(54) Title: STREPTOCOCCUS PNEUMONIAE CAPSULAR POLYSACCHARIDE GENES AND FLANKING REGIONS

Disclosed is the identification, cloning and sequencing of flanking DNA regions common to all polysaccharide capsule types in Streptococcus pneumoniae. Also disclosed are particular type-specific genes and gene products that direct the formation of the S. pneumoniae serotype-specific polysaccharide capsule. Methods are provided for detecting S. pneumoniae and for constructing gene cassettes that may be transferred as a unit during transformation and used to direct the expression of specific serotypes of S. pneumoniae capsules.
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DESCRIPTION

STREPTOCOCCUS PNEUMONIAE CAPSULAR
POLYSACCHARIDE GENES AND FLANKING REGIONS

BACKGROUND OF THE INVENTION

The present application is a continuation-in-part of co-pending U.S. Patent Application Serial No. 08/243,546, filed May 16, 1994. The entire text and figures of which disclosure is specifically incorporated herein by reference without disclaimer. The government owns certain rights in the present invention pursuant to grant number AI28457 from the Public Health Service and T32 AI07041-13 from the National Institutes of Health.

1. Field of the Invention

The present invention relates generally to the fields of bacterial capsule formation and the genes responsible for polysaccharide synthesis. More particularly, it concerns the genes and gene products that direct the formation of the Streptococcus pneumoniae serotype-specific polysaccharide capsule. The present invention also includes the identification of non-type specific gene sequences, flanking the capsule genes, and their use for the directed expression of specific serotypes of S. pneumoniae capsules.

2. Description of the Related Art

Infections due to S. pneumoniae are among the top ten causes of death in the United States. The normal populations most affected are young children and the elderly: pneumococcal pneumoniae, mainly affecting the elderly, causes >40,000 deaths per year among ~500,000
cases and represents 60 to 80% of all bacterial pneumonias; pneumococcal meningitis, with ~4000 cases/year, represents 11% of the total meningitis cases and has a fatality rate of >30% - greater than twice that of the two other leading causes, *N. meningitidis* and *H. influenzae*; bacteremia, usually following pneumonias or meningitis, accounts for >35,000 cases per year (>30% fatal); and otitis media, the most frequent reason for pediatric office visits after well-child care, is caused by *S. pneumoniae* in ~50% of cases (ACIP, 1981; ACIP, 1989; Austrian, 1984; Burke *et al.*, 1971; Center for Disease Control, 1978; Johnston and Sell, 1964; Koch and Dennison, 1974).

Other populations have an even higher incidence of pneumococcal infections: approximately 30% of sickle cell children will have severe pneumococcal infections in the first three years of life and ~35% of those will die (Overturf, *et al.*, 1977; Powars, *et al.*, 1981; Powars, 1975); in both adults and children with HIV infections, *S. pneumoniae* is the major cause of invasive bacterial respiratory disease (Janoff *et al.*, 1992). Patients with lymphomas, Hodgkins disease, multiple myeloma, splenectomy, and other debilitating diseases or immunologic deficiencies, are particularly susceptible to serious pneumococcal disease, as are those with chronic illnesses such as diabetes mellitus and heart disease. Furthermore, strains of *S. pneumoniae* are emerging that harbor resistances to multiple antibiotics, including penicillin (Appelbaum, 1992; Jacobs *et al.*, 1978; Landesman *et al.*, 1982).

The polysaccharide capsule of *S. pneumoniae* is the major virulence determinant of this organism. Despite early studies of the genetics, pathogenesis, and immunology of capsular polysaccharides, it remains unclear why certain capsular types appear to have a
greater capacity to cause disease. Of the more than 80 known capsular serotypes, 23 account for more than 90% of all pneumococcal infections.

In children, the most prevalent types are 3, 6, 14, 19, and 23, (Gray and Dillon, 1986), whereas in adults types 1, 3, 4, 6, 7, 8, 9, 12, 14, 18, 19, 23 prevail (Finland and Barnes, 1977). In assays of opsonophagocytosis (Branconier and Odeberg, 1982; Giebink et al., 1977; Knecht et al., 1970), complement activation and deposition (Fine, 1975; Gordon et al., 1986; Hostetter, 1986; Stephens et al., 1977; Winkelstein et al., 1980; Winkelstein et al., 1976), and mouse virulence (Briles et al., 1992; Briles et al., 1986; Knecht et al., 1970; MacLeod, 1965; Walter et al., 1941; Yother et al., 1982), levels of virulence have frequently been found to vary with the type of capsule expressed. For example, isolates expressing type 3, 4, and 19 capsules are highly resistant to phagocytosis, whereas those expressing types 6A, 14, 23 and 37 are significantly less resistant (Branconier and Odeberg, 1982; Hostetter, 1986; Knecht et al., 1970; Wood and Smith, 1949).

The importance of the capsule also results from the fact that anti-capsular antibodies are highly protective against infection. Nonetheless, the current polysaccharide-based vaccine is not particularly useful in some of the populations most affected by pneumococcal disease, e.g., the very young and the elderly, because of poor or absent immune response to polysaccharide antigens.

The ability to produce improved vaccines and therapies for pneumococcal infections will most likely be the result of a better understanding of the basic pathogenic mechanisms of the organism. This understanding necessarily includes the genetic basis for the expression
of serotype-specific polysaccharides and the role of capsular type per se in pathogenesis.

Some 85 different serotypes of Streptococcus pneumoniae, differing in the structure of the polysaccharide produced, have been identified (van Dam et al., 1990). The basis for the emergence of new capsule types remains obscure. Whether influenced by mutation, recombination, or immune selection, genetic exchange of DNA is likely to have played a major role in the evolution of capsule types. It is known that pneumococcal capsule types can be changed through genetic transformation in vitro (Dawson, 1930; Dawson and Sia, 1931; Langvad-Nielson, 1944; Avery et al., 1944).

Epidemiological studies suggest that a significant degree of genetic exchange occurs in vivo (Crain et al., 1990; Coffey et al., 1991; Versalovic et al., 1993). However, the mechanism by which capsule types are exchanged is not fully understood.

Extensive study was made of the genetics of capsular polysaccharide synthesis in S. pneumoniae using spontaneous mutants with defects in biosynthetic functions (Effrussi-Taylor, 1951; Ravin, 1960; Bernheimer and Wermundsen; 1972). The results of these studies indicated that the genes for polysaccharide synthesis were closely linked and could be transferred as a unit during genetic transformation. A cassette-type model of capsule type change based on this data has been proposed (Taylor, 1949; Austrian et al., 1959; Bernheimer and Wermundsen, 1972). According to the model, the type-specific genes for each capsule type would be present only in the genome of a strain of that capsule type and would show little homology to the type-specific genes of other capsule types. The type-specific genes would be located in homologous sites in the different chromosomes, clustered together between regions of highly
homologous flanking DNA. During transformation, recombination would occur in the flanking regions, resulting in the replacement of the recipient’s type-specific region by that of the donor.

The clustering of capsule biosynthetic genes proposed by the model is analogous to the organization that has been observed in the gram negative bacteria *Escherichia coli* (K antigens) (Roberts et al., 1988), *Neisseria meningitidis* (Frosch et al., 1989), and *Haemophilus influenzae* (Kroll et al., 1989). For each of these organisms, the type-specific region encoding biosynthetic functions (region 2) is flanked by highly homologous regions necessary for polysaccharide translocation (region 1) and modification (region 3). Since *H. influenzae*, like *S. pneumonias*, is naturally transformable, it has been proposed that capsule type change in this pathogen may occur by transformation with the type-specific gene cluster from a different serotype (Zwahlen et al., 1989).

The one exception to the cassette model of capsule type change in *S. pneumonias* is binary capsule formation. When non-encapsulated mutants have been transformed with chromosomal DNA from a strain of a different capsule type, most of the encapsulated transformants express the capsule type of the donor. However, at a frequency 10 to 100 times lower, encapsulated transformants are obtained which express both capsules (Bernheimer and Wermundsen, 1972). In some of these transformants, the second set of capsule genes is closely linked to the original set. However, these strains are unstable, and, at high frequency, lose the ability to produce the original capsule type. In binary strains in which the acquired capsule genes are unlinked to the original genes, binary capsule production is stable (Bernheimer and Wermundsen, 1969). Elucidation of the mechanism of binary capsule
type formation may be the key to understanding novel capsule type creation in *S. pneumoniae*.

It is clear that a better understanding of the genetics of capsular polysaccharide synthesis in *Streptococcus pneumoniae* is needed. The identification of type-specific capsular genes and the ability to transfer them, singly or as a gene cassette, to desired recipients, will elucidate the role of capsular types in virulence and allow easy identification of *S. pneumoniae* serotype. This ability will improve existing methods of diagnosis, identifying not only the presence of *S. pneumoniae* but also the capsular type of the invading strain. Furthermore, it will allow construction of strains producing elevated levels of capsular polysaccharides for improved vaccines.

**SUMMARY OF THE INVENTION**

20 Capsular Polysaccharide Genes and Flanking Regions

The present invention arises out of the discovery and sequence characterization of a gene family that confers on *S. pneumoniae* the ability to produce type-specific capsules that define the serotype of the organism. The inventors refer to this gene family as the capsule synthesis or *cps* genes. These genes encode the various enzymatic functions of capsule synthesis and determine the particular structure of the capsule polysaccharide that is produced, and thereby define serotype. These genes, designated *cpsB, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, tnpA*, and *pIPA*, map to specific DNA segments of sizes believed to range from about 0.5 kb to greater than 10 kb that appear to be type-specific for *S. pneumoniae*. Based upon the findings of the inventors, many type-specific genes may be distinguished on the
basis of restriction fragment length polymorphism (RFLP) analysis.

The present invention also includes the discovery and sequence characterization of non-type specific DNA regions that flank both sides of the cps locus. These flanking DNA segments can be used to identify the location of cps flanking DNA from any strain of S. pneumoniae. This invention thus provides the ability to identify the cps locus within all strains and allows for the subsequent isolation and characterization of all genetic elements involved in determining S. pneumoniae serotype.

The classification of S. pneumoniae strains is based on serological analysis of cell surface structures. 85 distinct serotypes have been identified to date based on the formation of surface molecules. The formation of the cell surface of S. pneumoniae, and in particular its polysaccharide capsule, has, until now, eluded characterization at the molecular genetic level. However, studies of the biosynthesis of the polysaccharide capsule have revealed that at least some of the genes are likely to include enzymes involved in the preparation of the sugar backbones for incorporation into the saccharide backbone, such as UDP-glucose dehydrogenase.

As mentioned above, these polymorphic "type-specific" sequence regions were found to be bounded or flanked by "non-type specific" regions having sequence elements that are apparently shared among the various subtypes. These regions, referred to as the left and right flanking regions, extend for, at least, 1 to 3 kb on either side of the cps genes. Thus, in type 3, the entire length of the capsule synthesis genes, including the non-type specific flanking regions, and any DNA
sequences in between, is greater than 9 kb. In other capsule types the length is on the order of about 5 to 20 kb, with the maximum length being related to the complexity of the polysaccharide encoded. Importantly, it is these flanking regions that allow recombination and integration of the type specific capsule genes to occur. Thus, when a selected cps gene or genes is positioned between the flanking regions, the resultant construct can be stably integrated into a S. pneumoniae host.

The present discoveries concerning the cps gene regions, the identification of conserved flanking regions, and the construction of erythromycin resistant insertions in adjacent, non-type specific DNA elements, allows for the changing of capsular serotypes by "cassette-in" the biosynthetic genes for different serotypes. This methodology may be employed for generating high yield capsular polysaccharide producing strains of different (heterologous) serotypes. For a high yielding strain, the existing serotype biosynthetic genes may be deleted and a different serotype's genes inserted ("cassetteted in"). These other serotypes may come from strains where their natural genetic background gives only poor or moderate levels of capsular production.

As used herein, the term "gene cassette" or simply "cassette", is intended to refer to any DNA segment flanked by one, or both, or part of, the cps flanking regions or a cps genetic element or DNA sequence which is found between the flanking regions.

Prior to the present invention, the foregoing underlying mechanism of genetic recombination of the capsule synthesis genes was unknown, as were the specific sequences involved. A principal contribution of the present inventors is the specific characterization of the
individual genes and flanking regions, allowing for their manipulation and individual transfer to hosts. Now, discrete nucleic acid segments, or cassettes, containing a cps flanking region, gene or genes, can be readily prepared and easily used in transformation.

Hybridization Probes and Primers

Accordingly, in certain embodiments the invention concerns nucleic acid segments that hybridize with cps genes and/or flanking regions. The nucleic acid segments will generally be less than about 20 kb in length, and preferably less than about 15 kb in length, or even 10 kb, and will comprise a non-type specific S. pneumoniae cps gene flanking region, and/or a type-specific cps gene, of sufficient length to allow hybridization with a pneumococcal cps flanking region and/or gene. Nucleic acid segments that are capable of hybridizing with the 5' flanking region, the 3' flanking region, to both flanking regions, to one or more of the genes designated cpsA, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, tnpA and plpA, and to one or more genes in combination with one or more flanking regions, are encompassed by the invention.

Nucleic acid segments that include a first sequence portion capable of hybridizing to the 5' cps gene flanking region and a second sequence portion capable of hybridizing with the 3' cps gene flanking region form one aspect of the invention. Such nucleic acid segments may be combined with one or more cps genes and may be constructed to form a genetic unit in which the gene (or genes) is located between the two flanking regions. The gene(s) may be from a different cps serotype to the flanking regions or they may be from the same cps serotype to the flanking regions. Such genetic units may be termed cassettes and may also encompasses the form of a circular DNA segment, plasmid, cosmid, or phage. An
isolated fragment of DNA containing DNA such as might be generated by restriction digestion, ligation, and/or PCR methodologies would also be included. The nucleic acid segment, or cassette, may also include other regions of DNA, such as restriction or cloning sites, PCR primers, promoters, antibiotic resistance genes, and the like, as necessary or desired to make a functional genetic unit.

In order to have utility in connection with the present invention, all that is required is that such a nucleic acid segment or genetic unit include a region of sufficient complementarity and size to allow selective cross-hybridization with the target flanking region or gene sequence.

In general, shorter and intermediate length nucleic acid fragments will be useful as hybridization probes and primers, and in particular, for use in PCR, where the primers will allow generation of the entire intervening cps sequence.

Thus flanking region and gene fragments on the order of at least about 14-15, 20, 30, 40, 50, or 100 to 200, contiguous complementary nucleotides are contemplated, although sequences of 500, 1,000, or more, nucleotides in length may also be used. The DNA segments may, of course, be of any length within the stated ranges. This is the meaning of "about" in this context, with "about" meaning a range longer or shorter than the stated length, extending to the previously quoted longer and shorter lengths (with about 14 or so still being the minimum length). The ranges thus encompass 1 to 4, 1 to 9, 1 to 49, 1 to 99, and the like, nucleotides in length.

