public Health, Oslo, Norway) was used as a source of genomic DNA template for amplifying the polynucleotide encoding fulllength CbPA. Full length CbpA was amplified with PCR 10 primers SJ533 and SJ537 described below.

The degenerate forward primer SJ533 was designed based on the CbpA N-terminal sequence XENEG provided by H.R. Masure (St. Jude Childern's Research Hospital, 15 Memphis, TN). The SJ533 primer = 5' GGC GGA TCC ATG GA(A,G) AA(C,T) GA(A,G) GG 3'. It incorporates both BamHI and NcoI restriction sites and an ATG start codon.

20 The 3' reverse primer SJ537 = 5' GCC GTC GAC TTA GTT TAC CCA TTC ACC ATT GGC 3'

This primer incorporates a SalI restriction site for cloning purposes, and the natural stop codon from CbpA, and is based on type 4 and R6X sequence generated in-25 house.

PCR product generated from genomic DNA template with SJ533 and SJ537 was digested with BamHI and SalI, and cloned into the pQE30 expression vector (Qiagen, Inc.) 30 digested with BamHI, XbaI, and SmaI. The type R6X template resulted in full-length vector PMI581 and the type 4 template DNA resulted in full-length vector PMI580.

35 B. Vector for CbpA Truncate Protein (NR1XR2) The naturally occurring PvuII site in the end on the second amino repeat (nucleic acid 1228 of Type 4 sequence) was exploited to create a truncated version of

CbpA, containing only the amino terminus of the gene. то create the truncate clone, the full-length clone PMI580 (Type 4) or PMI581 (R6X) was digested with PvuII and XbaI, and the amino terminus along with a portion of the 5 expression vector was isolated by size on an agarose gel. pQE30 was digested with XbaI and SmaI, and the band corresponding to the other half of the vector was also size selected on an agarose gel. The two halves were ligated and clones identified by restriction digest, then 10 expressed. In this instance, the stop codon utilized is in the expression vector, so the protein expressed is larger than the predicted size due to additional amino acids at the 5' and 3' end of the cloning site.

15 C. Vector for CbpA Truncate Protein (NR1X)

A similar strategy was used to express only the first amino repeat of CbpA. Here the naturally occurring XmnI site between the two amino repeats (nucleic acid 856 of Type 4 sequence) was utilized. CbpA full-length clone 20 PMI580 was digested with XmnI and AatII. Expression vector pQE30 was digested with AatII and SmaI. Once again, the two sized fragments were ligated, and clones were screened by restriction digest and expressed.

Example 2

Expression of CbpA Truncate Protein From Expression Vectors

All proteins are expressed from the vectors 30 described in Examples 1A-1C in the Qia expressionist System (Qiagen) using the *E. coli* expression vector pQE30, and the amino terminus His6 tagged proteins are detected by western analysis using both anti-Histidine antibodies and gene specific antibodies.

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The expressed CBP truncates were purified as follows.

A single colony was selected from plated bacteria

containing the recombinant plasmid and grown overnight at 37° in 6.0 ml LB buffer with 50 ug/ml Kanamycin and 100 ug/ml Ampicillin. This 6.0 ml culture was added to 1L LB with antibiotics at above concentrations. The culture

5 was shaken at 37°C until $A_{600} = -0.400$. 1M IPTG was added to the 1L culture to give a final concentration of 1mM. The culture was then shaken at 37°C for 3-4 h. The 1L culture was spun 15 min. in 250 ml conical tubes at 4000 rpm in a model J-6B centrifuge. The supernatant was 10 discarded and the pellet stored at -20°C until use.

The 1L pellet was resuspended in 25 ml 50 mM NaH,PO, 10mM Tris, 6M GuC1, 300mM NaC1, pH 8.0 (Buffer This mixture was then rotated at room temperature A). The mixture was then subjected to 30 minutes. 15 for sonication (VibraCell Sonicator, Sonics and Materials, Inc., Danbury, CT) using the microtip, two times, for 30 sec., at 50% Duty Cycle and with the output setting at 7. The mixture was spun 5 min. at 10K in a JA20 rotor and the supernatant removed and discarded. The supernatant 20 was loaded on a 10 ml Talon (Clonetech, Palo Alto, CA) resin column attached to a GradiFrac System (Pharmacia Biotech, Upsala, Sweden). The column was equilibrated with 100 ml Buffer A and washed with 200 ml of this 25 buffer. A volume based pH gradient using 100% 50 mM NaH₂PO₄, 8M Urea, 20mM MES, pH 6.0 (Buffer B) as the final target buffer was run over a total volume of 100 Protein eluted at ~ 30% Buffer B. Eluted peaks were ml. collected and pooled.

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For refolding, dialysis was carried out with a 2L volume of PBS at room temperature for approximately 3 hr. using dialysis tubing with a molecular weight cutoff of 14,000. The sample was then dialyzed overnight in 2L of PBS at 4°C. Additional buffer exchange was accomplished during the concentration of the protein using Centriprep-30 spin columns by adding PBS to the spun retentate and

respinning. The protein concentration was determined using the BCA protein assay and the purity visualized using a Coomassie stained 4-20% SDS-PAGE gel.

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Example 3

Passive Protection with Anti-CbpA Truncate NR1XR2 Antisera

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A. Generation of Rabbit Immune Serum

immune serum against CbpA truncate Rabbit was generated at Covance (Denver, PA). Following collection of preimmune serum, a New Zealand white rabbit (#ME101) immunized with 250 CbpA truncate was μq NR1XR2 15 (containing both alpha helix I and alpha helix II amino acid N-terminal repeats that are prepared from 483:58) in Complete Freund's Adjuvant. The rabbit was given a boost of 125 µg CbpA truncate in Incomplete Freund's Adjuvant on day 21 and bled on days 31 and 52.

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B. Passive Protection in Mice

C3H/HeJ mice (5 mice/group) were passively immunized intraperitoneally with 100 μ l of a 1:2 dilution of rabbit sera in sterile PBS (preimmune or day 31 immune sera).

- 25 One hour after administration of serum, mice were challenged with 1600 cfu virulent serotype 6B S. pneumoniae, strain SP317 (obtained from H.R. Masure). Mice were monitored for 14 days for survival.
- 30 Eighty percent of the mice immunized with rabbit immune serum raised against CbpA truncate NR1XR2 protein survived the challenge for 14 days (Figure 2). All mice immunized with preimmune rabbit serum were dead by day 7.

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C. Passive Protection in Mice (Higher Challenge Dose)

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C3H/HeJ mice (10 mice/group) passively were immunized intraperitoneally with 100 μ l of a 1:2 dilution of rabbit sera in sterile PBS (preimmune or day 52 immune sera). One hour after administration of serum, mice were challenged with 3450 cfu virulent serotvpe 6B S. pneumoniae, strain SP317. Mice were monitored for 10 days for survival.

One hundred percent of the mice immunized with 10 rabbit immune serum raised against CbpA truncate NR1XR2 protein survived the challenge for ten days (Figure 3). Ninety percent of the mice immunized with preimmune rabbit serum were dead at day 10.

15 D. Passive Protection in Mice (Against High Virulence)

mice/group) C3H/HeJ (10 mice were passively immunized intraperitoneally with 100 μ l of a 1:2 dilution of rabbit sera in sterile PBS (preimmune or day 52 immune sera). One hour after administration of serum, mice were challenged with 580 cfu virulent serotype 6B S. pneumoniae, strain SPSJ2 (provided by P. Flynn, St. Jude Children's Research Hospital, Memphis, TN). Mice were monitored for 10 days for survival.

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Fifty percent of the mice immunized with rabbit immune serum raised against CbpA truncate NR1XR2 protein survived the challenge for 10 days (Figure 4). All of the mice immunized with preimmune rabbit serum were dead at day 8.

These data demonstrate that antibodies specific for CbpA protective against systemic are pneumococcal infection. The data further indicate that the cholinebinding region is not necessary for protection, 35 as antibody specific for truncated protein NR1XR2, lacking the choline-binding repeats, was sufficient for In addition, serum protection. directed against

recombinant CbpA protein based on a serotype 4 sequence, was still protective against challenge with two different strains of serotype 6B.

Example 4

Active Protection with Anti-CbpA Truncates NR1X and NR1XR2

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Active Protection With NR1X Truncate Vaccination Α. (10/group)C3H/HeJ mice were immunized intraperitoneally with CbpA truncate protein NR1X (15µq in 50 µl PBS, plus 50 µl Complete Freund's Adjuvant). Α 10 sham immunized mice received of PBS aroup and A second immunization was administered four adjuvant. weeks later, 15 μg protein i.p. with Incomplete Freund's 15 Adjuvant (sham group received PBS plus IFA). Blood was drawn (retro-orbital bleed) at weeks 3, 6, and 9 for analysis of immune response. The ELISA end point anti-CbpA truncate titer of pooled sera from the 10 CbpA 20 immunized mice at 9 weeks was 4,096,000. No antibody was detected in sera from sham immunized mice. Mice were challenged at week 10 with 560 CFU serotype 6B S. pneumoniae strain SPSJ2. Mice were monitored for 14 days for survival. 25

Eighty percent of the mice immunized with CbpA truncate protein NR1X survived the challenge for 14 Days (results shown in Figure 5). All sham immunized mice were dead by day 8.

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Β. Active Protection With NR1XR2 Truncate Vaccination

C3H/HeJ mice (10/group)immunized were intraperitoneally with CbpA truncate protein NR1XR2 (15µg in 50 µl PBS, plus 50 µl Complete Freund's Adjuvant). A 35 group of 10 control immunized mice received pneumococcal recombinant protein SP90 and adjuvant. Α second

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immunization was administered four weeks later, 15µg protein i.p. with Incomplete Freund's Adjuvant. Blood was drawn (retro-orbital bleed) at weeks 3, 6, and 9 for analysis of immune response. The ELISA end point anti-CbpA truncate titer of pooled sera from the 10 CbpA immunized mice at 9 weeks was 4,096,000. Mice were challenged at week 10 with 680 CFU serotype 6B S. pneumoniae strain SPSJ2. Mice were monitored for 14 days for survival.