Longer nucleic acid segments and fragments having on the order of up to 1,000, 2,000, 3,000, 5,000, 10,000, 15,000 or longer in length will also have particular
utilities in addition to their function in hybridization embodiments. In particular, longer nucleic acid segments and fragments including selected cps coding sequences may be employed in the expression of recombinant proteins, and nucleic acid segments that include one or more cps gene sequences positioned between the flanking regions (cassette constructs) may be used in gene transfer embodiments as described above. The DNA segments may, of course, be of any length within the ranges stated above, so long as they function to achieve the desired effect. "About" in this context therefore indicates ranges of from 1 to 999, or 1 to 4,999, and the like, nucleotides longer or shorter than the stated length.

Nucleic Acid and Amino Acid Sequences

Exemplary flanking regions sequences, as disclosed herein, are set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4 and SEQ ID NO:6 (FIG. 7 and FIG. 8). The 5’ cps gene flanking region is represented by SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3 and SEQ ID NO:4. SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3 corresponds to regions of DNA sequenced in the upstream portion of the 5’ flanking region and SEQ ID NO:4 corresponds to the downstream portion of the 5’ flanking region and is termed the "repeat" region. DNA between SEQ ID NO: 1, SEQ ID NO:2, SEQ ID NO:3 and SEQ ID NO:4 are also part of the cps 5’ flanking region. SEQ ID NO:1 is 300 nucleotides in length and corresponds to the last 180 nucleotides of cpsB and the 5’ end of cpsC which begins immediately (FIG. 6A and FIG. 7). SEQ ID NO:2 is 261 nucleotides in length and corresponds to a 3’ end region of cpsC (FIG. 6B and FIG. 7). SEQ ID NO:3 is 262 nucleotides in length and corresponds to part of cpsE and the 5’ end of the repeat region (FIG. 6C and FIG. 7). SEQ ID NO:4 is 934 nucleotides in length, (nucleotide 1 through 934, FIG. 6D and FIG. 8). The 3’ cps gene
The inventors have found that certain *S. pneumoniae* nucleotide sequences described in the scientific literature correspond to stretches of sequences from the flanking regions of the present invention. For example, Guidolin et al. (1994) have sequenced 6,322 base pairs of the 19F *S. pneumoniae* serotype cps locus. Sequence analysis indicated that this region contained six complete open reading frames and one partial, which they named cps19fA to cps19fG. Southern hybridization revealed that cps19fA and cps19fB were conserved in 12 other *S. pneumoniae* serotypes tested, including serotype 3. cps19fC and cps19fD also hybridized to Type 3 *S. pneumoniae*. The sequences for cpsB, cpsC and cpsE (SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3), as disclosed herein, are about 99% identical to cps19fB, cps19fC and cps19fD respectively (Guidolin et al. 1994). However, cpsE is truncated at the 3' end with respect to the 19F gene (lacks ~280 nt). The site of the truncation is adjacent to the region designated as the "repeat" sequence (SEQ ID NO:4). Based on blotting data (see Example 17) part of the repetitive element is in SEQ ID NO:3. This sequence extends 132 nt upstream of the SacI site at the start of SEQ ID NO:4, as shown in FIG. 7. Although Guidolin et al. (1994) have sequenced this area in serotype 19f, they do not identify this sequence as being a gene flanking region and do not suggest its use as part of an *S. pneumoniae* capsular cassette.

Garcia et al. (1993) localized a 781 bp EcoRV subfragment of a gene (cap3-1) that they proposed was involved in the transformation to a capsulated phenotype. The first 52 nucleotides of the 781 Garcia et al. sequence correspond to nucleotides 883 to 934 of SEQ ID
NO:4. The next 443 nucleotides of the Garcia et al. sequence corresponds to nucleotides 1 to 443 of SEQ ID NO:5. However, the remaining 286 nucleotides of the 781 bp sequence do not show any homology with the instant sequences. As with the above, this article does not identify this sequence as being a gene flanking region and do not suggest its use as part of an S. pneumoniae capsular cassette. However, the same group (Arrecubieta et al., 1994) using probes in Southern blotting, identified the upstream region of their cap3A as a sequence common to all pneumoniae strains tested.

The amino acid sequences described by Garcia et al. (1993) include a putative protein of 138 amino acids (CAP3-1), transcribed in the same direction as CpsD, and another truncated open reading frame, transcribed in the opposite direction. The first 117 amino acid residues of CAP3-1 corresponds to the first 117 residues of CpsD (SEQ ID NO:11). CAP3-1 is reportedly similar to the amino-terminus of GDP-mannose dehydrogenase of Pseudomonas aeruginosa. The remaining 21 residues of CAP3-1 described by Garcia et al. (1993) have no homology with the remaining 277 amino acid residues of CpsD, suggesting that Garcia et al. had cloned a truncated gene in which the 5' end of the cpsD gene was aberrantly fused to an unidentified DNA sequence.

Pearce et al. (1994) described an S. pneumoniae protein and corresponding nucleotide sequence, that they proposed was a permease-like protein involved in peptide transport. They termed this protein PlpA. Nucleotides 484 to 1307 of SEQ ID NO:6 correspond to nucleotides 843 to 1672 of the plpA described by Pearce et al. (1994), and the amino acid SEQ ID NO:15 corresponds to residues 282 to 557 of their PlpA. Previously, Pearce et al. (1993) had described an S. pneumoniae protein they called Expl. The amino acid sequence of Expl corresponds to
residues 49 through 209 of SEQ ID NO:15. The inventors have found that the 'p1pA located adjacent to the type 3-specific genes lacks about one third of its 5' end when compared to p1pA genes located adjacent of the capsule genes of other types such as that used by Pearce et al. (1994). Neither Pearce et al. (1994) nor Pearce et al. (1993) identify this sequence as being involved in capsule synthesis in any way, nor do they suggest that it forms part of a common DNA flanking region.

Although certain stretches of nucleotide sequences may have been known in the art, their function, relationship to capsule synthesis and, particularly, their role as interchangeable flanking regions has not previously been described. An important feature of the invention is that the functional characterization of the flanking regions allows, for the first time, for the exchange of S. pneumoniae type-specific capsule genes to be manipulated and controlled. This is only possible in light of the inventors discovery of the conserved cps gene flanking regions. Nucleic acid segments, including cassettes and plasmids, that include both 5' flanking region sequences and 3' flanking region sequences are thus one preferred embodiment of the invention. PCR primers that have sequences corresponding to both flanking regions form another preferred embodiment of the invention.

Encoded within the upstream 5' flanking region (SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3) are the partially sequenced genes cpsB, cpsC and cpsE which encode for CpsB (SEQ ID NO:7), CpsC (SEQ ID NO:8 and SEQ ID NO:9) and CpsE (SEQ ID NO:10) (FIG. 6A, FIG. 6B, FIG. 6C and FIG. 7).

The invention also includes other cps gene sequences, either alone or in combination with the
flanking region sequences described above. The type-specific portions of the polycistronic cps gene locus operon, as disclosed herein, are encompassed within nucleotides 1 through 4951, SEQ ID NO:5 (nucleotides 935 through 5885, FIG. 6E, FIG. 6F, FIG. 6G, FIG. 6H, FIG. 7 and FIG. 8). Beside the open reading frames for the proteins, the sequences also contain putative promoters that direct the transcription of the genes of the cps locus. Other promoters, herein termed "recombinant promoters", may also be used to direct the expression of the cps genes in accordance with the invention.

The genes encoded within SEQ ID NO:5 and as disclosed herein, include *cpsD, cpsS, cpsU, cpsM* and part of *tnpA*.

*cpsD* is 1277 nucleotides in length, 1 through 1277 of SEQ ID NO:5 (935 through 2211, FIG. 6E and FIG. 8).

*cpsS* is 1267 nucleotides in length, 1277 through 2543 of SEQ ID NO:5 (2211 through 3477, FIG. 6F and FIG. 8).

*cpsU* is 1055 nucleotides in length, 2707 through 3761 of SEQ ID NO:5 (3641 through 4695, FIG. 6G and FIG. 8).

*cpsM* is 1194 nucleotides in length, 3758 through 4951 of SEQ ID NO:5 (4692 through 5885, FIG. 6H and FIG. 8).

CpsS is just downstream of CpsD, only 15 nucleotides separate a potential start codon for CpsS from the stop codon of CpsD. Other start codons for CpsS are at nucleotides 1311 and 1355 (SEQ ID NO:5). There is a large non-coding region between *cpsS* and *cpsU* (nucleotide 2543 through 2707, SEQ ID NO:5). Where as *cpsD* and *cpsS*
overlap by one nucleotide and cpsU and cpsM overlap by 4
nucleotides. All four genes are transcribed in the same
direction and cpsD and cpsS are in the same reading
frame.

Encoded within the 3' flanking region (SEQ ID NO:6)
is the truncated sequence for plpA, designated 'plpA,
which is 823 nucleotides in length, 484 through 1307 of
SEQ ID NO:6 (nucleotides 6370 through 7192, FIG. 6I and
FIG. 8). As mentioned above the 5' end is not present in
the plpA gene of type 3 S. pneumoniae. A partial
transposase sequence, tnpA, is contained between cpsM and
'plpA. It is transcribed in the opposite orientation to
all other genes described herein, and extends from
nucleotide 480 through 1, SEQ ID NO:6 to overlap with the
cpsM gene nucleotide 4951 through 4902, SEQ ID NO:5 a
total of 531 nucleotides (nucleotides 5836 through 6366,
FIG. 6J and FIG. 8).

cpsD encodes a 394 amino acid sequence (SEQ ID
NO:11) which is homologous to that of the UDP-glucose
dehydrogenase (HasB) from Streptococcus pyogenes. The
deduced amino acid sequence encoded by cpsS predicts a
protein of 416 residues (SEQ ID NO:12) which is
homologous to polysaccharide synthases. cpsU encodes a
306 amino acid sequence (SEQ ID NO:13) which is
homologous to glucose-1-phosphate uridylyltransferases
from several other bacterial species. cpsM encodes a 397
amino acid sequence (SEQ ID NO:14) which has homology
with phosphoglucomutases from several bacterial species.
However, it lacks approximately 25% of the C-terminal
present in other phosphomutases and may not encode a
functional protein. 'plpA encodes a 274 amino acid
sequence (SEQ ID NO:15) and tnpA encodes a 177 amino acid
sequence (SEQ ID NO:16).
Thus, in certain particular embodiments the present invention concerns the individual *cps* genes, including segments encoding sequences corresponding to one or more of the *cpsB*, *cpsC*, *cpsE*, *cpsD*, *cpsS*, *cpsU* and *cpsM* genes. *tnpA* and *plpA* gene constructs, when combined with other *cps* genes and flanking regions, are also encompassed by the invention. In further embodiments, the invention concerns the respective proteins and polypeptides encoded by the *cpsB*, *cpsC*, *cpsE*, *cpsD*, *cpsS*, *cpsU*, *cpsM*, *tnpA* and *plpA* genes. The proteins, polypeptides and peptides of the invention may be used in a variety of embodiments, including, e.g., in immunological protocols to generate antibodies that may, in turn, be used in diagnostic embodiments to detect *S. pneumoniae*.

It should be noted that in the definition of the genes and proteins, the term "*cps*" is not used to indicate that the gene or protein concerned has a defined role in capsule synthesis in all cases. Rather, it indicates that the gene is located between the *cps* gene flanking regions, i.e., within the *cps* operon, and in close association with other *cps* genes. It should also be noted that, the "*S. pneumoniae* gene region" refers to all genetic elements associated with the *cps* genes, including genes incorporated within the flanking regions. A "genetic element" refers to any DNA that may encode for a protein or polypeptide, regardless of functionality. The utility of the *cps* genes remains that, e.g., they may be used in the same diagnostic manner to identify *S. pneumoniae*.

While the present disclosure is exemplified in part through the cloning and sequencing of type 3 *cps* genes, the techniques are equally applicable to the *cps* genes of other capsule serotypes, including any one of the 85 serotypes. For example, using the techniques developed for the characterization of the type 3 *cps* gene region,
the inventors have proceeded to characterize the restriction maps for the type 2 type 6B strains (Example 16, FIG. 11). As expected, the maps of the non-type specific flanking regions were found to be identical from serotype to serotype, whereas the maps for the cps gene regions themselves were serotype specific.

Diagnostic Embodiments

The cps flanking region and gene sequences, and the encoded proteins, may be employed in diagnostic embodiments. For example, the amount of S. pneumoniae, or S. pneumoniae serotype, present within a biological sample, such as blood, serum or a swab from nose, ear or throat, may be determined by means of a molecular biological assay to determine the level of nucleic acid complementary to the cps loci, or even by means of an immunoassay to determine the level of one of the polypeptides encoded by a gene from this locus. The cps locus DNA segment used in molecular biological assays may include the non-type specific segments such as the 5' and 3' flanking regions, SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:6 and any sequence in between, or the region encoding various polypeptides, such as those incorporated within SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, ID NO:5 and SEQ ID NO:6.

In a molecular biological method for detecting S. pneumoniae, one would obtain nucleic acids from a suitable sample and analyze the nucleic acids to identify a specific nucleic acid segment complementary to the cps loci (whether type- or non-type-specific). The nucleic acid segment will generally be identified by sequence, which method generally includes either; identifying a transcript with a corresponding or complementary sequence, e.g., by Northern or Southern blotting using an appropriate probe or; identifying a transcript with two
or more shorter primers and amplifying with PCR technology.

**Blotting Techniques**

To detect *S. pneumoniae*, as may be used to identify a patient with otitis media, pneumococcal pneumonia or pneumococcal meningitis, using a method based upon hybridization and blotting techniques, one may use a probe with a sequence as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 or SEQ ID NO:6, including any sequence in between, or an equivalent thereof. This imparts an evident utility to the nucleic acid segments of the present invention.

To conduct such a diagnostic method, one would generally obtain sample nucleic acids from the sample and contact the sample nucleic acids with a nucleic acid segment corresponding to the *cps* loci disclosed herein, under conditions effective to allow hybridization of substantially complementary nucleic acids, and then detect the presence of any hybridized substantially complementary nucleic acid complexes that formed.

The presence of a substantially complementary nucleic acid sequence in a sample, or a significantly increased level of such a sequence, in comparison to the levels in a normal or "control" sample, will thus be indicative of a sample that harbors *S. pneumoniae*. Here, substantially complementary nucleic acid sequences are those that have relatively little sequence divergence and that are capable of hybridizing to the sequences disclosed herein under standard conditions.

Where a substantially complementary nucleic acid sequence, or a significantly increased level thereof, is detected in a clinical sample from a patient suspected of
having otitis media, pneumococcal pneumonia or pneumococcal meningitis, for example, this will be indicative of a patient that does have that particular disease. As used herein, the term "increased levels" is used to describe a significant increase in the amount of the cps loci nucleic acids detected in a given sample in comparison to that observed in a control sample, e.g., an equivalent sample from a normal healthy subject.

A variety of hybridization techniques and systems are known that can be used in connection with the S. pneumonae detection aspects of the invention, including diagnostic assays such as those described in Falkow et al., U.S. Patent 4,358,535.

In general, the "detection" of a cps locus is accomplished by attaching or incorporating a detectable label into the nucleic acid segment used as a probe and "contacting" a sample with the labeled probe. In such processes, an effective amount of a nucleic acid segment that comprises a detectable label (a probe), is brought into direct juxtaposition with a composition containing target nucleic acids. Hybridized nucleic acid complexes may then be identified by detecting the presence of the label, for example, by detecting a radio, enzymatic, fluorescent, or even chemiluminescent label.

Where one simply desires to distinguish S. pneumonae DNA from the DNA of other bacteria, it is contemplated that the non-type specific region sequences may be employed as probes. However, where one desires to distinguish among different S. pneumonae serotypes, it is contemplated that probes will include both type specific and non-type specific cps sequences. The type-specific sequences, being type specific, will selectively hybridize only to corresponding serotypes. Thus one can envision a battery of serotype specific cps nucleic acid
hybridization probes can be employed to distinguish and identify serotype DNA samples. In these instances, it is not believed to be necessary to employ restriction enzyme digestion prior to hybridization, but this can be employed where desired. Alternatively, only one non-type specific sequences may be employed as a "universal probe" that allows detection of restriction fragment length polymorphisms (RFLPs). Typically for RFLP detection, one will employ the more specific hybridization technique of Southern analysis wherein restriction digestion of the unknown or target DNA is carried out using an enzyme that will cleave either within or surrounding the cps gene region (FIG. 4 and FIG. 11) show restriction maps for several of the serotype cps gene regions).

Many suitable variations of hybridization technology are available for use in the detection of nucleic acids, as will be known to those of skill in the art. These include, for example, in situ hybridization, Southern blotting and Northern blotting. In situ hybridization describes the techniques wherein the target nucleic acids contacted with the probe sequences are those located within one or more cells, such as cells within a clinical sample or even cells grown in tissue culture. As is well known in the art, the cells are prepared for hybridization by fixation, e.g. chemical fixation, and placed in conditions that allow for the hybridization of a detectable probe with nucleic acids located within the fixed cell.

Alternatively, target nucleic acids may be separated from a cell or clinical sample prior to contact with a probe. Any of the wide variety of methods for isolating target nucleic acids may be employed, such as cesium chloride gradient centrifugation, chromatography (e.g., ion, affinity, magnetic), phenol extraction and the like. Most often, the isolated nucleic acids will be separated,
e.g., by size, using electrophoretic separation, followed by immobilization onto a solid matrix, prior to contact with the labelled probe. These prior separation techniques are frequently employed in the art and are generally encompassed by the terms "Southern blotting" and "Northern blotting". Although the execution of various techniques using labeled probes to detect cps locus DNA or RNA sequences in clinical samples are well known to those of skill in the art, a particularly preferred method is described in detail herein, in Example 4.

**PCR Techniques**

To detect *S. pneumoniae*, using a method based upon PCR technology of U.S. Patent 4,603,102 (incorporated herein by reference), one may also use one or more probes with a sequence as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3 and SEQ ID NO:4 (including sequence in between), SEQ ID NO:5 or SEQ ID NO:6, or an equivalent thereof.

To conduct such a diagnostic method, one would generally obtain sample nucleic acids from a suitable source and contact the sample nucleic acids with two probes or primers corresponding to the cps loci disclosed herein, under conditions which allow for hybridization and polymerization to occur. A pair of probes, one corresponding to the 5' flanking region and the other corresponding to the 3' flanking region, would be sufficient to detect the presence of *S. pneumoniae* in a sample and may even be used to indicate the amount of bacteria present. Furthermore the size of the isolated PCR product, when separated by any of the methods as described above, may be sufficient to identify the *S. pneumoniae* serotype. Alternatively the PCR product may be digested with one or more restriction enzymes, which
would enable individual serotypes to be distinguished, using the same principle as RFLP.

In further embodiments, it may be desired to employ other probes corresponding to type specific or non-type specific regions. A battery of serotype specific probes, probes corresponding to type specific DNA regions, may be employed in individual reactions, with a universal probe, a probe corresponding to non-type specific regions. The size of PCR products, with or without prior digestion with restriction enzymes, would distinguish and identify the *S. pneumoniae* serotype.

**Kits**

Kits for use in Southern and Northern blotting or PCR technology, to identify *S. pneumoniae* and/or individuals having, or at risk for developing, otitis media, pneumococcal pneumonia or pneumococcal meningitis, are also contemplated to fall within the scope of the present invention. Such kits will comprise, in suitable container means, cps nucleic acid probes; also, generally, unrelated probes for use as controls; and optionally, one or more restriction enzymes.

**Characterization of Streptococcus pneumoniae Serotypes**

In another embodiment of the invention, the non-type specific DNA may be used, to isolate and characterize the type-specific DNA sequence for all or any strain of *S. pneumoniae*. Consequently, the most suitable probes for diagnosing *S. pneumoniae* infection, for use in any molecular biological technique, could be found.

The present invention identifies the common flanking DNA which may be used in hybridizations to target the location of the type-specific *cps* genes for any strain of
S. pneumoniae. Once this location has been identified
the cps genes may then be isolated and characterized by
the use of conventional techniques, such as; chromosome
crawling, PCR technology, cloning and DNA sequencing, all
are disclosed herein. As mentioned above, this in turn
would enable suitable probes from each S. pneumoniae
strain to be chosen and then used for, diagnostic and
research purposes. Furthermore, the genetic elements
involved in determining S. pneumoniae serotype may be
elucidated, and an understanding of their effect on
virulence and evolutionary role may be achieved.

In still more particular embodiments, the invention
concerns the preparation and cloning of entire cps gene
regions encoding one or more specific cps genes of a
particular serotype, positioned within a "cassette" for
ease of manipulation, e.g. in plasmid preparation or host
cell introduction, etc. Thus, cps gene cassettes in
accordance with the present invention will typically
include left and right hand flanking regions to allow
homologous recombination in S. pneumoniae host cells.

A preferred method for preparing cps gene cassette
is through the application of PCR technology wherein left
and right hand primer corresponding to left and right
hand flanking region are employed to amplify the cps gene
coding regions. Of course, the primers employed will
typically include at their termini appropriate
restriction enzyme site. Thus, the resulting cassette
will preferably include at each terminus a restriction
enzyme site of choice. The site, of course, will depend
upon the vector that is ultimately employed for
manipulation or transformation but may be the specific
cleavage site for one or more of the restriction enzymes
shown in FIG. 4 and FIG. 7 or listed in Table 1. The
list is, of course, exemplary, and any restriction enzyme
could be employed, as is well known to those of skill in the art.

The present invention also contemplates more traditional approaches to cps gene regions cloning, such as by partial fragmentation of S. pneumoniae DNA followed by cloning into a recombinant cloning host, such as E. coli, and screening by hybridization and antibiotic selection (using a selection marker found on the plasmid or other vector employed for cloning). Of course, if the cloning host is not a S. pneumoniae host, one may employ either type specific or non-type specific cps gene sequences for screening. In these cases, cassettes that are developed may include enzyme restriction sites naturally found to occur in the flanking regions, such as an SphI site.

It is contemplated that virtually any type of host cell may be employed in connection with the present invention, depending on the particular application. For example, where one simply desires to manipulate cps gene sequences, any acceptable host may be employed, such as an E. coli, or even an appropriate eukaryotic host where desired. However, where one contemplated producing capsule polysaccharides, one will desire to employ a gram positive host such as bacillus, staphylococcus or streptococcal hosts. Particularly preferred will be S. pneumoniae host, in that it is contemplated that such hosts will be more readily amenable to manipulation to increase capsular polysaccharide production.

The inventors contemplate that a particular application, therefore, will be the use of recombinant hosts not only for the preparation and manipulation of cps gene sequences, but also in the large scale production of capsule saccharides and polysaccharides. These polysaccharides are useful for the production of
antigenic haptens and epitopes for use in the production of immunogens. Haptens and epitopes are described herein as portions of a molecule against which an immune response is directed. In conjunction with an molecule that elicits an immune response, that is an immunogen, an hapten-immunogen complex is able to elicit an immune response.

Generally speaking, where capsule production is required, one will employ a *S. pneumoniae* host cell into which a selected *cps* gene region is introduced or "cassetted in". First, a DNA segment that includes the selected *S. pneumoniae* *cps* gene(s) flanked by sufficient *S. pneumoniae* flanking regions to allow homologous recombination in the *S. pneumoniae* host is identified. It is contemplated that flanking regions on the order of 0.1 to 1 kb will be sufficient to allow recombination to occur. Once an appropriate DNA segment is introduced into the *S. pneumoniae* host, either as genomic DNA or as a recombinant vector (plasmid), transformed host expressing the introduced *cps* gene are selected.

DNA may be introduced into a suitable host by a variety of mechanisms, including natural transformation of *S. pneumoniae*, calcium mediated transformation or electroporation of *E. coli*. A particularly preferred method of bacterial transformation includes the steps of making an *S. pneumoniae* competent for transformation by growth in Todd Hewitt broth supplemented with calcium and bovine serum albumin. Alternatively electroporation may be employed, one makes the bacterial cells, such as the *E. coli* strain LE392, competent in 10% glycerol in water, adds the DNA, and electroporates the cells.

Where the *cps* gene DNA segment is introduced in the form of a plasmid, it would be preferable, at certain times, to employ a plasmid that is free of a *S.*
pneumoniae origin of replication. Thus, virtually any traditional plasmid (which are designed, e.g. for gram negative hosts such as E. coli) may be employed. The reason is that where the plasmid is free of a S. pneumoniae origin of replication only those clones that have successfully undergone homologous recombination with the recombinant cps gene region will be detected. Stated another way, in this case, there is no requirement of a S. pneumoniae origin of replication in order for homologous recombination to occur, and thus homologous recombinants are inherently selected for using such a cloning technique.

Alternatively it may be simpler to introduce the cassette into S. pneumoniae on a replicating plasmid with a S. pneumoniae origin of replication. In this way, higher levels of polysaccharide production as a result of the elevated copy number of the plasmid (10 to 20) as opposed to the low copy number of the chromosome (1 to 2) would be achieved and homologous recombination need not occur.

Additionally, it is contemplated that there will be some advantage to employing as the starting host a S. pneumoniae strain that is a high producer of its own inherent cps gene. These high producers will necessarily include the genetic environment to support high production of the newly introduced cps complex, and thus will likely be ideal hosts for such production. To achieve such recombinants, all that is required is that the heterologous gene containing the flanking regions, or a genome containing the flanking regions, is introduced into the host cell, and resultant recombinants wherein the homologous gene has been replaced is selected.

Although the invention contemplates in particular embodiments the introduction of an entire recombinant cps
gene region where capsule saccharide synthesis is desired, the invention also contemplates the introduction of one, two, three or so of the individual cps genes. It is contemplated that one or two genes, such as those that control the biosynthesis of sugar precursors may be sufficient to, in and of themselves, confer serotype specific saccharides production on the selected host, or can in of themselves, upregulate capsule production by existing cps genes of the host.

**BRIEF DESCRIPTION OF THE DRAWINGS**

The following drawings form part of the present specification and are included to further demonstrate certain aspects of the present invention. The invention may be better understood by reference to one or more of these drawings in combination with the detailed description of specific embodiments presented herein.

**FIG. 1.** ELISA for the detection of type 3 capsule. Wells of microtiter dishes were coated with crude extracts of capsule material. Type 3 capsule was detected with the monoclonal antibody 16.3 (Briles et al., 1981a). The type 2 strain D39 served as a negative control. Measurements were made in triplicate, and error bars represent standard errors. Values were standardized to protein content. Genotypic designations were based on linkage as determined by transformation mapping.

**FIG. 2.** Insertion-duplication restoration. The cloned fragment and the homologous fragment in the chromosome are represented by the open boxed regions. The dark block represents the mutation in the chromosome of the mutant strain. Insertion of the plasmid clone into the pneumococcal chromosome results in a duplication of the homologous fragment with the plasmid inserted in between. If the recombination occurs to the left of the
mutation as shown here, a wild type, full-length copy of the gene is created and function is restored. The plasmid clone leading to restoration spontaneously excises at low frequency through homologous recombination and can therefore be easily recovered by transformation to E. coli. pJD330 contains a 2.4 kb Sau3AI fragment.

**FIG. 3A.** Repair of capsule mutations by double crossover recombination event. Mutants were transformed with plasmid subclones of pJD330 and no selection for Ery$^R$ was made. The box in the chromosome represents the region in the mutant chromosome homologous to the fragment cloned in pJD330. The plasmid represents a subclone of pJD330 capable of restoring encapsulation in the mutant strain.

**FIG. 3B.** Deletion analysis to locate the site of the cpsA1, capD4, and Rx1 mutations. The mutations were mapped by transformation with plasmid clones containing the indicated fragments and screening for the mucoid colony phenotype. Identical fragments repaired the cpsA1, capD4, and Rx1 mutations. No selection was made for insertion of the plasmids, thus these numbers represent double crossover events. The actual frequencies of repair, shown for the cpsA1-containing mutant, are mainly a reflection of the transformation efficiency of the recipient. No encapsulated transformants were obtained when pJY4163 (no insert) was used for transformation. Restriction sites are: M, MunI; P, PstI; P, PvuII; R, RsaI; S3, Sau3A I; Ss, SspI; Xb, XbaI.

**FIG. 4.** Physical and genetic map of the type 3 capsule region of *S. pneumoniae* WU2. The restriction map was developed by probing chromosomal digests of WU2 with pJD330 and pJD366 and by probing chromosomal digests of JD770 with pJD330 and pJY4163. The location of primary
clones pJD330 and pJD366 are indicated above the map. Subcloned fragments used to target insertion-duplication mutations are indicated below the map. Insertion of plasmids containing shaded fragments led to loss of capsule production. + or - at the end of a fragment indicates the presence or absence of transcription detected at that point in the chromosome. Insertions of pJD330 and pJD366 are in the orientation to detect transcription to the left. Clones pJD351 and pJD364, which contain the pJD330 and pJD366 fragments, respectively, in the opposite orientation, were also used for the transcription studies. The other plasmids used for insertions were pJD356, pJD337, pJD369, pJD359, pJD362, pJD361, pJD357, and pJD374, in the order shown.


**FIG. 5.** Schematic representation of the capsule regions in these strains. Insertions in JD871 and JD872 result from incorporation of pJD366. The insertion in JD803 is pJD330. The shaded square symbol represents type 2 specific DNA; the open square symbol represents type 3 specific DNA; the hatched square symbol represents flanking DNA common to both type 2 and type 3 and; the black square symbol represents pJY4163 or pJY4164. The locations of the probes used are indicated below the map.

**FIG. 6A.** DNA sequence of the 5' flanking region including partial sequence of \textit{cps3B} and \textit{cps3C} (SEQ ID NO:1) encoding for SEQ ID NO:7 and SEQ ID NO:8.
FIG. 6B. DNA sequence of the 5' flanking region including partial sequence of cps3C (SEQ ID NO:2), encoding for SEQ ID NO:9.