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Fifty percent of the mice immunized with CbpA truncate protein NR1XR2 survived the challenge for 14 days (results shown in Figure 6). All control immunized mice were dead by day 9.

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These data demonstrate that immunization with recombinant CbpA truncate proteins elicit production of specific antibodies capable of protecting against systemic pneumococcal infection and death. The data 20 further indicate that the choline-binding region is not immunogens necessary for protection, as the were truncated proteins NR1X and NR1XR2. Additionally, the results suggest that a single amino terminal repeat may be sufficient to elicit a protective response. Cross 25 demonstrated protection is as the recombinant pneumococcal protein was generated based on serotype 4 DNA sequence and protection was observed following challenge with a serotype 6B isolate.

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Numerous modifications and variations of the present invention are possible in light of the above teachings and, therefore, within the scope of the appended claims, the invention may be practiced otherwise than as particularly described.

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WHAT IS CLAIMED IS:

	1. A vaccine against bacterial infections
	comprising an immunogen which is a polypeptide truncate
5	of a pneumococcal surface-binding protein, analog or
	variant having a highly conserved immunogenic alpha-
	helical portion with respect to different types of
	pneumococcal bacteria, which polypeptide does not include
	a choline-binding portion.

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2. A vaccine according to claim 1, wherein the amino acid sequence of said alpha-helical portion has at least 75 % identity with respect to the amino acid sequence of SEQ ID NO:1.

3. A vaccine according to claim 1, wherein the amino acid sequence of said alpha-helical portion has at least 85 % identity with respect to the amino acid sequence of SEQ ID NO:1.

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4. A vaccine according to claim 1, wherein the amino acid sequence of said alpha-helical portion has at least 90 % identity with respect to the amino acid sequence of a member consisting of:

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(a) the amino acid sequence of SEQ ID NO:1, and(b) the amino acid sequence of SEQ ID NO:19.

5. A vaccine according to claim 1, wherein the amino acid sequence of said alpha-helical portion has 30 at least 95 % identity with respect to the amino acid sequence of a member selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO:1, and

(b) the amino acid sequence of SEQ ID NO:19.

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6. A vaccine according to claim 1, wherein said vaccine is for preventing or treating otitis media, sepsis, meningitis and lobar pneumonia infections.

7. A vaccine according to claim 6, wherein said vaccine is for invasive infections.

 8. A vaccine according to claim 6, wherein
 5 said vaccine is for otitis media infections caused by S. pneumoniae.

9. A vaccine according to claim 1, wherein said polypeptide truncate comprise an amino acid sequence
 which has at least 90 % identity with respect to a member selected from the group consisting of the amino acid sequences of each of SEQ ID NOS:3 to 18.

- 10. A vaccine according to claim 1, wherein 15 said polypeptide truncate comprise an amino acid sequence which has at least 95 % identity with respect to a member selected from the group consisting of the amino acid sequences of each of SEQ ID NOS:3 to 18.
- 20 11. An antibody raised against an immunogen which is a polypeptide truncate of a pneumococcal surface-binding protein, analog or variant having a highly conserved immunogenic alpha-helical portion with respect to different types of pneumococcal bacteria, 25 which polypeptide does not include a choline-binding portion.

12. An antibody according to claim 11, wherein the amino acid sequence of said alpha-helical portion has
30 at least 85 % identity with respect to the amino acid sequence of SEQ ID NO:1.

13. An antibody according to claim 11, wherein the amino acid sequence of said alpha-helical portion has
35 at least 90 % identity with respect to the amino acid sequence of a member selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO:1,

and

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(b) the amino acid sequence of SEQ ID NO:19.

14. An antibody according to claim 11, wherein the amino acid sequence of said alpha-helical portion has 5 at least 95 % identity with respect to the amino acid sequence of a member selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO:1, and

10

(b) the amino acid sequence of SEQ ID NO:19.

15. An antibody according to claim 11, wherein said polypeptide truncate comprise an amino acid sequence which has at least 95 % identity with respect to a member 15 selected from the group consisting of the amino acid sequences of each of SEQ ID NOS:3 to 18.

16. An antibody according to claim 11, wherein said antibody is an antibody that will detect S.
20 pneumoniae infections.

17. An antibody according to claim 15, wherein said antibody is effective for the prevention and/or treatment of S. pneumoniae infections.

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18. An antibody according to claim 15, wherein said antibody is effective for the prevention and/or treatment of pneumococcal infections caused by types 1-5, 6A, 6B, 7F, 8, 9N, 9V, 10A, 11A, 12F, 14, 15B, 17F, 18C, 19F, 19A, 20, 22F, 23F and 33F *S. pneumoniae* bacteria.

19. A method for preventing and/or treating pneumococcal infections in a host comprising immunizing said host with a member selected from the group consisting of:

(a) a vaccine according to claim 2, and

(b) at least one antibody raised against an immunogen which is a polypeptide truncate of a pneumococcal surface-binding protein, analog or variant

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comprising an amino acid sequence that is has at least 90 % identity to the amino acid sequence of a member selected from the group consisting of SEQ ID NO:3 to 18, which polypeptide does not include a choline-binding portion.

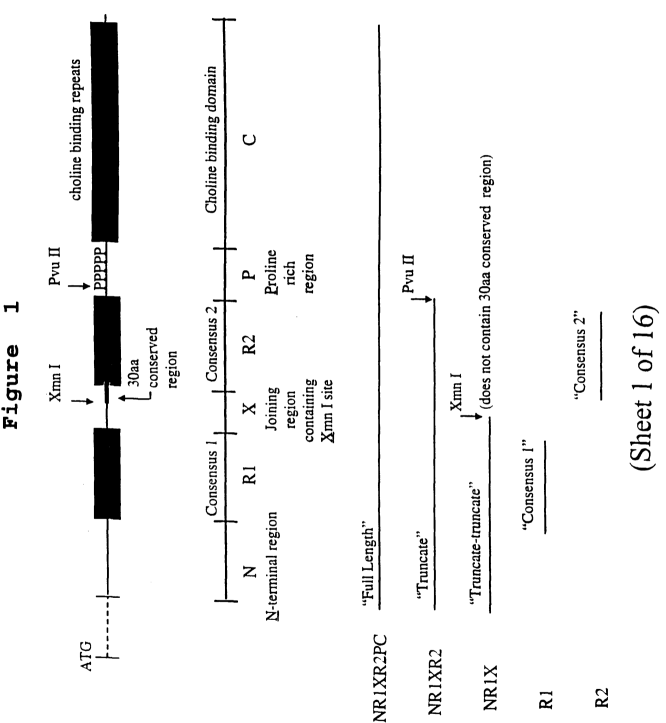
20. A polypeptide comprising an amino acid sequence which has at least 90 % identity with respect to a member selected from the group consisting of the amino
10 acid sequences of each of SEQ ID NOS:3 to 18.

21. An isolated polynucleotide comprising polynucleotide sequence having at least 90 % identity to a member selected from the group consisting of:

(a) a polynucleotide coding sequence encoding
 a polypeptide comprising a member selected from the group
 consisting of the amino acid sequences of each of SEQ ID
 NOS:3 to 18, and

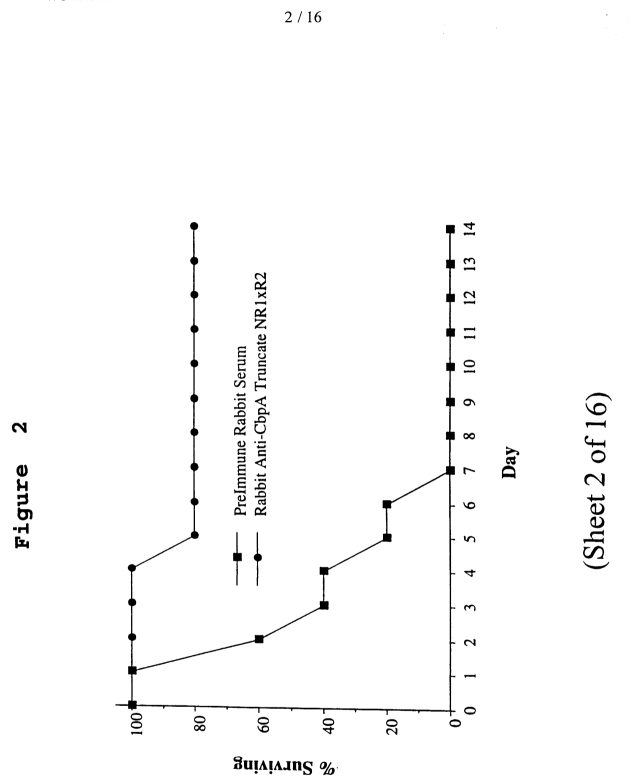
(b) and the complement of (a).



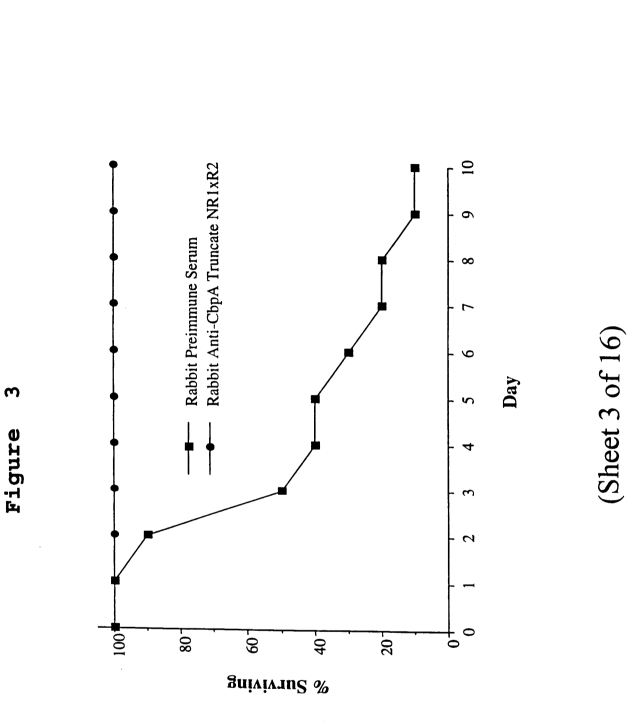


Substitute Sheet (Rule 26)

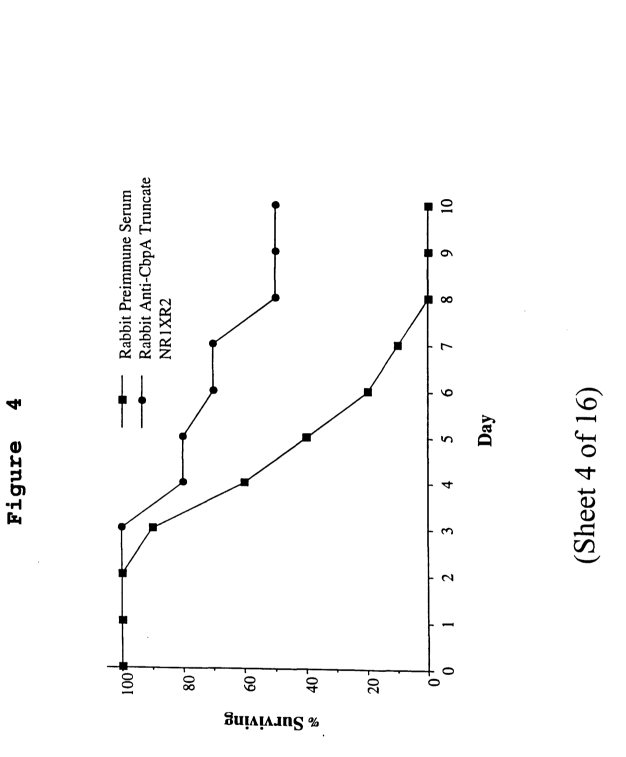
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Substitute Sheet (Rule 26)

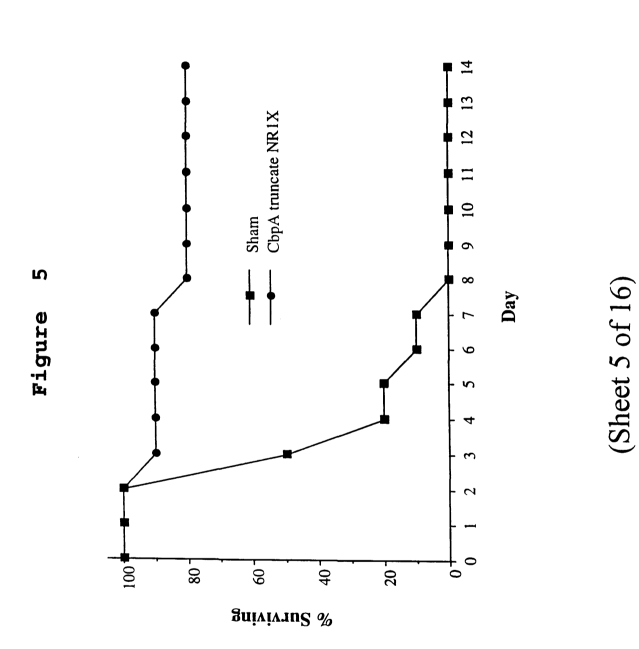


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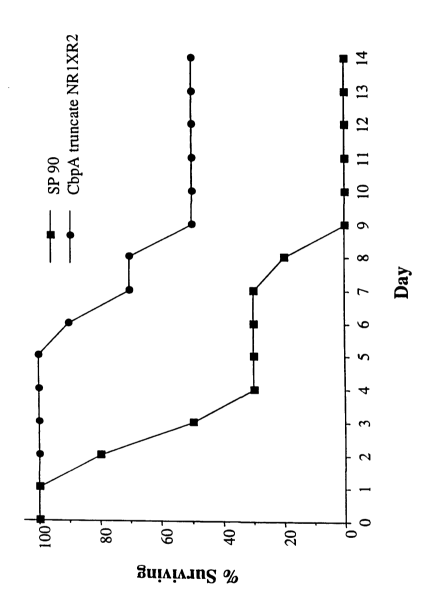
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Substitute Sheet (Rule 26)

Figure 6



Majority Norway4 Arrcc13400(1) Arrcc1733(2) Arrcc1733(2) Arrcc6 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc68 Arrcc1173(2) Arrcc68 Arrcc70 Arrcc68 Arrcc1173(2) Arrcc68 Arrcc70 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc68 Arrcc1173(2) Arrcc1173	Majority Nurway4 Aucco1733400(1) Aucco1733400(1) Aucco Aucca
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Figure

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Substitute Sheet (Rule 26)

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(Sheet 7 of 16)

Decoration 'Decoration #1': Box residues that differ from the Consensus. Decoration 'Decoration #3': Box residues that match the Consensus exactly.

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		Norway4	ATCC33400(1)	ATCC11733(2)	ATCC2	ATCC4	ATCC6B	ATCC18C	R6X(2)	SPB105(6B)	SPB328(23F)	SPB331(14)	SPB365(23F)	SPB609(6B)	SPR332(9V)	SPSJ2(6B)p	SPSJ9(14)	SPSJ12(19A)
		-	2	3	4	5	9	7	œ	6	10	11	12	13	14	15	16	17
	17	37.0	62.0	65.0	60.2	37.0	38.0	34.0	65.0	67.0	81.0	81.0	82.0	29.1	63.5	65.0	41.0	T
	16	23.3	37.6	46.5	38.8	23.3	64.1	60.2				45.1	45.1	66.3	42.7	46.5	T	,
	15	31.7	63.4		78.6	31.7	40.6	36.6	100.0	62.0 42.0	60.4	60.4	60.4	30.2	59.4	T	,	
	14	32.3	59.4	59.4 100.0	67.7	32.3	41.7	38.5	59.4	65.6	63.5	63.5	64.6	23.3	T	,		
	13	30.2	27.9 59.4 63.4	30.2	24.4	30.2	51.2	48.8	60.4 60.4 60.4 30.2 59.4 100.0 46.5	70.0 70.0 71.0 27.9 65.6	27.9 63.5 60.4 45.1	27.9	27.9	T				
	12	37.3		60.4	63.3	37.3	37.3	34.3	60.4	71.0		99.0	T					
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(Sheet 8 of 16)

Substitute Sheet (Rule 26)

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Figure

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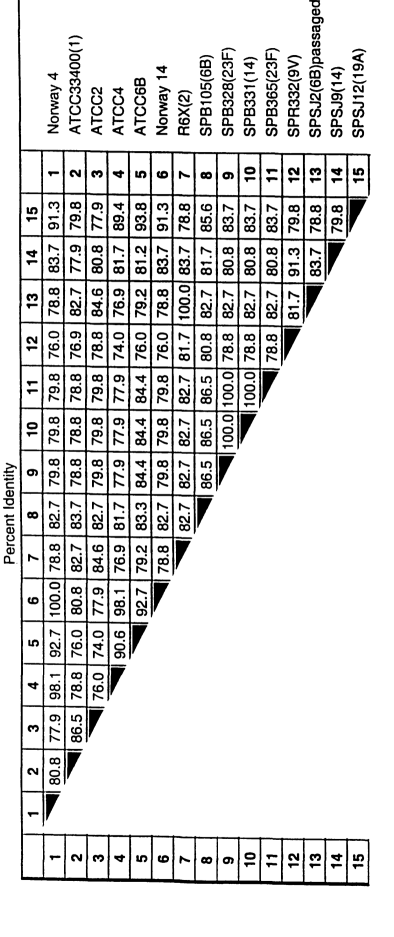
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(Sheet 9 of 16)

Decoration 'Decoration #3': Box residues that match the Consensus exactly.

Decoration 'Decoration #2': Box residues that differ from the Consensus.

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(Sheet 10 of 16)