FIG. 6C. DNA sequence of the 5' flanking region including partial sequence of cps3E (SEQ ID NO:3), encoding for SEQ ID NO:10.

FIG. 6D. DNA sequence of the "repeat" upstream flanking DNA (SEQ ID NO:4).

FIG. 6E. DNA sequence of the region containing cps3D (nucleotides 1 through 1277, SEQ ID NO:5). The -35 and -10 hexamers of potential σ-70 type promoters upstream of cps3D are underlined and labeled above the sequence. Putative ribosome binding sites are underlined. The precise locations of endpoints of insertion mutations shown in FIG. 7 are indicated by triangles below the sequence and are labeled with the name of the strain containing the given mutation. The locations of spontaneous mutations in cps3D are labeled with the sequence of the mutation and the name of the strain containing the mutation. The sequence from the PvuII-SspI fragment of A66R2 began at nucleotide 1921, thus it is possible that additional mutations are present from the PvuII site to this point. Selected restriction sites are shown.

FIG. 6F. DNA sequence of the region containing cps3S (nucleotides 1277 through 2543, SEQ ID NO:5). The precise locations of endpoints of insertion mutations shown in FIG. 7 are indicated by triangles below the sequence and are labeled with the name of the strain containing the given mutation.

FIG. 6G. DNA sequence of the region containing cps3U (nucleotides 2707 through 3806, SEQ ID NO:5). The
-35 and -10 hexamers of potential σ-70 type promoters upstream of cps3U are underlined and labeled above the sequence. A short region of dyad symmetry just upstream of cps3U is overlined. Putative ribosome binding sites are underlined. The precise locations of endpoints of insertion mutations shown in FIG. 7 are indicated by triangles below the sequence and are labeled with the name of the strain containing the given mutation. Selected restriction sites are shown.

FIG. 6H. DNA sequence of the region containing cps3M (3746 through 4951, SEQ ID NO:5) with corresponding amino acid sequences. Note that the first line of FIG. 6H overlaps the last line of FIG. 6G.

FIG. 6I. DNA sequence of the region containing the 3' flanking region including 'pIpa (SEQ ID NO:6) with corresponding amino acid sequences (SEQ ID NO:15).

FIG. 6J. The DNA sequence for a partial transposase A, tnpA, located between 'pIpa and overlapping cpsM. The open reading frame is in the opposite orientation starting at nucleotide 6366 through 5836 of FIG. 6I.

FIG. 7. Map of the chromosomal region involved in the biosynthesis of S. pneumoniae type 3 capsular polysaccharide. Triangles indicate the endpoints of insertion mutations. Filled triangles represent insertions which resulted in loss of capsule production. Open triangles represent mutations which did not result in loss of capsule production. Restriction enzyme sites are: Bg, BglII; Ev, EcoRV; H, HindIII; P, PstI; Pv, PvuII; S, SacI; Sa, SalI; Sp, Sphi. Also included is a diagram showing the position of SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3, including the corresponding amino acid sequences SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9 and SEQ ID NO:10.
FIG. 8. Map of the chromosomal region involved in the biosynthesis of \textit{S. pneumoniae} type 3 capsular polysaccharide showing the relationship between SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6, and DNA sequence as described in FIG. 6D through FIG. 6J.

FIG. 9. Location of insertion mutations in the type 3-specific region of the \textit{S. pneumoniae} WU2 chromosome. Schematic illustration of the insertions. The schematic was derived from Southern blot analysis such as that shown in FIG. 11a and FIG. 11b. The ability of the strains to produce type 3 capsule is indicated. Restriction sites are: F, FspI; H, HindIII; K, KpnI; Ms, MscI; P, PstI; Pv, PvuII, X, XbaI.

FIG. 10. Biosynthetic pathway for type 3 capsular polysaccharide. The functions of the proteins encoded by the type 3-specific genes, based on homologies, genetic, and biochemical data are shown. Additional functions may be necessary for capsule transport or attachment.

FIG. 11. Chromosome maps of the capsule regions in strains of types 2, 3, and 6B. The SacI-HindIII fragment (pJD377) from type 3 used for the probe is shown below the maps. Restriction sites are Bg, BglII; F, FspI; H, HindIII; S, SalI; Sac, SacI; Sp, SphI.

FIG. 12. Production of type 3 capsule. Buoyant densities of parents and derivatives expressing the type 3 capsule. Densities were determined by centrifugation on 0 to 50\% Percoll gradients for 30 min at 8,000 x g. Samples were grown in duplicate, and the density of each sample was determined in duplicate gradients. The results shown were obtained with bacteria grown on solid medium. Identical results were obtained from growth in liquid culture.
FIG. 13. Total capsule production. Triplicate cultures of each strain were grown to an OD₆₀₀ of ~0.5. Duplicate wells of polyvinyl plates were coated with either supernatant fluids or cell sonicates. Total capsule contents of the type 3 parent and the derivatives were determined by using a monoclonal antibody to type 3 capsule. See Table 2, footnote d, for explanation of strain designations.

FIG. 14A. All studies were done in BALB/ByJ mice. See Table 7, footnote d, for explanation of strain designations. Virulence of type 2 derivatives. The parental strains JD770 (3/3) and D39 (2/2) and the derivatives JD803 (2/3) and JD804 (2/3) had similar LD₅₀s (50 to 75 CFU). For time-to-death studies, groups of mice were infected i.p. with doses approximately 5- to 20-fold above the LD₅₀. The observed times to death were not related to the dose received. Each circle represents an individual mouse. The median times to death for D39 (2/2) and for derivatives JD803 (2/3) and JD804 (2/3) were 31.5 and 33.0 h, respectively (not significantly different). All three values differed significantly (P < 0.005) from that of the type 3 parent JD770 (52h). The values for JD803 and JD804 did not differ from each other.

FIG. 14B. All studies were done in BALB/ByJ mice. See Table 7, footnote d, for explanation of strain designations. Virulence of the type 5 derivatives. Mice were infected i.v. with doses of 10³ to 10⁶ CFU (10-fold increments) of each type 3 derivative. Doses of 10⁰ to 10² and 10³ to 10⁶ CFU were used fr the parental strains DBL5 (5/5) and JD770 (3/3), respectively. The total number of mice used per dose is listed beside each datum point. TK5010* represents the combined data for TK5010 (5/3), TK5011 (5/3), and TK5012 (5/3). The derivatives did not differ from each other but did differ
significantly from the parental strains JD770 ($P < 0.0005$) and DBL5 ($P < 0.0001$).

**FIG. 14C.** All studies were done in BALB/ByJ mice. See Table 7, footnote d, for explanation of strain designations. Virulence of type 6B derivatives. Mice were infected i.p. with doses of $10^1$ to $10^6$ CFU of the type 6B derivatives. Doses of $10^0$ to $10^3$ CFU and of $10^3$ to $10^6$ CFU were examined for the parent strains JD770 (3/3) and DBL1 (6B/6B), respectively. The total number of mice used per dose is listed beside each datum point. TK3028* represents the combined data for TK3026 (6B/3) and TK3028 (6B/3), which did not differ fro each other. However, these strains did differ significantly from the parental strains JD770 (3/3) ($P < 0.003$) and DBL1 (6B/6B) ($P < 0.0005$).

**FIG. 15A.** Model for the transfer of type-specific genes. Cassette type-recombination. Replacement of the recipient's type-specific genes with those of the donor results from homologous recombination between common regions that flank the type-specific cassettes. The open ellipsoid symbol represents sequence containing repeated element; the black oblong symbol represents common DNA upstream of type-specific cassettes and; the open oblong symbol represents common DNA (including plpA) downstream of type-specific cassettes.

**FIG. 15B.** Model for the transfer of type-specific genes. Binary encapsulation by recombination involving homology at only one end. Integration at one end of the type-specific cassette would occur via homologous recombination through the repeated element. Integration at the other end would result from an apparent illegitimate recombination. Linkage of the two type-specific cassettes would result if the integration occurred in a repeat element in or closely linked to the
recipient's capsule genes. Symbols represent the same as in FIG. 15A.

FIG. 15C. Model for the transfer of type-specific genes. Binary encapsulation via a transposition-like event. Type-specific cassettes flanked by the repeated element would resolve out of the chromosome and be transferred to recipient cells as circular intermediates. Recombination into the recipient chromosome could occur at a repeat element unlinked (as shown) or linked to the recipient's type-specific genes. Transfer of linear DNA could also yield binary strains as a result of recombination between the two repeat elements that flank the type-specific genes and two repeat elements that are closely linked in the recipient chromosome. Symbols represent the same as in FIG. 15A.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present inventors have determined a genetic and physical map that encompasses the region responsible for the synthesis of the polysaccharide capsule of S. pneumoniae. The polysaccharide capsule of S. pneumoniae is a potent defense against the immune response of the host organism and is directly involved in bacterial virulence. The capsule locus, cps, is basically composed of two functional regions: a central region that contains the genes responsible for capsular biosynthesis and is described herein as the type-specific region, and the non-type specific regions that flank the central biosynthetic type-specific region.

S. pneumoniae has evolved a complex 'antigenic shift' mechanism that allows the bacteria to evade the host immune system. The antigenic shift of S. pneumoniae occurs via homologous recombination of a type-specific cassette that is replaced through natural transformation.
S. pneumoniae is naturally competent allowing for the acquisition of chromosomal DNA from exogenous sources, such as other S. pneumoniae. Disclosed herein is evidence identifying the non-type specific regions as being responsible for providing the sequence identity that allows for homologous recombination cross-over points.

The present inventors have identified and cloned the region of the S. pneumoniae chromosome that contains genes involved in the production of type 3 capsular polysaccharide, and that is specific to type 3 strains. They have also cloned approximately 1-3 kb of DNA flanking both sides of this region and found it to be common to all capsular serotypes examined. A genetic and physical map of the region is presented in FIG. 6A, FIG. 6B, FIG. 6C, FIG. 6D, FIG. 6E, FIG. 6F, FIG. 6G, FIG. 6H, FIG. 6I, and FIG. 6J. A simplified version of which is shown in FIG. 7 and FIG. 8. The sites of insertion mutations made within the region are shown in FIG. 7. The regions found by hybridization studies to be specific to type 3 or common to all capsule types are also indicated in FIG. 7. The cloning of the upstream region, creation of insertion mutations, sequence analysis of the region, hybridization analyses using the upstream region, and an in vitro assay of type 3 capsule polymerization are described in the following examples.

The DNA sequence of the region containing the nine genes; cps3E, cps3C, cps3E, cps3D, cps3S, cps3U, cps3M, ‘pIpA, tnpA and the flanking DNA was determined and is presented in FIG. 6A, FIG. 6B, FIG. 6C, FIG. 6D, FIG. 6E, FIG. 6F, FIG. 6G, FIG. 6H, FIG. 6I, and FIG. 6J (SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:5 and SEQ ID NO:6) along with the deduced amino acid sequences (SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ
Based on genetic, molecular, and biochemical data the inventors have been able to assign putative functions to the type-specific genes in the pathway for type 3 capsular polysaccharide biosynthesis. Two of the genes, \(\text{cps3D}\) and \(\text{cps3S}\), are required for capsule synthesis. There is substantial evidence to indicate that \(\text{cps3D}\) encodes UDP-glucose dehydrogenase. Described herein is genetic evidence to indicate that several mutations causing the capsule-negative phenotype are located in the gene for UDP-glucose dehydrogenase. The predicted amino acid sequence has characteristics consistent with this function. Cps3D shows a high degree of homology to HasB, which is the UDP-glucose dehydrogenase of \(S.\ pyogenes\) (Dougherty & van de Rijn, 1993). Within Cps3D are sequences homologous to the active site and the NAD-binding site in known UDP-glucose dehydrogenases. It is not possible to perform the standard UDP-glucose dehydrogenase assay on extracts of \(S.\ pneumoniae\) due to the presence of a NADH oxidase, which copurifies with the enzyme (Smith, et al., 1960; Smith, et al., 1958). However, extracts from \(\text{cps3D}\) mutants could synthesize type 3 capsule in vitro if supplied with UDP-glucuronic acid, i.e., they lacked the ability to convert UDP-glucose to UDP-glucuronic acid and thus lack UDP-glucose dehydrogenase activity.

Cps3S is a new member of a family of polysaccharide synthases. All of these polysaccharide synthases for which the structures of the polysaccharides are known produce \(\beta\) (1-4) linked polysaccharides. Thus, it is possible that Cps3S forms the \(\beta\) (1-4) linkage in the disaccharide cellobiuronic acid (\(\text{glcA} \beta\) (1-4) \(\text{glc}\)), and that a second enzyme is required to polymerize (i.e., create the \(\beta\) (1-3) linkages) the disaccharides into the
full length polysaccharide. However, HasA, the enzyme most closely related to Cps3S, creates both linkages, a β (1-4) and a β (1-3), in the production of hyaluronic acid capsule (DeAngelis, et al., 1993). HasA has recently been shown to be sufficient for hyaluronic acid synthesis in heterologous bacteria, given the nucleotide sugar substituents (DeAngelis, et al., 1993a). Because the inventors did not find another required enzyme in the type 3-specific region, Cps3S, like HasA, may synthesize the polysaccharide by monomer addition.

Neither cps3U nor cps3M appears to be required for type 3 capsule synthesis. Cps3M and Cps3U should function to convert glucose-6-phosphate into glucose-1-phosphate, and glucose-1-phosphate into UDP-glucose, respectively (FIG. 10). Since UDP-glucose is necessary for the production of essential cell constituents, including teichoic acid and lipoteichoic acid (Austrian, et al., 1959), the products of other genes may complement the functions lost in Cps3U and Cps3M mutants. There are at least two plausible reasons for the retention of these genes in the type-specific region. One explanation is that their functions cannot be fully duplicated by the second enzymes. For example, they may play a role in regulating the amount of polysaccharide produced. Under given conditions, such as during infection, increased production of capsule could be advantageous. The large noncoding region upstream of cps3U might be a site at which regulation of cps3U and cps3M occurs. It should also be noted that the reactions carried out by Cps3M and Cps3U are each reversible, and the enzymes might be more active in the reverse reaction. Therefore, Cps3U and Cps3M might function to limit the amount of capsule produced.

Another possible explanation is that these genes were obtained along with the necessary type-specific
genes in a horizontal transfer from another organism and have not been lost. This theory is consistent with the hybridization data indicating that none of the type-specific genes could be detected even at low stringency in strains of six other pneumococcal types, including types with related capsular polysaccharide structures (Dillard, et al., 1994). However, if these genes serve no necessary function, it is surprising that they have been maintained in the type 3 cassettes of multiple strains; i.e., the restriction maps of the type 3 regions of five non-clonal type 3 strains are identical, and all have cps3U and cps3M.