0(1) 0(1) 3) 35 35 35 31 31 31 31 31 31 31 31 31 31 31 31 31	Majority 01 / 1	Normary Arrcc33400(1) Arrcc4 Arrcc6B	Norway14 R6X(2) SPB105 (6B) SPB328 (23F) SPB331 (14)	SPB365 (23F) SPB609 (6B) SPR332 (9V) SPSJ2 (6B) passaged SPSJ12 (19A) SPSJ12 (19A)		
SEODK PKGRAKRGVPGELATPDKKENDAKSSDSSVGEETLPS - EOGK PKGRAKRGVPGELATPDKKENDAKSSDSSVGEETLPS EVKNKLKBRTKRÅVPGELATPDKKENDAKSSDSSVGEETLPS - ECGK PKGRPKRAVPGELATPDKKENDAKSSDSSVGEETLPS - ECGK PKGRPKRAVPGELATPDKKENDAKSSDSSVGEETLPS - ECGK PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECGK PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECGR PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECGR PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECGR PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECGR PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECOGR PKGRPKRGVPGELATPDKKENDAKSSDSSVGEETLPS - ECOGR PKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR PKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR DKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR DKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR DKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR DKRRVEVLGELATPDKKENDAKSSDSSVGEETLPS - ECOGR DKRRVKENDAKSSDSSVGEETLPS - ECOGR DKRRVKENDAKSSDSSVGEETLPS - ECOGR DKRRVKENDAKSSDSSVGEETLPS - ECOGR DKRRVKENDAKSSDSSVGEETLPS - ECOGR DKRKENDAKSSDSSVGEETLPS - ECOGR DKRKENDAKSSDSSVGEETLPS - ECOGR DKRKENDAKSSDSSVGEETLPS - ECOGR DKRKENDAKSSDSSVGEETLPS - ECOGR DKKENDAKSSDSSVGEETLPS - ECOR DKKENDAKSSDSSVGEETLPS - ECOR DKKENDAKSSDSSVGEETLPS - ECOR DKKENDAKSSDSSVGEETLPS - ECOR	Majori		Norward Norward R6X(2) SPB105 SPB328 SPB331	358452 CERAR2 SEBAR2 SECTOR SE	Box residues that differ from the Consensus. Box residues that match the Consensus exactly.	(Sheet 11 of 16)
$\begin{array}{c} \mathbf{D} \mathbf{A} \mathbf{K} \mathbf{L} \\ \mathbf{D} \mathbf{A} \mathbf{L} \\ \mathbf{D} \mathbf{A} \mathbf{L} \\ \mathbf{D} \mathbf{L} \mathbf{L} \\ \mathbf{D} \mathbf{A} \mathbf{L} \\ \mathbf{D} \mathbf{L} \\ \mathbf{D} \mathbf{L} \\ \mathbf{D} \mathbf{L} \\$	P S L K P	<u>, , , , , , , , , , , , , , , , , , , </u>	. 4 .9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	888888 88888 888888 888888 888888 888888	Decoration 'Decoration #1': Box r Decoration 'Decoration #2': Box r	

Substitute Sheet (Rule 26)

Figure 11

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		Norway4	ATCC33400(1)	ATCC2	ATCC4	ATCC6B	Norway14	R6X(2)	SPB105(6B)	SPB328(23F)	SPB331(14)	SPB365(23F)	SPB609(6B)	SPR332(9V)	SPSJ2(6B)passaged	SPSJ9(14)	SPSJ12(19A)	
		-	2	3	4	2	9	2	8	6	9	11	12	13	14	15	16	
	16	76.0	46.3	69.8	76.0	70.4	76.0	66.7	66.7	64.8	64.8	64.8	66.7	88.7	66.7	88.7	7	
	15	76.0	50.9	71.7	76.0	71.7	76.0	66.0	67.9	67.9	69.8	67.9	75.5	92.5	66.0	7		
	14	86.0	43.1 50.9	7.17 71.7	84.0	62.7	86.0	100.0	79.7	81.4	81.4	81.4	78.0	66.0	T	I		
	13	76.0	49.1	69.8	76.0	71.7	84.0 76.0 86.0 76.0 76.0	66.0 100.0 66.0	69.8	67.9	67.9	69.8	69.8	7				
	12	84.0	48.3	69.8	84.0 76.0 84.0 76.0	67.7 71.7 62.7 71.7 70.4	84.0	78.0	83.9	78.0	79.7	78.0	T	1				
	11	82.0	43.1	69.8	80.0	66.1	82.0	81.4	78.0	98.3	98.3	7						
У	10	82.0	44.8	69.8	80.0	64.4	82.0 82.0	81.4	78.0	98.3	T							
ldentit	6	82.0	43.1	69.8	80.0	64.4	82.0	81.4	78.0	7								
Percent Identity	æ	86.0	44.8	71.7 71.7	86.0	62.7 64.5 64.4	86.0	79.7	T									
۵.	7	86.0	43.1	71.7	84.0	62.7	86.0	7										
	9	100.0	52.0	76.0	98.0	82.0	7											
	2	82.0	55.2	81.1	82.0	1												
	4	98.0	52.0	76.0														
	3	76.0	58.5															
	2	52.0																
	-																	
		1	7	e	4	5	9	7	8	6	10	11	12	13	14	15	16	

12 Figure

(Sheet 12 of 16)

Substitute Sheet (Rule 26)

									aged		13 /	16												iged				
Majority		ATCC334 ATCC334	ATTC2	ATCC6B	Norwery14	KOX(2) SPB105(6B)	SPB328 (23F) SPB331 (14)	SPB365 (23F) SPR332 (9V)	SPSJ2 (6B) passaged	SPSJ12 (19A) Consensus A	Majority		Norwery 4	ATCC33400(1)	ATCC11733 (2)	ATTC2	ATCC6B	ATCC18C-3	Norway14	R6X (2) SDR105 (6R)	SPB328 (23F)	SPB331(14)	SPB365 (23F) SPR332 (9V)	SPSJ2 (6B) passaged	SPSJ9(14) SPSJ12(19A)	Consensus A		
<u> </u>	· 3-	VEVKAELELVKEEAN VKVKEAELELVKEEAK VKVKEAELELVKEEAK	VIXV XIZAELELV XEEAX VEV X X A FLELV XEEAX	· · · · · · · · · · · · · · · · · · ·		VKV KEAELELVKEEAK VKV KEAELELVKEEAK	V E V K K A E L E L V K E E A K V E V K K A E L E L V K E E A K	VEVKKAELELVKEEAK VKVKEAELELVKEEAK	VIXIV XIEAELELVKEEAK VEVKKAELELVKEEAK	រេខាត																		
YKTLELEIAESD	- 6	X X T L E L E L A E S D X X T L E L E L A E S D X X T L E L E I A E E D	YKTLDLEIAESD VKTLFLEIAESD			Y K T L D L E I A E S D Y K T L E L E I A E S D	Y K T L E L E I A E S D Y K T L E L E I A E S D	Y K T L E L E I A E S D Y K T L E L E I A E S D	YKTLDLEIAESD YKTLELEIAESD	Y K T L E L E L A E S D Y K T L E L E L A E S D Y K T L E L E L A E S D	K K A E E E - A K R K A	100	KKAEEE-AKRKA	ккаее – – акк ки	KKAEEE-AKRKA	KKAEEU-AKRKA KKAFEEU-AKFKA	KKAEEE-AKRKA	А - - -	KAEEE-AK	K K A E E E - A K K K A K K A E E I A K R K A	RAEEDDAKRR	KAEEEEAKRR	KRAEEEEAAKRRA KKAFEEI-AKRKA	KAEEE-AKRK	KKAEEE-AKRKA KKAEEE-AKRKA	AEEB-AKR	. SI	
K E E D R R N Y P T N T	R-1	K E E U K K N Y F T N T K E E U R R N Y P T N T K E E U R R N Y P T N T	ម ម ម	រ យក្រ រ យ ផ	ង ខ្មារ ខ្មារ	មា ខា	о о ы ы ы ы	យ	а н 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		ΑΤΚΓΕΚΙΚΤΟΚ	- 8-	ATRL	ATRLE	А Т Т Т Т Т Т Т	А Т К Т К Г Г К Г	ATRL	ATRLEKIKTD	A T R L E	A T A T A T A T A T A T A T A T A T A T	ATRLENIKTD	ATRL	4 4 4 4	ATRLENIKT	ATRL ATRL	ATRLEKIKT	differ from the Consensus	
EEAKKKAEDO	· 23- 0 	LEEAKKKAEUQK LAEABKKAKOQK EEAKKKAEDOK	АЕАЕККАКОО ЕЕАККАКОО			BEAEKKAKAO DEAEKKAKADO	E E A K K K A E D Q E E A K K K A E D Q	EEAKKKAEDO EEADKKAKAOO	E E A E K K A K D O	EEAKKKAEDO	A K V E S K K A	- 8-	KAEVESKKA	KAKVESKKA	KIEKVESKKA Vieteka	K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K A E < K	K A B V E S K Q A	KAEVESKOA	K ALEVESKKA V v v v v v v v v v v v v v v v v v v v	K A K V E S K K A K A K V E S K K A	KAKVESKKA	KAKVESKKA	K A K V E S K K A K A K V E S K K A	KAKVESKKA	A K A E V E S K K A E A K A E V E S K K A E	KAKVESKKA	: Box residues that d	
<u> </u>		L KKVAEAEKKV L KKVAEAEKKV L EKVAEAKKV	K K V A E A E K K V K K V A E A E K K V	K K V A E A E K K V K K V A E A E K K V	KKVAEAEKKV	KKVAEAEKKV KKVAEAEKKV	K K V A E A E K K V K K V A E A E K K V	K K V A E A E K K V K K V A E A E K K V	K K V A E A E K K V K K V A E A E K K V	K K V A E A E K K V K K V A E A E K K V	A N E E K I K O		EPLENEEKUKO	E P O N E E K I K O	ESRNEGUIKO	EPRNEEKVKO	EPRNEEKVKO	EPRNEEKVKO	E P R N E E KUKO		ESRNEEKIKO	ESRNEEKIKO	E S R N E E K I K Q	EPRDEEKIKO	<u>GS</u> RNEEKVKO EPRNEEKVKO	ESRNEEKIKO	Decoration 'Decoration 1'	
													Ø	6	6	06	5 10	6	6	d (d	0	6	30	6	ଟ ଟ	ស	Ä	

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Figure

Substitute Sheet (Rule 26)

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											• •	/ 1	0						
		Norway 4	ATCC33400(1)	ATCC11733(2)	ATCC2	ATCC4	ATCC6B	ATCC18C-3	Norway14	R6X(2)	SPB105(6B)	SPB328(23F)	SPB331(14)	SPB365(23F)	SPR332(9V)	SPSJ2(6B)passaged	SPSJ9(14)	SPSJ12(19A)	Consensus A
		1	2	3	4	5	9	7	8	o	9	=	12	13	14	15	16	17	18
	18	97.1	89.2	91.3	91.3	97.1	95.1	93.2	97.1	92.2	92.2	93.2	94.2	93.2	95.1	92.2	93.2	94.2	T
	17	97.1	88.2	85.4	93.2 92.2 90.3	97.1	95.1	93.2	97.1	91.3	93.2	87.4	87.4 88.3 94.2		95.1	91.3	94.2	7	
	16	94.2	86.3	84.5	92.2	91.3 94.2 97.1	92.2	92.2	94.2	89.3	91.3	87.4 86.4 87.4	87.4	86.4 87.4	91.3	89.3	T	,	
	15	91.3	91.2	89.3	93.2	91.3	91.3 92.2	87.4	91.3	100.0	92.2	87.4	88.3	87.4	94.2	T	,		
	14	92.2	91.2	90.3	93.2	92.2	90.3	88.3	92.2	94.2			89.3	88.3	T	,			
	13	90.3	86.3	86.4	93.2 95.1 84.5 85.4 84.5	90.3 92.2	90.3	86.4	90.3	87.4 94.2	86.4 85.4 96.1	99.0 100.0 88.3	99.0	7	,				
	12	91.3	87.3	87.4	85.4	100.0 91.3 91.3 90.3 91.3	91.3	87.4	91.3	88.3	86.4	99.0	T						
Y	11	90.3	86.3	86.4	84.5	90.3	90.3	86.4	90.3	87.4	85.4	1							
	10	91.3	91.2	87.4	95.1	91.3	89.3	87.4	91.3	92.2	7								
בורבווו ותכוווווא	6	91.3	91.2	89.3	93.2	91.3	91.3	87.4	91.3	7									
-	8	96.1 100.0	88.2	88.3	90.3	100.0	98.1	96.1	7										
	2	96.1	84.3	86.4	86.4	96.1	96.1												
	9	98.1	86.3	88.3	88.3	98.1													
	2	100.0	88.2	88.3	90.3														
	4	90.3	90.2	86.4															
	3	88.3	84.3	Ī															
	2	88.2	T																
	-																		
		-	2	9	4	5	9	7	8	6	10	=	12	13	14	15	16	17	18
		-	-	-	-			-											

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Figure 14

Percent Identity

Substitute Sheet (Rule 26)

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W	O 99/51266	5													PC7	ſ/U	S99/076
				saged	15	5 / 16	Ď								saged		
	Norway4 ATCC33400(1) ATCC11733(2) ATCC2 ATCC2			, se	SPSJ9(14) SPSJ12(19A) Consensus B	Majority		Norway4 ATCC33400(1) ATCC3173400	ATCC2 ATCC4	ATCC6B ATCC18C-3	Norway14	R6X (2) SPB105 (6B)	SPB328 (23F) SPB331 (14)	SPB365 (23F) SPR332 (9V)	SBC	SPSJ12 (19A)	Consensus B
<u>KKVAEAEKKYEEAKKKAEDQKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK</u> 10 20 30 40 50 60	K K V A E A E K K V E E A K K K A E D Q K E E D R N Y P T N T Y K T L E L E I A E S D V E V K A E L E L V K E E A K K K V A E A E K K V A E A F K A K D Q K E E D R R N Y P T N T Y K T L E L E I A E S D V K V K E A E L E L V K E E A K E K V A E A E K K V E E A K K A E D Q K E E D R R N Y P T N T Y K T L E L E I A E E D V K V K E A E L E L V K E E A K K K V A E A E K K V A E A B K A K D Q K E E D R R N Y P T N T Y K T L E L E I A E E D V K V K E A E L E L V K E E A K K K V A E A E K K V A E A B K A K D Q K E E D R R N Y P T N T Y K T L D L E I A E S D V K V K E A E L E L V K E E A K	KVAEAEKKVEEAKKKAEDQKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK KVAEAEKKVEEAKKKAEDQKEEDRRNYPTIJTYKTLELEIAESDVEVKKAELELVKEEAK KVAEAEKVEEAKKKAEDQKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK KVAEAEKVVEEAKKKAEDQKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK	KKVAEAEKKVEEAEKKKAEDOKEEDRRNYPTNTYKTLDLEIAESDVKVKEAELELVKEEAK KKVAEAEKKVEEAKKKAEDOKEEDRRNYPTNTYKTLELEIAESDVKVKAELELVKEEAK KKVAEAEKVVEEAKKKAEDOKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK	KKVAEAEKKVEEADKKAKAOKEDRRNYPTNTYKTLELEIAESDVEVKEAELELVKEEAK KKVAEAEKKVEEADKKAKAOKEDRRNYPTNTYKTLELEIAESDVKVKEAELELVKEEAK	KKVAEAEKKVEEA EKKA KDQKEED HRNYPT JTYKTLELEIAESDVEVKKAELELVKEEAK KKVAEAEKKVEEA DKKA KAQKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK KKVAEAEKKVEEAKKAEDOKEEDRRNYPTNTYKTLELEIAESDVEVKKAELELVKEEAK	E P R N E E K I K Q A K A K V E S K K A E A T R L E K I K T D R K K A E E E - A K R K A A E E D K V K E K P A		D R K K A E E E – A K R K A A E E D K V K E K P A D R K K A E E – – A K R K V A E E D K V K E K P A D P K V S E E – – A K R K V A E E D K V K E K P A	GERNEEKINGAKABVESKKAEATRLEKIKTDRKKAEEE-AKRKAAEEDKVKEKPA	E PRNEEKVKOAKABVESKOAEATRLENIKTDRKKAEEE – AKRKAAEEDKVKEKPA E PRNEEKVKOAKABVESKOAEATRLENIKTDRKKAEEE – AKRKAAEEDKVKEKPA	EPRNEEKUKOAKAEVESKKAEATRLEKIKTDRKKAEEE-AKRKAAEEDKVKEKPA	E PRDEEKIKQAKAKVESKKAEATRLENIKTDRKKAEEE - AKRKAAEEDKVKEKPA E PRDEEKIKQAKAKVESKKAEATRLENIKTDRKKAEEE - AKRKAAEEDKVKEKPA	ESRNEEKIKOVKAKVESKKAEATRLENIKTDRKRAEEFEFAKRAAEEDKVKEKPA FSRNEEKIKOVKAKVESKKAFATRLENIKTDRKKAEEFFFAKRRAAEFDKVKEKPA	ESRNEEKIKOVKAKVESKKAEATRLENIKTDRKRAEEBEAKRRAAEEDKVKEKPA FSRNEEKIKOAKAKVESKKAEATRLEKIKTDRKKAEEB-AKRKAAEED-AKRKAAEEDA	EPRIDEEKIKQAKAKVESKKAEATRLENIKTDRKKAEEE-AKRKAAEEDKVKEKPA	EPRNEEKUKOAKAEVESKKAEATRLEKIKTDRKKAEEE-AKRKAAEED-AKRAAEEDKVKEKPA	EPRNEEKIKOAKAKVESKKAEATRLEKIKTDRKKAEEEL-AKRKAAEEDKVKEKPA coration 'Decoration #1': Box residues that differ from the Consensus.

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Figure

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Decoration '': Box residues that match the Consensus exactly.

		Norway4	ATCC33400(1)	ATCC11733(2)	ATCC2	ATCC4	ATCC6B	ATCC18C-3	Norway14	R6X(2)	SPB105(6B)	SPB328(23F)	SPB331(14)	SPB365(23F)	SPR332(9V)	SPSJ2(6B)passaged	SPSJ9(14)	SPSJ12(19A)	Consensus B	
		Ļ	2	3	4	5	9	7	80	6	9	÷	12	13	14	15	16	17	-18	
	18	98.2	91.2	91.2	91.2	98.2	96.5	94.7	98.2	93.9	93.9	93.0	93.9	93.0	94.7	93.9	93.0	92.6	Ţ	
	17	97.4	89.4	86.8	91.2	97.4	95.6	93.9	97.4	92.1	92.1	_	89.5	88.6	95.6	92.1	94.7	T	•	
	16	94.7	87.6	86.0		94.7	93.0	93.0	94.7		_	87.7	88.6	87.7	32.1	90.4	Ţ	ľ		
	15		92.0	90.4	93.9 93.0	92.1	92.1	88.6	92.1	100.0 90.4	100.0 90.4	38.6	89.5	88.6	94.7 92.1	Ī	J.			
	14	93.0 92.1	92.0	91.2	93.9	93.0	91.2	89.5	93.0 92.1	94.7 1	94.7 1	99.1 100.0 89.5 88.6 87.7 88.6	90.4	89.5 8	Ī	ľ				
	13	91.2	86.7	87.7	86.0	91.2	91.2	87.7	91.2	88.6	88.6 5	0.00	99.1 9		r					
	12	92.1	87.6	88.6		92.1	92.1		92.1 9	89.5 8	89.5 8	9.1 1		•						
	11	91.2	86.7 8	87.7	86.0 86.8	91.2	91.2 5	87.7 88.6	91.2 9	88.6 8	88.6 8	6								
Percent Identity	10	92.1	92.0	90.4	93.9 8	92.1 9	92.1 9	88.6 8		100.0 8		7								
rcent l	6	92.1	92.0	90.4 5			92.1 9	88.6 8	92.1 92.1	Ī	/									
Ре	8	100.0	89.4 5	89.5 5	91.2 93.9	100.0 92.1	98.2 9	96.5 8	6	/										
	7	96.5 1	85.8 8	87.7 8	87.7 9	96.5 1	96.5 9	6	/											
	9	98.2 9	87.6 8	89.5 8	89.5 8	98.2 9	6 7	7												
	5	100.0 9	89.4 8	89.5 8	91.2 8	6 7	7													
	4	91.2 1	91.2 8	87.7 8	6	1														
	3	89.5 9	85.8 9	®	1															
	2	89.4 8	80 	7																
	1	8	1																	
		-	2	3	4	S	9		æ	6	0	-	2	<u>س</u>	4	2		2		
					-	-					-	-	Ŧ	₽	Ť	Ŧ	16	-	8	

16 Figure

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(Sheet 16 of 16)

Substitute Sheet (Rule 26)