There are three requirements for a DNA region to be considered a gene cassette: 1) more than one copy of a gene or set of genes must exist, each specifying the production of a different, but related, product; 2) each copy must be flanked by DNA which is common to all the copies; and 3) cassettes must undergo recombination resulting in the replacement of one copy by another. More than 80 different capsular serotypes of S. pneumoniae have been identified, and the structures of more than half of the polysaccharides have been determined (van Dam, et al., 1990).

The presence of multiple types implies that as many different sets of genes exist. The inventors have shown that all the necessary genes specific for the production of capsules of types 2, 3, and 6B (Example 16) are closely linked to an approximate 1.2 kb fragment present in all capsule types examined. This fragment (corresponding to SEQ ID NO:6 and part of SEQ ID NO:5, see FIG. 4 and Example 5), cloned from the region flanking the type 3-specific genes, contains a gene with a sequence virtually identical to a gene fragment from type 2 strain, described by Pearce et al., and designated p1pA (Pearce, et al., 1993). However, the flanking
region from type 3 strain is distinct from the sequence described by Pearce et al. (1993) in that it is missing about one third of the 5' end of the gene, designated 'plpA. Furthermore Pearce et al. did not identify the location of the plpA gene nor did they attempt to define the sequences on either side of the gene.

The mapping studies reported here confirm that the regions to the right of this fragment are common for at least 4 kb. The regions map differently to the left of the fragment (Example 16), implying that these regions contain the type-specific genes in types 2 and 6B, as shown herein for type 3.

The upstream left flanking region from type 3, SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3 and SEQ ID NO:4 is common to all capsule types examined (2, 3 and 6B, the Repeat region was also examined in 5, 6A, 8, 9 and 22; Example 17). However, the presence of multiple copies of the Repeat fragment (SEQ ID NO:4) has made the linkage in other types difficult to determine and it is possible that the repeat region may not flank the type-specific genes in other types.

Previous workers have provided biochemical evidence of replacement of capsule gene cassettes. When a type 1 and a type 2 strain were each transformed to type 14 or type 23, they no longer produced UDP-glucuronic acid, implying that the transformants had lost the UDP-glucose dehydrogenase gene (Austrian, et al., 1959). Similarly, a type 1 strain transformed to type 3 encapsulation no longer epimerized UDP-glucuronic acid to UDP-galacturonic acid. Molecular evidence is presented herein for the replacement of the type-specific genes for a type 3 strain transformed to type 2 encapsulation and a type 2 strain transformed to type 3 (Example 6). Together with
Example 18, these observations provide strong evidence for a cassette organization of the type-specific genes.

Since the proposal was put forth that capsule genes are exchanged through a cassette-type recombination, there has always been one glaring exception - binary encapsulation. At low frequency, strains of certain capsule types transformed with DNA from strains of certain other capsule types were found to produce both polysaccharides (Austrian, et al., 1959). Evidently, cassette-type recombination had not occurred in these transformants since the genes for the original capsule had been maintained. Bernheimer et al., found that stable binary strains contained the second set of type-specific genes at a site unlinked to the recipient's type-specific genes (Bernheimer, et al., 1967). Unstable binary strains frequently lost the donor type-specific genes, which were usually located at a site linked to the recipient type-specific genes (Bernheimer, et al., 1967; Bernheimer and Wermundsen, 1969).

Based on the hybridization data described here concerning the flanking regions and replacement of type-specific genes, as well as the work of Bernheimer concerning transformation to binary capsule types (Bernheimer, et al., 1968; Bernheimer, et al., 1967; Bernheimer, et al., 1969; Bernheimer and Wermundsen, 1972), the inventors can now propose models for capsule type change and binary capsule type formation. Cassette-type recombination would result from crossover events in the homologous flanking regions, leading to replacement of the type-specific genes. The left crossover could take place in the repeated element in strains containing this region linked to the type-specific genes but would occur in flanking DNA further upstream in strains that did not contain the repeat. This type of recombination is shown in Fig 19A.
The finding of a repeated element upstream of the type 3 capsule genes (SEQ ID NO:4; Example 17) may provide an explanation for binary encapsulation. It is clear that at least one of the copies of the repeated fragment in types 2 and 6B is unlinked to the capsule genes, since neither of the type-specific cassettes could be moved with a marker inserted in this location. In type 3, a 2.2 kb HindIII fragment containing the repeat element is linked to the type-specific genes but based on transformation studies, an 8 kb fragment is not. The frequency of binary capsule transformants observed by Bernheimer et al., was significantly lower (10-1000 fold) than related transformations resulting in replacement, leading them to suggest that the recombination event involved strong homology at only one end. Once integrated at the "atypical" location (unlinked to the type-specific cassette), the genes for the second capsule type could not be moved to the normal location, except by transformation of a strain containing the genes of that type in the normal location, again suggesting that the non-type specific flanking DNA on at least one end had been lost (Bernheimer, et al., 1967; Bernheimer and Wermundsen, 1972).

Since finding that part of the left flanking DNA (SEQ ID NO:4) is repeated in chromosomes of several strains, whereas the right flanking region is only present in one location, it is proposed that the repeated element of the left flanking region may be involved in the recombination that results in binary capsule type formation. The mechanism proposed by Bernheimer et al., for stable binary strains could involve homologous recombination at a repeated element unlinked to the capsule locus; the recombination at the other end of the capsule genes would occur by an apparent illegitimate recombination event, as shown in FIG. 15B.
An alternative possibility for the generation of stable binary strains, shown in FIG. 15C, involves a transposition-like event that could result if certain type-specific genes are flanked on both sides by the repeated element. Unstable binary strains could result from either type of integration occurring at repeated elements in, or closely linked to, the recipient's type-specific genes. Instability could result from recombination through genes common to both capsule types, as suggested by Bernheimer et al., for the UDP-glucose dehydrogenases of types 1 and 3. Results presented here provide the basis for examining these possibilities. Binary strains containing the two sets of genes linked are of particular interest since they might recombine to form a novel capsule type. Examination of strains producing related capsule structures may help elucidate the possible mechanisms involved in novel capsule type formation.

Epidemiological studies have indicated that capsule type varies independently of other factors, suggesting that a substantial amount of genetic exchange has occurred (Crain et al., 1990; Coffey et al., 1991; Versalovic et al., 1993). Nonetheless, virulence of clinical isolates appears to correlate with the capsule type expressed (Briles et al., 1992). Taken together, these data suggest that the capsule type has a prominent role in determining virulence. However, epidemiological studies cannot demonstrate a causal relationship between capsule type and virulence due to the variability in the genetic backgrounds of the different serotypes. The characterization of the S. pneumoniae capsule locus described here has facilitated the construction of isogenic strains differing only in capsule type. These strains have been used to evaluate the role of capsule type in virulence (Example 18). The cloning of capsule genes and elucidation of the genetic organization of the
capsule locus is a significant step toward understanding antigenic variation and virulence in this pathogen.

Cloning of the Type 3 region

There are a number of reasons for first cloning the type 3 specific genes: the type 3 capsule has a relatively simple structure that is expected to require a small number of genes for its synthesis; production of type 3 capsule is an easily identifiable phenotype; and finally, the availability of antibodies specific for the type 3 polysaccharide capsule allowed rapid screening for the presence of the capsule, of a large number of isolates, using an ELISA assay as described herein in Example 1. The approach provided and disclosed allowed for the first molecular genetic map of the cps gene locus.

The cloning of additional type-specific genes has been accomplished using the information derived from the present invention. Taking advantage of the non-type specific region one can isolate the DNA encoding other type-specific genes by simply obtaining a strain of S. pneumoniae known to have a type-specific capsule. Polymerase chain reaction using primers specific for opposite flanking regions and directed toward the opposite flanking non-type specific region are used to amplify the type-specific gene cassette. Where the size of the type-specific region is unknown, restriction fragment length polymorphism analysis, using probes specific for either or both of the non-type specific regions may be used to determine the size.

Antibodies specific for type-specific antigenic epitopes may be used with the present invention to distinguish and evaluate the stability of the S. pneumoniae strain prior to, and after cloning of the
region. It will also be used for verifying the directed transfer of type-specific genes to a prospective host. Preferred hosts for polysaccharide capsule production will be gram positive bacteria, in particular members of the *Streptococcus*, *Bacillus* and even *Staphylococcus* species.

Selection of Hosts for Type-Specific Capsule Production

The present invention provides methods for the selection, isolation and transformation of *Streptococcus* sp. with a type-specific capsule polysaccharide gene locus. A *cps* locus can now be isolated and used to specifically change the capsular phenotype of a selected host organism. The preferred host organism for use with the present invention is a bacteria that produces high amounts of the capsular polysaccharide.

Once a suitable high producing host is identified, it will be used to carry the type-specific genes of choice, as shown in Examples 6 and 18. The organisms can be converted to other serotypes by transforming the high producing recipient bacteria with a gene cassette or with intact genomic DNA. A gene cassette, as previously mentioned, is a segment of DNA comprising of one or more genes flanked by specific DNA sequences which enables incorporation of the cassette into a recipient's cell chromosome at a specific site or locus via homologous recombination. A cassette may contain type-specific genes, either alone or in combination with non-type specific genes. Of course, the preferred construct for transfection will be a cassette containing the non-type specific flanking regions.

A cassette of the *cps* locus comprising of the *cps* genes and the 5' and 3' flanking regions donated from any one of the 85 *S. pneumoniae* serotypes may be transformed
into a recipient *S. pneumoniae* also belonging to any one of the 85 serotypes. During transformation, recombination would occur in the flanking regions, resulting in the replacement of the recipient’s type-specific region by that of the donor. The capsule type of the recipient would be expected to change to that of the donor.

The introduction of a gene cassette comprising of the *cps* locus or DNA segment or genetic element, into *S. pneumoniae* may be performed by a variety of methods. A particularly preferred embodiment would be to digest the donor *S. pneumoniae* genomic DNA with one or more restriction enzymes such as those described in Table 1, for example, and then separate the entire *cps* locus from the rest of the genomic DNA by gel purification. This specific DNA segment, *cps* locus or genetic element, may then be ligated into a vector such as a plasmid or cosmid or bacteriophage, and transformed by various methods into the recipient *S. pneumoniae*. Alternatively, the donor *S. pneumoniae*’s entire genomic DNA may be naturally transformed into the recipient by a suitable method, e.g., as described in Example 1. Further still, the donor’s genomic DNA may be digested with one or more restriction enzymes and then ligated into a plasmid, cosmid or bacteriophage, without selecting specifically for the *cps* locus. This may then be transformed into the recipient *S. pneumoniae*. 
### TABLE 1

**RESTRICTION ENZYMES**

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Recognition Sequence</th>
</tr>
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<td>GACGT/C</td>
</tr>
<tr>
<td>Acc I</td>
<td>GT/MKAC</td>
</tr>
<tr>
<td>5 Acc II</td>
<td>CG/CG</td>
</tr>
<tr>
<td>Acc III</td>
<td>T/CCGGA</td>
</tr>
<tr>
<td>AcI I</td>
<td>CCGC(2/2)</td>
</tr>
<tr>
<td>AcY I</td>
<td>GR/GYGC</td>
</tr>
<tr>
<td>Afl II</td>
<td>C/TTAAG</td>
</tr>
<tr>
<td>10 Afl III</td>
<td>A/CRYGT</td>
</tr>
<tr>
<td>Age I</td>
<td>A/CCGCT</td>
</tr>
<tr>
<td>Aha III</td>
<td>TTT/AAA</td>
</tr>
<tr>
<td>Alu I</td>
<td>AG/CT</td>
</tr>
<tr>
<td>AlwN I</td>
<td>CAGNNN/CTG</td>
</tr>
<tr>
<td>15 Aoc I</td>
<td>CC/TNAGG</td>
</tr>
<tr>
<td>Apa I</td>
<td>GGGCC/C</td>
</tr>
<tr>
<td>ApaB I</td>
<td>GCANNNNN/TGC</td>
</tr>
<tr>
<td>ApaL I</td>
<td>G/TGCAC</td>
</tr>
<tr>
<td>Asc I</td>
<td>GG/GCGGCC</td>
</tr>
<tr>
<td>20 Asu I</td>
<td>G/GNCC</td>
</tr>
<tr>
<td>Asu II</td>
<td>TT/CGAA</td>
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<tr>
<td>Ava II</td>
<td>G/GWCC</td>
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<td>25 Avr III</td>
<td>C/CTAGG</td>
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<td>Bcc I</td>
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<td>Bcsef I</td>
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<tr>
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<td>Sequence</td>
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<td>CGAT/CG</td>
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<td>BsaB I</td>
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</tr>
<tr>
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</tr>
<tr>
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<td>GTCTC(1/5)</td>
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<tr>
<td>Bsp50 I</td>
<td>CG/CG</td>
</tr>
<tr>
<td>BspG I</td>
<td>CG/CGCTGGAC</td>
</tr>
<tr>
<td>BspH I</td>
<td>T/CATGA</td>
</tr>
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<tr>
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<td>GAGCGG(-3/-3)</td>
</tr>
<tr>
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</tr>
<tr>
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<td>CC/WGG</td>
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<tr>
<td>BstX I</td>
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</tr>
<tr>
<td>Dpn I</td>
<td>GA/TC</td>
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</table>
Dra I       TTT/AAA
Dra II      TG/GNCCY
Dra III     CACNNN/GTG
Drd I       GACNNN/NNGTC
5 Drd II    GAACCA
Dsa I       C/CRYGG
Eam1105 I   GACNNN/NNGTC
Eci I       TCCGCC
Eco3 II     GGTCTC(1/5)
10 Eco47 III AGC/GCT
Eco52 I     C/GGCCG
Eco57 I     CTGAAG(16/14)
EcoN I      CCTNN/NNNAGG
EcoR I      G/AATTC
15 EcoR II  /CCWGG
Ecor V      GAT/ATC
Esp I       GC/TNAGC
Esp3 I      CGTCTC(1/5)
Fau I       CCCGC(4/6)
20 Fin I     GTCCC
Fnu4H I     GC/NGC
FnuD II     CG/CY
Fok I       GGATG(9/13)
Fse I       GGCCCG/CC
25 Fsi I     R/AAATTY
Gdi II      YGGCCG(-5/-1)
Gsu I       CTGGAG(16/14)
Hae I       WGG/CCW
Hae II      RGCGC/Y
30 Hae III   GG/CC
Hga I       GACGC(5/10)
HgiA I      GWGCW/C
HgaC I      G/GYRCC
HgiE II     ACCNNNNNNNGGT
35 HgiJ II  GRGCY/C
<table>
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<tr>
<th>Enzyme</th>
<th>Recognition Sequence</th>
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<td>Hha I</td>
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<td>CC/TNAGG</td>
</tr>
<tr>
<td>Mwo I</td>
<td>GCNNNNN/NNGC</td>
</tr>
<tr>
<td>Nae I</td>
<td>GCC/GGC</td>
</tr>
<tr>
<td>Nar I</td>
<td>GG/CGCC</td>
</tr>
<tr>
<td>Nci I</td>
<td>CC/SGG</td>
</tr>
<tr>
<td>Nco I</td>
<td>C/CATGG</td>
</tr>
<tr>
<td>Nde I</td>
<td>CA/TATG</td>
</tr>
<tr>
<td>Nhe I</td>
<td>G/CTAGC</td>
</tr>
<tr>
<td>Nla III</td>
<td>CATG/</td>
</tr>
<tr>
<td>Nla IV</td>
<td>GGN/NCC</td>
</tr>
</tbody>
</table>
Not I  GC/GGCCGC
Nru I  TCG/CGA
Nsi I  ATGCA/T
Nsp I  RCATG/Y
NspB II CMG/CKG
Pac I  TTAAT/TA
Pal I  GG/CC
Pfll 108 I  TCGTAG
PflM I  CCANNNN/NTGG
Ple I  GAGTC(4/5)
PmaC I  CAC/GTG
Pme I  GTTT/AAAC
PpuM I  RG/GWCCY
PshA I  GACNN/NNGTC
PspA I  C/CCGGG
Pst I  CTGCA/G
Pvu I  CGAT/CG
Pvu II  CAG/CTG
RleA I  CCCACA(12/9)
Rsa I  GT/AC
Rsr II  CG/GWCCG
Sac I  GAGCT/C
Sac II  CCGC/GG
Sal I  G/TGCAG
Sap I  GCTCTTC(1/4)
Sau3A I  /GATC
Sau96 I  G/GNCC
Sau I  CC/TNAGG
Sca I  AGT/ACT
ScrF I  CC/NGG
Sdu I  GDGCH/C
Sec I  C/CNNGG
SfaN I  GATC/(5/9)
Sfc I  CTYRAG
Sfe I  C/TYRAG
Sfi I  GGCCNNNN/NGGC
SgrA I  CR/CCGGYG
Sma I  CCC/GGG
Sna I  CTATAAC
5  SnaB I  TAC/GTA
Spe I  A/CTAGT
Sph I  GCATG/C
Spl I  C/GTACG
Srf I  GCCC/GGGC
10  Sse838 I  CCTGCA/GG
Ssp I  AAT/ATT
Stu I  AGG/CCT
Sty I  C/CWGGG
Swa I  ATTT/AAAT
15  Taq I  T/CGA
Taq II  GACCGA(11/9)
Tfi I  GAWTC
Tsp45 I  GTSAC
Tsp E I  AATT
20  Tth111 I  GACN/NNGTC
Tth111 II  CAARCA(11/9)
Vsp I  AT/TAAT
Xba I  T/CTGAGA
Xcm I  CCANNNNN/NNNNTGG
25  Xho I  C/TCGAG
Xho II  R/GATCY
Xma I  C/CCGGG
Xma III  C/GGCCG
Xmn I  GAANN/NNTTC
30
Nucleic Acid Hybridization