SEQUENCE LISTING <110> Wizemann, Theresa M. Koenig, Scott Johnson, Leslie S <120> Derivatives of Choline Binding Proteins for Vaccines <130> 469201-364 <140> <141> <150> US 60/085,743 <151> 1998-05-15 <160> 38 <170> MS-Word (DOS Text) <210> 1 <211> 103 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 1 Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Lys 1 5 10 15 Ala Glu Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr 20 25 30 Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys 35 40 45 Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn 50 55 60 Glu Glu Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala 65 70 75 80 Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu 85 90 95 Glu Glu Ala Lys Arg Lys Ala 100 <210> 2 <211> 141 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae

<400> 2

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Tyr Pro Thr Ile Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Gly Ser Arg Asn Glu Glu Lys Ile Lys Lys Ala Lys Ala Glu Val Glu Ser Lys Lys Ala Glu Ala Thr Lys Leu Glu Glu Ile Lys Thr Glu Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Glu Ala Glu Glu Glu Val Lys Asn Lys Leu Lys Lys Arg Thr Lys Arg Gly Ala Phe Gly Glu Pro Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Asp Ser Ser Val Val Lys Lys Ser Ser Lys Pro Ile Leu Lys Ser Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Pro Gln Asn Glu Glu Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Ala Lys Arg Lys Val Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Lys Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Ala Glu Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu Glu Asp Tyr Ala Arg Arg Ser Glu Glu Glu Tyr Asn Pro Leu Asp Leu Thr Ala Pro Ala Lys

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Glu Lys Met Leu Arg Glu Ile Gln Leu Asp Arg Arg Lys His Thr Gln 35 40 45
Asn Val Ala Leu Asn Ile Lys Leu Ser Ala Ile Lys Thr Lys Tyr Leu 50
Arg Glu Leu Asn Val Leu Glu Glu Lys Ser Lys Asp Glu Leu Pro Ser 65 70 75 80
Glu Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Lys Phe Lys Lys Asp 85 90 95
Thr Leu Lys Pro Gly Glu Lys Val Ala Glu Ala Lys Lys Lys Val Glu 100 105 110
Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Asn 115 120 125
Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Phe 130 135 140
Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala 145 150 155 160
Lys Glu Ser Arg Asn Glu Gly Thr Ile Lys Gln Ala Lys Glu Lys Val 165 170 175
Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp 180 185 190
Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp 195 200 205
Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Thr 210 215 220
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Streptococcus pneumoniae

<400> 5 Thr Glu L 1	ys Glu	Val 1 5	Chr Thr	Gln	Val	Pro 10	Thr	Tyr	Ser	Asn	Met 15	Ala
Lys Thr G	lu His 20	Arg I	Lys Ala	Ala	Lys 25	Gln	Val	Val	Asp	Glu 30	Thr	Ile
Glu Lys M	et Leu 35	Arg G	3lu Ile	Gln 40	Leu	Asp	Arg	Arg	Lys 45	His	Thr	Gln
Asn Phe A 50	la Phe	Asn M	let Lys 55		Ser	Ala	Ile	Lys 60	Thr	Glu	Tyr	Leu
Tyr Gly L 65	eu Lys	Glu I	ys Ser 70	Glu	Ala	Glu	Leu 75	Pro	Ser	Glu	Val	Lys 80
Ala Lys L	eu Asp	Ala A 85	ala Phe	Glu	Gln	Phe 90	Lys	Lys	Asp	Thr	Leu 95	Lys
Pro Gly G	lu Lys 100	Val A	ala Glu	Ala	Lys 105	Lys	Lys	Val	Ala	Glu 110	Ala	Glu
Lys Lys A 1	la Lys 15	Ala G	ln Lys	Glu 120	Glu	Asp	Arg	Arg	Asn 125	Tyr	Pro	Thr
Asn Thr T 130	yr Lys	Thr L	eu Glu 135		Glu	Ile	Ala	Glu 140	Ser	Asp	Val	Glu
Val Lys Ly 145	ys Ala		eu Glu	Leu	Leu	Lys	Glu 155	Glu	Ala	Lys	Thr	Arg 160
Asn Glu A	sp Thr	Ile A 165	Asn Gln	Ala	Lys	A la 170	Lys	Val	Glu	Ser	Lys 175	Lys
Ala Glu A	la Thr 180	Leu L	ys Glu	Glu	Ile 185	Lys	Thr	Asp	Arg	Lys 190	Lys	Ala
Glu Glu G 1	lu Ala 95	Lys A	arg Lys	Ala 200	Glu	Ala	Glu	Glu	As p 205	Lys	Val	Lys
Asp Lys Lo 210	eu Lys	Arg A	rg Thr 215		Arg	Ala	Val	Pro 220	Gly	Glu	Pro	Ala
Thr Phe P 225	he Lys		Slu Asn 230	Asp	Ala	Lys	Ser 235	Ser	Asp	Ser	Ser	Val 240
Gly Glu G	lu Thr	Leu P 245	Pro Ser	Pro	Ser	Leu 250	Lys	Ser	Gly	Lys	Lys 255	Val
Ala Glu A	la Glu 260	Lys I	ys Val	Ala	Glu 265	Ala	Glu	Lys	Lys	Ala 270	Lys	Asp
Gln Lys G 2	lu Glu 75	Asp A	Arg Arg	Asn 280	Tyr	Pro	Thr	Asn	Thr 285	Thr	Lys	Thr
Leu Asp L 290	eu Glu	Ile A	Ala Glu 295		Asp	Val	Lys	Val 300	Lys	Glu	Ala	Glu
Leu Glu L 305	eu Val	-	Glu Glu 310	Ala	Lys	Gly	Ser 315	Arg	Asn	Glu	Glu	Lys 320

5

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Ile Asn Gln Ala Lys Ala Glu Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Thr Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Thr Glu Glu Pro Glu Asn Pro Ala Pro Ala Pro Lys Pro Glu Lys Pro Ala Glu Gln Pro Lys Ala Glu Lys Thr Asp Asp Gln Gln Ala Glu Glu <210> 6 <211> 446 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 6 Thr Glu Asn Glu Gly Ala Thr Gln Val Pro Thr Ser Ser Asn Arg Ala Asn Glu Ser Gln Ala Glu Gln Gly Glu Gln Pro Lys Lys Leu Asp Ser Glu Arg Asp Lys Ala Arg Lys Glu Val Glu Glu Tyr Val Lys Lys Ile Val Gly Glu Ser Tyr Ala Lys Ser Thr Lys Lys Arg His Thr Ile Thr Val Ala Leu Val Asn Glu Leu Asn Asn Ile Lys Asn Glu Tyr Leu Asn Lys Ile Val Glu Ser Thr Ser Glu Ser Gln Leu Gln Ile Leu Met Met Glu Ser Arg Ser Lys Val Asp Glu Ala Val Ser Lys Phe Glu Lys Asp Ser Ser Ser Ser Ser Ser Ser Asp Ser Ser Thr Lys Pro Glu Ala Ser Asp Thr Ala Lys Pro Asn Lys Pro Thr Glu Pro Gly Glu Lys Val Ala Glu Ala Lys Lys Lys Val Glu Glu Val Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Ile Thr Tyr Lys Thr Leu

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Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala G 180 185 190	lu Leu						
Glu Leu Val Lys Val Lys Ala Asn Glu Pro Arg Asp Lys Gln Ly 195 200 205	ys Ile						
Lys Gln Ala Glu Ala Glu Val Glu Ser Lys Gln Ala Glu Ala T 210 215 220	hr Arg						
Leu Lys Lys Ile Lys Thr Asp Arg Glu Glu Ala Glu Glu Glu A 225 230 235	la Lys 240						
Arg Arg Ala Asp Ala Lys Glu Gln Gly Lys Pro Lys Gly Arg Pro245250250250	ro Lys 55						
Arg Gly Val Pro Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu As260265270	sn Asp						
Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Pro Se275280285	er Pro						
Ser Leu Lys Pro Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Va290295300	al Glu						
Glu Ala Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg An 305 310 315	rg Asn 320						
Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala G 325 330 33	lu Ser 35						
Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Gl 340 345 350	lu Ala						
Lys Glu Pro Arg Asn Glu Glu Lys Val Lys Gln Ala Lys Ala G 355 360 365	lu Val						
Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Th 370 375 380	hr Asp						
Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Gl 385 390 395	lu Asp 400						
Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Al 405 410 41	la Pro 15						
Lys Thr Glu Lys Pro Ala Pro Ala Pro Lys Pro Glu Asn Pro Al 420 425 430	la Glu						
Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu Glu 435 440 445							
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Lys Gln Ala Lys Ala Glu Val Glu Ser Lys Gln Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Ala Pro Lys Asp Glu Lys Pro Ala Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu Glu <210> 8 <211> 219 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 8 Glu Gly Val Arg Ser Gly Asn Asn Ser Thr Val Thr Ser Ser Gly Gln Asp Ile Ser Lys Lys Tyr Ala Asp Glu Val Glu Ser His Leu Gln Ser Ile Leu Lys Asp Val Asn Lys Asn Leu Lys Lys Val Gln His Thr Gln Asn Ala Asp Phe Asn Lys Lys Leu Ser Lys Ile Lys Pro Lys Tyr Leu Tyr Glu Leu Lys Cys Leu Glu Glu Lys Ser Glu Ala Glu Leu Thr Ser Lys Pro Lys Asn Lys Arg Arg Val Thr Ala Ala Phe Glu Gln Phe Lys Lys Asp Thr Leu Ser Thr Glu Pro Glu Lys Lys Val Ala Glu Ala Lys Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Lys Asp Arg Arg Asn Tyr Pro Thr Ile Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Pro Arg Asn Glu Glu Lys Val Lys Gln Ala