The DNA sequences disclosed herein will find utility as probes or primers in nucleic acid hybridization embodiments. As such, it is contemplated that oligonucleotide fragments corresponding to the sequence(s) of SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3 (including sequences in between), SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6 for stretches of between about 10-14 nucleotides to about 20 or to about 30 nucleotides will find particular utility, with even longer sequences, e.g., 40, 50, 100, 200, 500, and even up to full length, being more preferred for certain embodiments. The ability of such nucleic acid probes to specifically hybridize to non-type-specific and to type-specific-encoding sequences will enable them to be of use in a variety of embodiments. For example, the probes can be used in a variety of assays for detecting the presence of complementary sequences in a given sample, as may be used, for example to isolate related type-specific genes. Alternatively, one may use the non-type-specific regions to aid in the isolation and cloning of additional type-specific cassettes. However, other uses are envisioned, including the use of the sequence information for the preparation of mutant species primers, or primers for use in preparing other genetic constructions.

Nucleic acid molecules having stretches of about 10-14, 20, 30, 50, or even of about 100-200 nucleotides or so, complementary to SEQ ID NO:1, SEQ ID NO:2 and SEQ ID NO:3 (including sequences in between), SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6, will have utility as hybridization probes. These probes will be useful in a variety of hybridization embodiments, such as Southern and Northern blotting in connection with analyzing genomic structure and organization of type-specific genes or both linked and non-linked regulatory genes in diverse
strains of *S. pneumoniae*. The total size of fragment, as well as the size of the complementary stretch(es), will ultimately depend on the intended use or application of the particular nucleic acid segment. Smaller fragments will generally find use in hybridization embodiments, wherein the length of the complementary region may be varied, such as between about 10-14 and about 100 nucleotides, or even up to full length according to the complementary sequences one wishes to detect.

The use of a hybridization probe of about 10-14 nucleotides in length allows the formation of a duplex molecule that is both stable and selective. Molecules having complementary sequences over stretches greater than 10-14 bases in length are generally preferred, though, in order to increase stability and selectivity of the hybrid, and thereby improve the quality and degree of specific hybrid molecules obtained. One will generally prefer to design nucleic acid molecules having gene-complementary stretches of 15 to 20 nucleotides, or even longer where desired. Such fragments may be readily prepared by, for example, directly synthesizing the fragment by chemical means, by application of nucleic acid reproduction technology, such as the PCR technology of U.S. Patent 4,603,102 (incorporated herein by reference) or by introducing selected sequences into recombinant vectors for recombinant production.

Accordingly, the nucleotide sequences of the invention may be used for their ability to selectively form duplex molecules with complementary stretches of non-type- and of type-specific genes. Depending on the application envisioned, one will desire to employ varying conditions of hybridization to achieve varying degrees of selectivity of probe towards target sequence. Such hybridization conditions are standard in the art and include low stringency and high stringency. For
applications requiring high selectivity, one will typically desire to employ relatively stringent conditions to form the hybrids, e.g., one will select relatively low salt and/or high temperature conditions, such as provided by 0.02M-0.15M NaCl at temperatures of 50°C to 70°C. One particular example is using the QuickHyb® system (Stratagene's Illuminator™ Nonradioactive Detection System) at 68°C. Such selective conditions tolerate little, if any, mismatch between the probe and the template or target strand, and would be particularly suitable for isolating other genes encoding gene products that are involved in the production of capsule polysaccharides. A preferred embodiment for hybridization conditions is described in detail in Example 4. Further standard hybridization conditions can be found in Sambrook et al., (1989), and are known to those of skill in the art.

Of course, for some applications, for example, where one desires to prepare mutants employing a mutant primer strand hybridized to an underlying template or where one seeks to isolate type-specific-encoding sequences from related species, functional equivalents, or the like, less stringent hybridization conditions will typically be needed in order to allow formation of the heteroduplex.

One may also desire to employ other hybridization techniques and to change salt conditions such as varying the amount of salt from between about 0.15M-0.9M. Other parameters that can be modified may be temperature such as those ranging from 20°C to 55°C to optimize the signal-to-noise ratio to reduce unwanted background. The techniques for optimizing hybridization conditions are well known to those of skill in the art and are generally also described within the instruction manual for various reagents and apparatus.
In any case, it is generally appreciated that conditions can be rendered more stringent by the addition of increasing amounts of formamide, which serves to destabilize the hybrid duplex in the same manner as increased temperature. Thus, hybridization conditions can be readily manipulated as is known to those of skill in the art, and thus will generally be a method of choice depending on the desired results.

In certain embodiments, it will be advantageous to employ nucleic acid sequences of the present invention in combination with an appropriate means, such as a label, for determining hybridization. A wide variety of appropriate indicator means are known in the art, including fluorescent, radioactive, enzymatic or other ligands, such as avidin/biotin, which are capable of giving a detectable signal. In preferred embodiments, one will likely desire to employ a fluorescent label or an enzyme tag, such as urease, alkaline phosphatase or peroxidase, instead of radioactive or other environmental undesirable reagents. In the case of enzyme tags, colorimetric indicator substrates are known which can be employed to provide a means visible to the human eye or spectrophotometrically, to identify specific hybridization with complementary nucleic acid-containing samples.

In general, it is envisioned that the hybridization probes described herein will be useful both as reagents in solution hybridization as well as in embodiments employing a solid phase. In embodiments involving a solid phase, the test DNA (or RNA) is adsorbed or otherwise affixed to a selected matrix or surface. This fixed, single-stranded nucleic acid is then subjected to specific hybridization with selected probes under desired conditions. The selected conditions will depend on the particular circumstances based on the particular criteria
required (depending, for example, on the G+C contents, type of target nucleic acid, source of nucleic acid, size of hybridization probe, etc.). Following washing of the hybridized surface so as to remove nonspecifically bound probe molecules, specific hybridization is detected, or even quantified, by means of the label.

Longer DNA segments will often find particular utility in the recombinant production of peptides or proteins. DNA segments which encode peptide antigens from about 15 to about 50 amino acids in length, or more preferably, from about 15 to about 30 amino acids in length are contemplated to be particularly useful, as are DNA segments encoding entire cps locus encoded proteins, such as those of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:5 and SEQ ID NO:6. DNA segments encoding peptides will generally have a minimum coding length in the order of about 45 to about 150, or to about 90 nucleotides.

The nucleic acid segments of the present invention, regardless of the length of the coding sequence itself, may be combined with other DNA sequences, such as promoters, repressors, attenuators, additional restriction enzyme sites, multiple cloning sites, other coding segments, and the like, such that their overall length may vary considerably. It is contemplated that a nucleic acid fragment of almost any length may be employed, with the total length preferably being limited by the ease of preparation and use in the intended recombinant DNA protocol. For example, nucleic acid fragments may be prepared in accordance with the present invention which are up to about 10,000 base pairs in length, with segments of about 5,000 or 3,000 being preferred and segments of about 1,000 base pairs in length being particularly preferred.
It will be understood that this invention is not limited to the particular nucleic acid sequences of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6, or to the particular amino acid sequences of SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15 and SEQ ID NO:16. Therefore, DNA segments prepared in accordance with the present invention may also encode biologically functional equivalent proteins or peptides which have variant amino acid sequences. Such sequences may arise as a consequence of codon redundancy and functional equivalency which are known to occur naturally within nucleic acid sequences and the proteins thus encoded. Alternatively, functionally equivalent proteins or peptides may be created via the application of recombinant DNA technology, in which changes in the protein structure may be engineered, based on considerations of the properties of the amino acids being exchanged.

DNA segments encoding a gene, including the cpsB, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, plpA and tnpA genes may be introduced into recombinant host cells and employed for expressing and producing the type-specific proteins for use in producing type-specific capsule polysaccharides. Alternatively, through the application of genetic engineering techniques, subportions or derivatives of selected type-specific gene locus genes may be employed. Equally, through the application of site-directed mutagenesis techniques, one may re-engineer DNA segments of the present invention to alter the coding sequence, e.g., to introduce improvements to the antigenicity of the protein or to test mutants in order to examine the production of capsule polysaccharides at the molecular level. Where desired, one may also prepare fusion peptides, e.g., where the type-specific coding
regions are aligned within the same expression unit with
other proteins or peptides having desired functions, such
as for immunodetection purposes (e.g., enzyme label
coding regions).

Screening Method Type-Specific Genes

Screening for type-specific genes provides another
utility for the cps loci of the present invention. A
type-specific screening protocol will allow for the
epidemiological identification of S. pneumoniae and its
serotypes at the molecular level. By using one or both
of the non-type specific regions as probes one can
determine the presence of S. pneumoniae from a small
sample by immobilizing DNA from the sample onto a solid
matrix, for example a slot blot using nitrocellulose, and
hybridizing thereto a probe as described in the present
invention.

Using either or both of the non-type specific
regions of the present invention as a probe or probes one
may also screen southern blots. The screening of
southern blots may allow one to determine not only the
presence of S. pneumoniae but also the exact genotype of
S. pneumoniae present in the sample. In conjunction with
densitometric analysis of a southern blot containing
multiple serotypes on may determine not only the relative
frequency of serotypes within a sample, but in addition
one may examine the changing characteristics of the
serotypes by examining samples taken at distinct time
periods.

It also allows the clinician to determine if a
patient is having a recurrence of a particular serotype,
if the patient is susceptible to a particular serotype or
types, or if a new serotype is increasing in the
population.
Site-Specific Mutagenesis

Site-specific mutagenesis, also known as site-directed mutagenesis, is a technique useful in the preparation of changes, directed by the laboratory technician, that change the characteristics of genes and their gene products, for the addition of restriction sites, for modifying the activity of promoters, repressors, attenuators, and for directed changes affecting recombination. All of these changes may be produced through specific mutagenesis of the underlying non-type- and type-specific DNA of the present invention. The technique further provides a ready ability to prepare and test sequence variants, for example, incorporating one or more of the foregoing considerations, by introducing one or more nucleotide sequence changes into the DNA. Site-specific mutagenesis allows the production of mutants through the use of specific oligonucleotide sequences which encode the DNA sequence of the desired mutation, as well as a sufficient number of adjacent nucleotides, to provide a primer sequence of sufficient size and sequence complexity to form a stable duplex on both sides of the deletion junction being traversed. Typically, a primer of about 17 to 25 nucleotides in length is preferred, with about 5 to 10 residues on both sides of the junction of the sequence being altered.

In general, the technique of site-specific mutagenesis is well known in the art, as exemplified by various publications. As will be appreciated, the technique typically employs a phage vector which exists in both a single stranded and double stranded form. Typical vectors useful in site-directed mutagenesis include vectors such as the M13 phage. These phage are readily commercially available and their use is generally well known to those skilled in the art. Double stranded plasmids are also routinely employed in site directed
mutagenesis which eliminates the step of transferring the gene of interest from a plasmid to a phage.

In general, site-directed mutagenesis in accordance herewith is performed by first obtaining a single-stranded vector or melting apart the two strands of a double stranded vector which includes within its sequence a DNA sequence which encodes the type-specific protein or proteins encoded by the type-specific gene locus. An oligonucleotide primer bearing the desired mutated sequence is prepared, generally synthetically. This primer is then annealed with the single-stranded vector, and subjected to DNA polymerizing enzymes such as E. coli polymerase I Klenow fragment, in order to complete the synthesis of the mutation-bearing strand. Thus, a heteroduplex is formed wherein one strand encodes the original non-mutated sequence and the second strand bears the desired mutation. This heteroduplex vector is then used to transform appropriate cells, such as E. coli cells, and clones are selected which include recombinant vectors bearing the mutated sequence arrangement.

The preparation of sequence variants of type-specific genes using site-directed mutagenesis is provided as a means of producing potentially useful species, for example a strain having enhanced production of type-specific capsular polysaccharides, and is not meant to be limiting as there are other ways in which sequence variants of other type-specific genes may be obtained. For example, recombinant vectors encoding other type-specific genes, as described herein using the non-type specific regions of the capsule polysaccharide gene cassette are encompassed.
Biological Functional Equivalents

Even though the invention has been described with a certain degree of particularity, it is evident that many alternatives, modifications, and variations will be apparent to those skilled in the art in light of the foregoing disclosure. Accordingly, it is intended that all such alternatives, modifications, and variations which fall within the spirit and the scope of the invention be embraced by the defined claims.