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Lys Ala Glu Val Glu Ser Lys Gln Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala <210> 9 <211> 446 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400>9Thr Glu Asn Glu Gly Ala Thr Gln Val Pro Thr Ser Ser Asn Arg Ala Asn Glu Ser Gln Ala Glu Gln Gly Glu Gln Pro Lys Lys Leu Asp Ser Glu Arg Asp Lys Ala Arg Lys Glu Val Glu Glu Tyr Val Lys Lys Ile Val Gly Glu Ser Tyr Ala Lys Ser Thr Lys Lys Arg His Thr Ile Thr Val Ala Leu Val Asn Glu Leu Asn Asn Ile Lys Asn Glu Tyr Leu Asn Lys Ile Val Glu Ser Thr Ser Glu Ser Gln Leu Gln Ile Leu Met Met Glu Ser Arg Ser Lys Val Asp Glu Ala Val Ser Lys Phe Glu Lys Asp Ser Ser Ser Ser Ser Ser Ser Asp Ser Ser Thr Lys Pro Glu Ala Ser Asp Thr Ala Lys Pro Asn Lys Pro Thr Glu Pro Gly Glu Lys Val Ala Glu Ala Lys Lys Val Glu Glu Ala Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Ile Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Val Lys Ala Asn Glu Pro Arg Asp Glu Gln Lys Ile Lys Gln Ala Glu Ala Glu Val Glu Ser Lys Gln Ala Glu Ala Thr Arg

	210					215					220								
Leu 225	Lys	Lys	Ile	Lys	Thr 230	Asp	Arg	Glu	Glu	Ala 235	Glu	Glu	Glu	Ala	Lys 240				
Arg	Arg	Ala	Asp	Ala 245	Lys	Glu	Gln	Gly	Lys 250	Pro	Lys	Gly	Arg	Ala 255	Lys				
Arg	Gly	Val	Pro 260	Gly	Glu	Leu	Ala	Thr 265	Pro	Asp	Lys	Lys	Glu 270	Asn	Asp				
Ala	Lys	Ser 275	Ser	Asp	Ser	Ser	Val 280	Gly	Glu	Glu	Thr	Leu 285	Pro	Ser	Pro				
Ser	Leu 290	Lys	Pro	Glu	Lys	Lys 295	Val	Ala	Glu	Ala	Glu 300	Lys	Lys	Val	Glu				
Glu 305	Ala	Lys	Lys	Lys	Ala 310	Glu	Asp	Gln	Lys	Glu 315	Glu	Asp	Arg	Arg	Asn 320				
Tyr	Pro	Thr	Asn	Thr 325	Tyr	Lys	Thr	Leu	Glu 330	Leu	Glu	Ile	Ala	Glu 335	Ser				
Asp	Val	Glu	Val 340	Lys	Lys	Ala	Glu	Leu 345	Glu	Leu	Val	Lys	Glu 350	Glu	Ala				
Lys	Glu	Pro 355	Arg	Asn	Glu	Glu	Lys 360	Val	Lys	Gln	Ala	Lys 365	Ala	Glu	Val				
Glu	Ser 370	Lys	Lys	Ala	Glu	Ala 375	Thr	Arg	Leu	Glu	Lys 380	Ile	Lys	Thr	Asp				
Arg 385	Lys	Lys	Ala	Glu	Glu 390	Glu	Ala	Lys	Arg	Lys 395	Ala	Ala	Glu	Glu	Asp 400				
Lys	Val	Lys	Glu	Lys 405	Pro	Ala	Glu	Gln	Pro 410	Gln	Pro	Ala	Pro	Ala 415	Pro				
Lys	Ala	Glu	Lys 420	Pro	Ala	Pro	Ala	Pro 425	Lys	Pro	Glu	Asn	Pro 430	Ala	Glu				
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)> 1(Glu		Glu	Gly 5	Ser	Thr	Gln	Ala	Ala 10	Thr	Ser	Ser	Asn	Met 15	Ala				
Lys	Thr	Glu	His 20	Arg	Lys	Ala	Ala	Lys 25	Gln	Val	Val	Asp	Glu 30	Tyr	Ile				
Glu	Lys	Met	Leu	Arg	Glu	Ile	Gln	Leu	Asp	Arg	Arg	Lys	His	Thr	Gln				

Asn Val Ala Leu Asn Ile Lys Leu Ser Ala Ile Lys Thr Lys Tyr Leu Arg Glu Leu Asn Val Leu Glu Glu Lys Ser Lys Asp Glu Leu Pro Ser Glu Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Lys Phe Lys Lys Asp Thr Leu Lys Pro Gly Glu Lys Val Ala Glu Ala Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Phe Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Gly Thr Ile Lys Gln Ala Lys Glu Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Asp Ala Lys Leu Lys Glu Ala Asn Val Ala Thr Ser Asp Gln Gly Lys Pro Lys Gly Arg Ala Lys Arg Gly Val Pro Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Pro Ser Ser Ser Leu Lys Ser Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Asp Leu Glu Ile Ala Glu Ser Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Pro Arg Asp Glu Glu Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Asp Asp Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu

Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Thr Gln Pro Glu Lys Pro Ala Pro Lys Pro Glu Lys Pro Ala Glu Gln Pro Lys Ala Glu Lys Thr Asp Asp Gln Gln Ala Glu Glu <210> 11 <211> 425 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 11 Thr Glu Lys Glu Val Thr Thr Gln Val Ala Thr Ser Ser Asn Arg Ala Asn Glu Ser Gln Ala Gly His Arg Lys Ala Ala Glu Gln Phe Asp Glu Tyr Ile Lys Thr Met Ile Gln Leu Asp Arg Arg Lys His Thr Gln Asn Phe Ala Leu Asn Ile Lys Leu Ser Arg Ile Lys Thr Glu Tyr Leu Arg Lys Leu Asn Val Leu Glu Glu Lys Ser Lys Ala Glu Leu Pro Ser Glu Thr Lys Lys Glu Ile Asp Ala Ala Phe Glu Gln Phe Lys Lys Asp Thr Asn Arg Thr Lys Lys Thr Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Ala Lys Ala Gln Lys Glu Glu Asp His Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asp Asp Glu Lys Ile Lys Gln Ala Glu Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Glu Lys Ala Glu Glu Glu Ala Lys Arg Arg Ala Glu Ala Lys Leu Lys Glu Ala Val Glu Lys Asn Val Ala Thr Ser Glu Gln Asp Lys Pro Lys

Gly Arg Arg Lys Arg Gly Val Pro Gly Glu Gln Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Ala Leu Pro Ser Pro Ser Leu Lys Pro Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Ala Lys Ala Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Lys Val Lys Glu Ser Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Glu Lys Val Asn Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Thr Glu Glu Pro Glu Asn Pro Ala Pro Ala Pro Lys Pro Glu Lys Pro Ala Glu Gln Pro Lys Ala Glu Lys Thr Asp Asp Gln Gln Ala Glu Glu <210> 12 <211> 426 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 12 Thr Glu Lys Glu Val Thr Thr Gln Val Ala Thr Ser Ser Asn Lys Ala Asn Lys Ser Gln Thr Glu His Met Lys Ala Ala Lys Gln Val Asp Glu Tyr Ile Lys Lys Lys Ile Gln Leu Asp Arg Arg Lys His Thr Gln Asn Val Gly Leu Leu Thr Lys Leu Gly Val Ile Lys Thr Glu Tyr Leu His

Gly Leu Ser Val Ser Lys Lys Ser Glu Ala Glu Leu Pro Ser Glu Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Gln Phe Lys Lys Asp Thr Leu Pro Thr Glu Pro Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Lys Asp Leu Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Asp Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asp Glu Lys Lys Ile Asn Gln Ala Lys Ala Lys Val Glu Asn Lys Lys Ala Glu Ala Thr Arg Leu Lys Asn Ile Lys Thr Asp Arg Glu Lys Ala Glu Glu Ala Lys Arg Arg Ala Asp Ala Lys Leu Gln Glu Ala Asn Val Ala Thr Ser Glu Gln Asp Lys Ser Lys Arg Arg Ala Lys Arg Glu Val Leu Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Thr Ser Pro Ser Leu Lys Pro Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Glu Lys Ile Lys Gln Val Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Glu Ala Lys Arg Arg Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Thr Glu Glu Pro Glu Asn Pro Ala

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Pro Ala Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Lys Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu Glu <210> 13 <211> 425 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 13 Thr Glu Lys Glu Val Thr Thr Gln Val Ala Thr Ser Ser Asn Lys Ala Asn Lys Ser Gln Thr Glu His Met Lys Ala Ala Lys Gln Val Asp Glu Tyr Ile Lys Lys Leu Gln Leu Asp Arg Arg Lys His Thr Gln Asn Val Gly Leu Leu Thr Lys Leu Gly Val Ile Lys Thr Glu Tyr Leu His Gly Leu Ser Val Ser Lys Lys Ser Glu Ala Glu Leu Pro Ser Glu Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Gln Phe Lys Lys Asp Thr Leu Pro Thr Glu Pro Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Lys Asp Leu Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Asp Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asp Glu Lys Lys Ile Asn Gln Ala Lys Ala Lys Val Glu Asn Lys Lys Ala Glu Ala Thr Arg Leu Lys Asn Ile Lys Thr Asp Arg Glu Lys Ala Glu Glu Ala Lys Arg Arg Ala Asp Ala Lys Leu Gln Glu Ala Asn Val Ala Thr Ser Glu Gln Asp Lys Ser Lys Arg Arg Ala Lys Arg Glu Val Phe Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu

Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Glu Glu Glu Thr Leu Thr Ser Pro Ser Leu Lys Pro Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Glu Lys Ile Lys Gln Val Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Glu Ala Lys Arg Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Thr Glu Glu Pro Glu Asn Pro Ala Pro Ala Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Lys Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu <210> 14 <211> 424 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 14 Thr Glu Lys Glu Val Thr Thr Gln Val Ala Thr Ser Ser Asn Arg Ala Asn Lys Ser Gln Thr Glu His Met Lys Ala Ala Lys Gln Val Asp Glu Tyr Ile Lys Lys Lys Leu Gln Leu Asp Arg Arg Lys His Thr Gln Asn Val Gly Leu Leu Thr Lys Leu Gly Val Ile Lys Thr Glu Tyr Leu His Gly Leu Ser Val Ser Lys Lys Ser Glu Ala Glu Leu Pro Ser Glu