As used in this application, the term "DNA segment" refers to a DNA molecule that has been isolated free of total genomic DNA of a particular species. Therefore, a DNA segment encoding the cps locus refers to a DNA segment that contains the 5' and/or 3' flanking regions, or the cpsB, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, tnpA or pIpA coding sequences, yet is isolated away from, or purified free from, total genomic DNA of S. pneumoniae. Included within the term "DNA segment", are DNA segments and smaller fragments of such segments, and also recombinant vectors, including, for example, plasmids, cosmids, phage, viruses, and the like.

Similarly, a DNA segment comprising an isolated or purified 5' or 3' flanking region, or cpsB, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, pIpA or even tnpA gene, refers to a DNA segment including the coding sequences and, in certain aspects, regulatory sequences, isolated substantially away from other naturally occurring genes or protein encoding sequences. In this respect, the term "gene" is used for simplicity to refer to a protein, polypeptide or peptide encoding unit. As will be understood by those in the art, this term includes both genomic sequences, cDNA sequences and smaller engineered gene segments that express, or may be adapted to express, proteins, polypeptides or peptides.
"Isolated substantially away from other coding sequences" means that the locus of interest, in this case the 5' or 3' flanking regions, or cps B, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, tnpA or plpA coding sequences, forms the significant part of the DNA segment, and that the DNA segment does not contain large portions of naturally-occurring coding DNA, such as large chromosomal fragments or other functional genes or cDNA coding regions. Of course, this refers to the DNA segment as originally isolated, and does not exclude genes or coding regions later added to the segment by the laboratory technician.

In particular embodiments, the invention concerns isolated DNA segments and recombinant vectors incorporating DNA sequences that include the 5' or 3' flanking regions denoted by SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3 and SEQ ID NO:4 or SEQ ID NO:6, respectively. DNA segments and vectors that incorporate DNA sequences that encode CpsB, CpsC, CpsE, CpsD, CpsS, CpsU, CpsM, PlpA or transposase A proteins that include within their amino acid sequences an amino acid sequence as set forth in SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, SEQ ID NO:10, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13, SEQ ID NO:14, SEQ ID NO:15 or SEQ ID NO:16 respectively are also included.

The term "a sequence as set forth in SEQ ID NO:7-16" means that the sequence substantially corresponds to a portion of SEQ ID NO:7-16 and has relatively few amino acids that are not identical to, or a biologically functional equivalent of, the amino acids of SEQ ID NO:7-16. The term "biologically functional equivalent" is well understood in the art and is further defined in detail herein. Accordingly, sequences that have between about 75% and about 85%; or more preferably, between about 86% and about 95%; or even more preferably, between
about 96% and about 99%; of amino acids that are identical or functionally equivalent to the amino acids of SEQ ID NO:7-16 will be sequences that are "essentially as set forth in SEQ ID NO:7-16."

Naturally, it will be understood that for the Cps proteins, the definition of "equivalents" in this sense does not extend to distinct, but homologous proteins, such as CpsD and HasB from *Streptococcus pyogenes*; CpsS and HasA from *S. pyogenes*, NodC from *Rhizobium meliloti*; nor CpsU and GtaB from *Bacillus subtilis*. Rather, the scope of equivalents contemplated are such that the changes made still result in a protein that is structurally and functionally a Cps protein.

In certain other embodiments, the invention concerns isolated DNA segments and recombinant vectors that include within their sequence a nucleic acid sequence essentially as set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4 and SEQ ID NO:6, for the flanking regions; and SEQ ID NO:5 for the type-specific encoding regions. The term "as set forth in SEQ ID NO:5" is used in the same sense as described above and means that the nucleic acid sequence substantially corresponds to a portion of SEQ ID NO:5 and has relatively few codons that are not identical, or functionally equivalent, to the codons of SEQ ID NO:5. The term "functionally equivalent codon" is used herein to refer to codons that encode the same amino acid, such as the six codons for arginine or serine, and also refers to codons that encode biologically equivalent amino acids (as in Table 2).

It will be understood that acid and nucleic acid sequences may include additional residues, such as additional N- or C-terminal amino acids or 5′ or 3′ sequences, and yet still be essentially as set forth in one of the sequences disclosed herein, so long as the
sequence meets the criteria set forth above, including the maintenance of biological protein activity where protein expression is concerned. The addition of terminal sequences particularly applies to coding nucleic acid sequences that may, for example, include various non-coding sequences flanking either of the 5' or 3' portions of the coding region, such as promoters.

Allowing for the degeneracy of the genetic code, sequences that have between about 75% and about 85%; or more preferably, between about 85% and about 95%; or even more preferably, between about 95% and about 99% of nucleotides that are identical to the nucleotides of SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 or SEQ ID NO:6 (SEQ ID NO:1-6), will be sequences that are "as set forth in SEQ ID NO:1-6". Sequences that are essentially the same as those set forth in SEQ ID NO:1-6 may also be functionally defined as sequences that are capable of hybridizing to a nucleic acid segment containing the complement of SEQ ID NO:1-6 under standard conditions. Suitable hybridization conditions will be well known to those of skill in the art and are clearly set forth herein, e.g., see Example 4.
Table 2. Amino Acids and the Corresponding Codons.

<table>
<thead>
<tr>
<th>Amino Acids</th>
<th>Codons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>Ala A</td>
</tr>
<tr>
<td>Cysteine</td>
<td>Cys C</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>Asp D</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>Glu E</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>Phe F</td>
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<tr>
<td>Glycine</td>
<td>Gly G</td>
</tr>
<tr>
<td>Histidine</td>
<td>His H</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>Ile I</td>
</tr>
<tr>
<td>Lysine</td>
<td>Lys K</td>
</tr>
<tr>
<td>Leucine</td>
<td>Leu L</td>
</tr>
<tr>
<td>Methionine</td>
<td>Met M</td>
</tr>
<tr>
<td>Asparagine</td>
<td>Asn N</td>
</tr>
<tr>
<td>Proline</td>
<td>Pro P</td>
</tr>
<tr>
<td>Glutamine</td>
<td>Gin Q</td>
</tr>
<tr>
<td>Arginine</td>
<td>Arg R</td>
</tr>
<tr>
<td>Serine</td>
<td>Ser S</td>
</tr>
<tr>
<td>Threonine</td>
<td>Thr T</td>
</tr>
<tr>
<td>Valine</td>
<td>Val V</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>Trp W</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>Tyr Y</td>
</tr>
</tbody>
</table>

Naturally, the present invention also encompasses DNA segments that are complementary, or essentially complementary, to the sequences set forth in SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6. Nucleic acid sequences that are "complementary" are those that are capable of base-pairing according to the standard Watson-Crick complementarity rules. As used herein, the term "complementary sequences" means nucleic acid sequences that are substantially complementary, as may be assessed.
by the same nucleotide comparison set forth above, or as defined as being capable of hybridizing to the nucleic acid segment of SEQ ID NO:1-6 under relatively stringent conditions such as those described herein as SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6.

The DNA segments of the present invention include those encoding biologically functional equivalent proteins and peptides. Such sequences may arise as a consequence of codon redundancy and functional equivalency that are known to occur naturally within nucleic acid sequences and the proteins thus encoded. Alternatively, functionally equivalent proteins or peptides may be created via the application of recombinant DNA technology, in which changes in the protein structure may be engineered, based on considerations of the properties of the amino acids being exchanged. Changes designed in the laboratory, may be introduced through the application of site-directed mutagenesis techniques, e.g., to introduce improvements to the antigenicity of the protein or to test S. pneumoniae mutants in order to examine capsular productivity at the molecular level.

If desired, one may also prepare fusion proteins and peptides, e.g., where the cpsB, cpsC, cpsE, cpsD, cpsS, cpsU, cpsM, plpA and tnpA coding regions are aligned within the same expression unit with other proteins or peptides having desired functions, such as for purification or immunodetection purposes (e.g., proteins that may be purified by affinity chromatography and enzyme label coding regions, respectively).

As mentioned above, modification and changes may be made in the structure of CpsB, CpsC, CpsE, CpsD, CpsS, CpsU, CpsM, plpA or tnpA and still obtain a molecule
having like or otherwise desirable characteristics. For example, certain amino acids may be substituted for other amino acids in a protein structure without appreciable loss of interactive binding capacity with structures such as, for example, antigen-binding regions of antibodies or binding sites on substrate molecules, receptors, or catalytic regulation of capsular polysaccharide production. Since it is the interactive capacity and nature of a protein that defines that protein’s biological functional activity, certain amino acid sequence substitutions can be made in a protein sequence (or, of course, its underlying DNA coding sequence) and nevertheless obtain a protein with like (agonistic) properties. Equally, the same considerations may be employed to create a protein or polypeptide with countervailing (e.g., antagonistic) properties. It is thus contemplated by the inventors that various changes may be made in the sequence of SEQ ID NO:7, SEQ ID NO:11, SEQ ID NO:12, SEQ ID NO:13 or SEQ ID NO:14, SEQ ID NO:15 or SEQ ID NO:16 proteins or peptides (or underlying DNA) without appreciable loss of their biological utility or activity.

It is also well understood by the skilled artisan that, inherent in the definition of a biologically functional equivalent protein or peptide, is the concept that there is a limit to the number of changes that may be made within a defined portion of the molecule and still result in a molecule with an acceptable level of equivalent biological activity. Biologically functional equivalent peptides are thus defined herein as those peptides in which certain, not most or all, of the amino acids may be substituted. In particular, the function of given protein must be retained to be an equivalent. Of course, a plurality of distinct proteins/peptides with different substitutions may easily be made and used in accordance with the invention.
It is also well understood that where certain residues are shown to be particularly important to the biological or structural properties of a protein or peptide, e.g., residues in active sites, such residues may not generally be exchanged. This is the case in the present invention, where CpsD has a putative NAD-binding site and active site region at residues 2 to 29 and 251-263 (SEQ ID NO:11) respectively.

Amino acid substitutions are generally based on the relative similarity of the amino acid side-chain substituents, for example, their hydrophobicity, hydrophilicity, charge, size, and the like. An analysis of the size, shape and type of the amino acid side-chain substituents reveals that arginine, lysine and histidine are all positively charged residues; that alanine, glycine and serine are all a similar size; and that phenylalanine, tryptophan and tyrosine all have a generally similar shape. Therefore, based upon these considerations, arginine, lysine and histidine; alanine, glycine and serine; and phenylalanine, tryptophan and tyrosine; are defined herein as biologically functional equivalents.

In making changes, the hydrophatic index of amino acids may be considered. Each amino acid has been assigned a hydrophatic index on the basis of their hydrophobicity and charge characteristics, these are: isoleucine (+4.5); valine (+4.2); leucine (+3.8); phenylalanine (+2.8); cysteine/cystine (+2.5); methionine (+1.9); alanine (+1.8); glycine (-0.4); threonine (-0.7); serine (-0.8); tryptophan (-0.9); tyrosine (-1.3); proline (-1.6); histidine (-3.2); glutamate (-3.5); glutamine (-3.5); aspartate (-3.5); asparagine (-3.5); lysine (-3.9); and arginine (-4.5).
The importance of the hydropathic amino acid index in conferring interactive biological function on a protein is generally understood in the art (Kyte & Doolittle, 1982, incorporated herein by reference). It is known that certain amino acids may be substituted for other amino acids having a similar hydropathic index or score and still retain a similar biological activity. In making changes based upon the hydropathic index, the substitution of amino acids whose hydropathic indices are within ±2 is preferred, those which are within ±1 are particularly preferred, and those within ±0.5 are even more particularly preferred.

It is also understood in the art that the substitution of like amino acids can be made effectively on the basis of hydrophilicity. U.S. Patent 4,554,101, incorporated herein by reference, states that the greatest local average hydrophilicity of a protein, as governed by the hydrophilicity of its adjacent amino acids, correlates with its immunogenicity and antigenicity, i.e. with a biological property of the protein. It is therefore understood that an amino acid can be substituted for another having a similar hydrophilicity value and still obtain a biologically equivalent, and in particular, an immunologically equivalent protein.

As detailed in U.S. Patent 4,554,101, the following hydrophilicity values have been assigned to amino acid residues: arginine (+3.0); lysine (+3.0); aspartate (+3.0 ± 1); glutamate (+3.0 ± 1); serine (+0.3); asparagine (+0.2); glutamine (+0.2); glycine (0); threonine (-0.4); proline (-0.5 ± 1); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine (-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); tryptophan (-3.4).
In making changes based upon similar hydrophilicity values, the substitution of amino acids whose hydrophilicity values are within ±2 is preferred, those which are within ±1 are particularly preferred, and those within ±0.5 are even more particularly preferred.

As outlined above, amino acid substitutions are generally therefore based on the relative similarity of the amino acid side-chain substituents, for example, their hydrophobicity, hydrophilicity, charge, size, and the like. Exemplary substitutions which take various of the foregoing characteristics into consideration are well known to those of skill in the art and include: arginine and lysine; glutamate and aspartate; serine and threonine; glutamine and asparagine; and valine, leucine and isoleucine.

Antibody Generation

Means for preparing and characterizing antibodies are well known in the art (See, e.g., Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, 1988; incorporated herein by reference). This invention thus contemplates the generation of antibodies against the proteins CpsB, CpsC, CpsE, CpsD, CpsS, CpsU, CpsM, PlpA and transposase A or peptides derived therefrom. The CpsB, CpsC, CpsE, CpsD, CpsS, CpsU, CpsM, PlpA and transposase A proteins or peptides may be obtained using standard methods of recombinant expression as is routinely in the art.

The methods for generating monoclonal antibodies (MAbs) generally begin along the same lines as those for preparing polyclonal antibodies. Briefly, a polyclonal antibody is prepared by immunizing an animal with an immunogenic composition in accordance with the present invention and collecting antisera from that immunized
animal. A wide range of animal species can be used for the production of antisera. Typically the animal used for production of anti-antisera is a rabbit, a mouse, a rat, a hamster, a guinea pig or a goat. Because of the relatively large blood volume of rabbits, a rabbit is a preferred choice for production of polyclonal antibodies.

As is well known in the art, a given composition may vary in its immunogenicity. It is often necessary therefore to boost the host immune system, as may be achieved by coupling a peptide or polypeptide immunogen to a carrier. Exemplary and preferred carriers are keyhole limpet hemocyanin (KLH) and bovine serum albumin (BSA). Other albumins such as ovalbumin, mouse serum albumin or rabbit serum albumin can also be used as carriers. Means for conjugating a polypeptide to a carrier protein are well known in the art and include glutaraldehyde, m-maleimidobencoyl-N-hydroxysuccinimide ester, carbodiimide and bis-biazotized benzidine.

As is also well known in the art, the immunogenicity of a particular immunogen composition can be enhanced by the use of non-specific stimulators of the immune response, known as adjuvants. Exemplary and preferred adjuvants include complete Freund’s adjuvant (a non-specific stimulator of the immune response containing killed Mycobacterium tuberculosis), incomplete Freund’s adjuvants and aluminum hydroxide adjuvant.