Ile	Lys	Ala	Lys	Leu 85	Asp	Ala	Ala	Phe	Glu 90	Gln	Phe	Lys	Lys	Asp 95	Thr
Leu	Pro	Thr	Glu 100	Pro	Gly	Lys	Lys	Val 105	Ala	Glu	Ala	Glu	Lys 110	Lys	Val
Glu	Glu	Ala 115	Lys	Lys	Lys	Ala	Glu 120	Asp	Gln	Lys	Glu	Lys 125	Asp	Leu	Arg
Asn	Tyr 130	Pro	Thr	Asn	Thr	Tyr 135	Lys	Thr	Leu	Glu	Leu 140	Asp	Ile	Ala	Glu
Ser 145	Asp	Val	Glu	Val	Lys 150	Lys	Ala	Glu	Leu	Glu 155	Leu	Val	Lys	Glu	Glu 160
Ala	Lys	Glu	Ser	Arg 165	Asp	Glu	Lys	Lys	Ile 170	Asn	Gln	Ala	Lys	Ala 175	Lys
Val	Glu	Asn	Lys 180	Lys	Ala	Glu	Ala	Thr 185	Arg	Leu	Lys	Asn	Ile 190	Lys	Thr
Asp	Arg	Glu 195	Lys	Ala	Glu	Glu	Ala 200	Lys	Arg	Arg	Ala	As p 205	Ala	Lys	Leu
Gln	Glu 210	Ala	Asn	Val	Ala	Thr 215	Ser	Glu	Gln	Asp	Lys 220	Ser	Lys	Arg	Arg
Ala 225	Lys	Arg	Glu	Val	Leu 230	Gly	Glu	Leu	Ala	Thr 235	Pro	Asp	Lys	Lys	Glu 240
Asn	Asp	Ala	Lys	Ser 245	Ser	Asp	Ser	Ser	Val 250	Gly	Glu	Glu	Thr	Leu 255	Thr
Ser	Pro	Ser	Leu 260	Lys	Pro	Glu	Lys	Lys 265	Val	Ala	Glu	Ala	Glu 270	Lys	Lys
Val	Glu	Glu 275	Ala	Lys	Lys	Lys	Ala 280	Glu	Asp	Gln	Lys	Glu 285	Glu	Asp	Arg
Arg	Asn 290	Tyr	Pro	Thr	Asn	Thr 295	Tyr	Lys	Thr	Leu	Glu 300	Leu	Glu	Ile	Ala
Glu 305	Ser	Asp	Val	Glu	Val 310	Lys	Lys	Ala	Glu	Leu 315	Glu	Leu	Val	Lys	Glu 320
Glu	Ala	Lys	Glu	Ser 325	Arg	Asn	Glu	Glu	Lys 330	Ile	Lys	Gln	Val	Lys 335	Ala
Lys	Val	Glu	Ser 340	Lys	Lys	Ala	Glu	Ala 345	Thr	Arg	Leu	Glu	Asn 350	Ile	Lys
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Glu	Glu 370	Asp	Lys	Val	Lys	Glu 375	Lys	Pro	Ala	Glu	Gln 380	Pro	Gln	Pro	Ala
Pro 385	Ala	Pro	Gln	Pro	Glu 390	Lys	Pro	Thr	Glu	Glu 395	Pro	Glu	Asn	Pro	Ala 400
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18

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....

Leu Lys Pro Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Asp Lys Lys Ala Lys Ala Gln Lys Glu Glu Asp Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Glu Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Ala Glu Glu Pro Glu Asn Pro Val Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu <210> 16 <211> 414 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 16 Thr Glu Asn Glu Gly Ser Thr Gln Ala Ala Thr Ser Ser Asn Met Ala Lys Thr Glu His Arg Lys Ala Ala Lys Gln Val Val Asp Glu Tyr Ile Glu Lys Met Leu Arg Glu Ile Gln Leu Asp Arg Arg Lys His Thr Gln Asn Val Ala Leu Asn Ile Lys Leu Ser Ala Ile Lys Thr Lys Tyr Leu Arg Glu Leu Asn Val Leu Glu Glu Lys Ser Lys Asp Glu Leu Pro Ser Glu Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Lys Glu Lys Lys Asp

Thr Leu Lys Pro Gly Glu Lys Val Ala Glu Ala Lys Lys Val Glu Glu Ala Lys Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Phe Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn Glu Gly Thr Ile Lys Gln Ala Lys Glu Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Asp Ala Lys Leu Lys Glu Ala Asn Val Ala Thr Ser Asp Gln Gly Lys Pro Lys Gly Arg Ala Lys Arg Gly Val Pro Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Pro Ser Ser Leu Lys Ser Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Asp Leu Glu Ile Ala **9**5 Glu Ser Asp Val Lys Val Lys Glu Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Pro Arg Asp Glu Glu Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Asn Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Thr Gln Pro Glu Lys Pro Ala Pro Lys Pro Glu Lys Pro Ala Glu Gln Pro Lys Ala Glu Lys Thr Asp Asp Gln Gln Ala Glu Glu

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Glu Lys Lys Ala Lys Asp Gln Lys Glu Glu Asp His Arg Asn Tyr Pro Thr Ile Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Gly Ser Arg Asn Glu Glu Lys Val Lys Gln Ala Lys Ala Glu Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Gln Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Ala Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala Glu Glu <210> 18 <211> 406 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA from the genome of Streptococcus pneumoniae <400> 18 Thr Glu Asn Glu Gly Thr Thr Gln Ala Pro Thr Ser Ser Asn Arg Gly Asn Glu Ser Gln Ala Glu His Met Lys Ala Ala Lys Gln Val Asp Glu Tyr Ile Glu Lys Met Leu Gln Leu Asp Arg Arg Lys His Thr Gln Asn Val Gly Leu Leu Thr Lys Leu Gly Ala Ile Lys Thr Glu Tyr Leu Arg Gly Leu Ser Val Ser Lys Glu Lys Ser Thr Ala Glu Leu Pro Ser Glu Ile Lys Glu Lys Leu Thr Ala Ala Phe Lys Gln Phe Lys Lys Asp Thr Leu Lys Pro Glu Lys Lys Val Ala Glu Ala Glu Lys Lys Val Ala Glu Ala Lys Lys Ala Glu Asp Gln Lys Glu Glu Asp Arg Asn Tyr

Pro Thr Ile Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Val Lys Ala Asn Glu Pro Arg Asp Glu Glu Lys Ile Lys Gln Ala Glu Ala Glu Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Lys Lys Ile Lys Thr Asp Arg Glu Lys Ala Glu Glu Glu Ala Lys Arg Arg Val Asp Ala Lys Glu Gln Asp Glu Ser Ser Lys Arg Arg Lys Ser Arg Val Lys Arg Gly Asp Leu Gly Glu Gln Ala Thr Pro Asp Lys Lys Glu Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Pro Ser Pro Ser Leu Lys Pro Gly Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Glu Ala Asp Lys Lys Ala Lys Ala Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Asn Thr Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Pro Arg Asn Glu Glu Lys Val Lys Gln Ala Lys Ala Glu Val Glu Ser Lys Lys Ala Glu Ala Thr Arg Leu Glu Lys Ile Lys Thr Asp Arg Lys Lys Ala Glu Glu Ala Lys Arg Lys Ala Ala Glu Glu Asp Lys Val Lys Glu Lys Pro Ala Glu Gln Pro Lys Pro Ala Pro Ala Pro Gln Pro Glu Lys Pro Ala Pro Lys Pro Glu Asn Pro Ala Glu Gln Pro Lys Ala Glu Lys Pro Ala Asp Gln Gln Ala <210> 19 <211> 114 <212> PRT <213> Artificial Sequence

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Streptococcus pneumoniae

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Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asn 50	
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65 70 75 80 Ile Lys Ala Lys Leu Asp Ala Ala Phe Glu Gln Phe Lys Lys Asp Thr 90 95 85 Leu Lys Thr Glu Pro Gly 100 <210> 37 <211> 55 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from a cDNA consensus sequence from the genome of Streptococcus pneumoniae <400> 37 Asp Ala Lys Leu Glu Ala Thr Ser Glu Gln Asp Lys Pro Lys Gly Arg 1 5 10 15 Ala Lys Arg Gly Val Pro Gly Glu Leu Ala Thr Pro Asp Lys Lys Glu 20 25 Asn Asp Ala Lys Ser Ser Asp Ser Ser Val Gly Glu Glu Thr Leu Pro 35 40 45 Ser Pro Ser Leu Lys Pro Glu 50 55 <210> 38 <211> 103 <212> PRT <213> Artificial Sequence <220> <223> Description: Amino acid sequence derived from consensus sequence of CDNA derived from the genome of Streptococcus pneumoniae <400> 38 Lys Lys Val Ala Glu Ala Glu Lys Lys Val Glu Ala Lys Lys Lys 1 5 10 15 Ala Lys Asp Gln Lys Glu Glu Asp Arg Arg Asn Tyr Pro Thr Ile Thr 20 25 30 Tyr Lys Thr Leu Glu Leu Glu Ile Ala Glu Ser Asp Val Glu Val Lys 35 40 45 Lys Ala Glu Leu Glu Leu Val Lys Glu Glu Ala Lys Glu Ser Arg Asp 50 55 60 Glu Gly Lys Ile Lys Gln Ala Lys Ala Lys Val Glu Ser Lys Lys Ala 65 70 75 80 Glu Ala Thr Arg Leu Lys Lys Ile Lys Thr Asp Arg Glu Lys Ala Glu 85 90 95 Glu Glu Ala Lys Arg Arg Ala 100