The amount of immunogen composition used in the production of polyclonal antibodies varies upon the nature of the immunogen as well as the animal used for immunization. A variety of routes can be used to administer the immunogen (subcutaneous, intramuscular, intradermal, intravenous and intraperitoneal). The production of polyclonal antibodies may be monitored by sampling blood of the immunized animal at various points
following immunization. A second, booster injection, may also be given. The process of boosting and titering is repeated until a suitable titer is achieved. When a desired level of immunogenicity is obtained, the immunized animal can be bled and the serum isolated and stored, and/or the animal can be used to generate MAb.

MAbs may be readily prepared through use of well-known techniques, such as those exemplified in U.S. Patent 4,196,265, incorporated herein by reference. Typically, this technique involves immunizing a suitable animal with a selected immunogen composition, e.g., a purified or partially purified CpsB, CpsC, CpsE, CpsD, CpsS, CpsU, CpsM, PlpA or transposase A protein, polypeptide or peptide. The immunizing composition is administered in a manner effective to stimulate antibody producing cells. Rodents such as mice and rats are preferred animals, however, the use of rabbit, sheep frog cells is also possible. The use of rats may provide certain advantages (Goding, 1986, pp. 60-61), but mice are preferred, with the BALB/c mouse being most preferred as this is most routinely used and generally gives a higher percentage of stable fusions.

Following immunization, somatic cells with the potential for producing antibodies, specifically B lymphocytes (B cells), are selected for use in the MAb generating protocol. These cells may be obtained from biopsied spleens, tonsils or lymph nodes, or from a peripheral blood sample. Spleen cells and peripheral blood cells are preferred, the former because they are a rich source of antibody-producing cells that are in the dividing plasmablast stage, and the latter because peripheral blood is easily accessible. Often, a panel of animals will have been immunized and the spleen of animal with the highest antibody titer will be removed and the spleen lymphocytes obtained by homogenizing the spleen.
with a syringe. Typically, a spleen from an immunized mouse contains approximately $5 \times 10^7$ to $2 \times 10^8$ lymphocytes.

The antibody-producing B lymphocytes from the immunized animal are then fused with cells of an immortal myeloma cell, generally one of the same species as the animal that was immunized. Myeloma cell lines suited for use in hybridoma-producing fusion procedures preferably are non-antibody-producing, have high fusion efficiency, and enzyme deficiencies that render them incapable of growing in certain selective media which support the growth of only the desired fused cells (hybridomas).

Any one of a number of myeloma cells may be used, as are known to those of skill in the art (Goding, pp. 65-66, 1986; Campbell, pp. 75-83, 1984). For example, where the immunized animal is a mouse, one may use P3-X63/Ag8, X63-Ag8.653, NS1/1.Ag 4 1, Sp210-Ag14, PO, NSO/U, MPC-11, MPC11-X45-GTG 1.7 and S194/5XX0 Bul; for rats, one may use R210.RCY3, Y3-Ag 1.2.3, IR983F and 4B210; and U-266, GM1500-GRG2, LICR-LON-HMy2 and UC729-6 are all useful in connection with human cell fusions.

One preferred murine myeloma cell is the NS-1 myeloma cell line (also termed P3-NS-1-Ag4-1), which is readily available from the NIGMS Human Genetic Mutant Cell Repository by requesting cell line repository number GM3573. Another mouse myeloma cell line that may be used is the 8-azaguanine-resistant mouse murine myeloma SP2/0 non-producer cell line.

Methods for generating hybrids of antibody-producing spleen or lymph node cells and myeloma cells usually comprise mixing somatic cells with myeloma cells in a 2:1 proportion, though the proportion may vary from about 20:1 to about 1:1, respectively, in the presence of an
agent or agents (chemical or electrical) that promote the fusion of cell membranes. Fusion methods using Sendai virus have been described by Kohler and Milstein (1975; 1976), and those using polyethylene glycol (PEG), such as 37% (v/v) PEG, by Gefter et al. (1977). The use of electrically induced fusion methods is also appropriate (Goding pp. 71-74, 1986).

Fusion procedures usually produce viable hybrids at low frequencies, about $1 \times 10^{-6}$ to $1 \times 10^{-8}$. However, this does not pose a problem, as the viable, fused hybrids are differentiated from the parental, unfused cells (particularly the unfused myeloma cells that would normally continue to divide indefinitely) by culturing in a selective medium. The selective medium is generally one that contains an agent that blocks the de novo synthesis of nucleotides in the tissue culture media. Exemplary and preferred agents are aminopterin, methotrexate, and azaserine. Aminopterin and methotrexate block de novo synthesis of both purines and pyrimidines, whereas azaserine blocks only purine synthesis. Where aminopterin or methotrexate is used, the media is supplemented with hypoxanthine and thymidine as a source of nucleotides (HAT medium). Where azaserine is used, the media is supplemented with hypoxanthine.

The preferred selection medium is HAT. Only cells capable of operating nucleotide salvage pathways are able to survive in HAT medium. The myeloma cells are defective in key enzymes of the salvage pathway, e.g., hypoxanthine phosphoribosyl transferase (HPRT), and they cannot survive. The B cells can operate this pathway, but they have a limited life span in culture and generally die within about two weeks. Therefore, the only cells that can survive in the selective media are those hybrids formed from myeloma and B cells.
This culturing provides a population of hybridomas from which specific hybridomas are selected. Typically, selection of hybridomas is performed by culturing the cells by single-clone dilution in microtiter plates, followed by testing the individual clonal supernatants (after about two to three weeks) for the desired reactivity. The assay should be sensitive, simple and rapid, such as radioimmunoassays, enzyme immunoassays, cytotoxicity assays, plaque assays, dot immunobinding assays, and the like.

The selected hybridomas would then be serially diluted and cloned into individual antibody-producing cell lines, which clones can then be propagated indefinitely to provide MAbs. The cell lines may be exploited for MAb production in two basic ways. A sample of the hybridoma can be injected (often into the peritoneal cavity) into a histocompatible animal of the type that was used to provide the somatic and myeloma cells for the original fusion. The injected animal develops tumors secreting the specific monoclonal antibody produced by the fused cell hybrid. The body fluids of the animal, such as serum or ascites fluid, can then be tapped to provide MAbs in high concentration. The individual cell lines could also be cultured in vitro, where the MAbs are naturally secreted into the culture medium from which they can be readily obtained in high concentrations. MAbs produced by either means may be further purified, if desired, using filtration, centrifugation and various chromatographic methods such as HPLC or affinity chromatography.

The following examples are included to demonstrate preferred embodiments of the invention. It should be appreciated by those of skill in the art that the techniques disclosed in the examples which follow represent techniques discovered by the inventor to
function well in the practice of the invention, and thus can be considered to constitute preferred modes for its practice. However, those of skill in the art should, in light of the present disclosure, appreciate that many changes can be made in the specific embodiments which are disclosed and still obtain a like or similar result without departing from the spirit and scope of the invention.

EXAMPLE 1

Isolation and Characterization of Capsule Mutants

A. Methods

1. Bacterial strains, plasmids, and culture conditions

The bacterial strains and plasmids used are listed herein in Table 3. Culture conditions for *S. pneumoniae* and *E. coli* were previously described (Dillard and Yother, 1991). Erythromycin was used at 0.3 µg/ml and streptomycin was used at 100 µg/ml in *S. pneumoniae* cultures where indicated. Chloramphenicol was used at 1 µg/ml to detect transcription in *S. pneumoniae* isolates carrying pJY4163/4164 chromosomal insertions.

Table 3. Bacterial strains and plasmids.

<table>
<thead>
<tr>
<th>Strain/Plasmid</th>
<th>Derivation and properties</th>
<th>Source/Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strain <em>S. pneumoniae</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>WU2</td>
<td>Type 3 encapsulated</td>
<td>Briles et al. (1991b)</td>
</tr>
<tr>
<td>D39</td>
<td>Type 2 encapsulated</td>
<td>Avery et al. (1944)</td>
</tr>
<tr>
<td>Rx1</td>
<td>Non-encapsulated</td>
<td>Ravin (1959)</td>
</tr>
<tr>
<td></td>
<td>mutant of R36A-A66 hybrid</td>
<td>Shoemaker and</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Guild (1974)</td>
</tr>
<tr>
<td>L8-2006</td>
<td>Type 1 encapsulated</td>
<td>Dillard and</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Yother (1994)</td>
</tr>
<tr>
<td>DBL5</td>
<td>Type 5 encapsulated</td>
<td>Yother et al. (1982)</td>
</tr>
<tr>
<td>Strain/Plasmid</td>
<td>Derivation and properties</td>
<td>Source/Reference</td>
</tr>
<tr>
<td>---------------</td>
<td>---------------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>BG9273</td>
<td>Type 6A encapsulated</td>
<td>Dillard and Yother (1994)</td>
</tr>
<tr>
<td>EP2809</td>
<td>Type 8 encapsulated</td>
<td>Dillard and Yother (1994)</td>
</tr>
<tr>
<td>BG5862</td>
<td>Type 9 encapsulated</td>
<td>Dillard and Yother (1994)</td>
</tr>
<tr>
<td>LM100</td>
<td>Type 22 encapsulated</td>
<td>Dillard and Yother (1994)</td>
</tr>
<tr>
<td>A66R₂</td>
<td>Non-encapsulated mutant of A66 (Type 3)</td>
<td>Muckerman et al. (1982)</td>
</tr>
<tr>
<td>661</td>
<td>Non-encapsulated mutant of A66 (Type 3)</td>
<td>Bernheimer and Wermundsen (1972)</td>
</tr>
<tr>
<td>JD531</td>
<td>Non-encapsulated mutant of WU2, Em⁺, cpsA1</td>
<td>This work</td>
</tr>
<tr>
<td>JD541</td>
<td>Non-encapsulated mutant of WU2, Em⁺, cpsA2</td>
<td>This work</td>
</tr>
<tr>
<td>JD542</td>
<td>Non-encapsulated mutant of WU2, Em⁺, cpsB1</td>
<td>This work</td>
</tr>
<tr>
<td>10 JD551</td>
<td>Non-encapsulated mutant of WU2, Em⁺, cpsB2</td>
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</tr>
<tr>
<td>JD571</td>
<td>Non-encapsulated mutant of WU2, Em⁺, cpsB3</td>
<td>This work</td>
</tr>
<tr>
<td>JD600</td>
<td>WU2 Str⁺</td>
<td>This work</td>
</tr>
<tr>
<td>JD611</td>
<td>JD600 × JD531, Em⁺, Str⁺, cpsA1</td>
<td>This work</td>
</tr>
<tr>
<td>JD614</td>
<td>JD600 × JD551, Em⁺, Str⁺, cpsB2</td>
<td>This work</td>
</tr>
<tr>
<td>15 JD619</td>
<td>JD600 × JD541, Em⁺, Str⁺, cpsA2</td>
<td>This work</td>
</tr>
<tr>
<td>JD692</td>
<td>JD600 × JD542, Em⁺, Str⁺, cpsB1</td>
<td>This work</td>
</tr>
<tr>
<td>JD816</td>
<td>JD600 × JD671, Em⁺, Str⁺, cpsB3</td>
<td>This work</td>
</tr>
<tr>
<td>JD636</td>
<td>WU2 Str⁺, RM⁺, Nov⁺</td>
<td>This work</td>
</tr>
<tr>
<td><strong>Strain/Plasmid</strong></td>
<td><strong>Derivation and properties</strong></td>
<td><strong>Source/Reference</strong></td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------</td>
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</tr>
<tr>
<td>JD752</td>
<td>Isolate of transformation pool W62 x JD811, type 3 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD770</td>
<td>pJD330 x WU2, type 3 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD803</td>
<td>JD770 x D39, type 3 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD871</td>
<td>pJD366 x D39, type 2 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD872</td>
<td>JD871 x WU2, type 2 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD875</td>
<td>pJD366 x DBL5, type 5 encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td>JD908</td>
<td>pJD369 x WU2, non-encapsulated, Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This work</td>
</tr>
<tr>
<td><em>E. coli</em> LE392</td>
<td>F&lt;sup&gt;-&lt;/sup&gt;hsdR514 (T&lt;sub&gt;4&lt;/sub&gt; &lt;sup&gt;+&lt;/sup&gt; M&lt;sub&gt;K&lt;/sub&gt; &lt;sup&gt;-&lt;/sup&gt;) supE44 supF&lt;sup&gt;E&lt;/sup&gt; supF&lt;sup&gt;L&lt;/sup&gt; λ (faciZ)&lt;sub&gt;Y&lt;/sub&gt; 6 galK2 galT22 met21 trpR55 λ</td>
<td>Tilghman et al. (1977)</td>
</tr>
<tr>
<td>10 pJY4163 and pJY4164</td>
<td>Lack origin of replication for <em>S. pneumoniae</em> Promoterless cat gene downstream of multiple cloning site (opposite orientations in 4163, 4164), Em&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Yother et al. (1992)</td>
</tr>
<tr>
<td>pJD330</td>
<td>WU2 Sau3A1 fragment cloned into pJY4163 BamHI site, isolated from JD752</td>
<td>This work</td>
</tr>
<tr>
<td>pJD337</td>
<td>pJY4163:: 1.5kb XbaI-PvuII fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>15 pJD343</td>
<td>pJY4164:: 0.4kb MucI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD345</td>
<td>pJY4164:: 1.1kb MucI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD351</td>
<td>pJY4164:: 2.4kb Sau3AI-Sau3AI fragment of pJD330 orientation opposite pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>Strain/Plasmid</td>
<td>Derivation and properties</td>
<td>Source/Reference</td>
</tr>
<tr>
<td>---------------</td>
<td>---------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>pJD353</td>
<td>pJY4164:: 1.6kb Sau3AI-XbaI fragment of pJD330</td>
<td>This work</td>
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<tr>
<td>pJD357</td>
<td>pJY4164:: 0.3kb MunI-Sau3AI fragment of JD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD359</td>
<td>pJY4164:: 0.6kb PvuII-HaeIII fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD361</td>
<td>pJY4164:: 0.45kb XbaI-PstI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD362</td>
<td>pJY4164:: 0.4kb HaeIII-MunI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD364</td>
<td>WU2 3.2kb HindIII fragment cloned into pJY4164 HindIII site</td>
<td>This work</td>
</tr>
<tr>
<td>pJD366</td>
<td>WU2 3.2kb HindIII fragment cloned into pJY4164 HindIII site, orientation opposite pJD364</td>
<td>This work</td>
</tr>
<tr>
<td>pJD368</td>
<td>pJD4164:: 0.45kb Rsai-MunI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD369</td>
<td>pJY4164:: 0.55 kb PvuII-MunI fragment of pJD330</td>
<td>This work</td>
</tr>
<tr>
<td>pJD374</td>
<td>WU2 1.2kb Sau3AI fragment cloned in pJY4163</td>
<td>This work</td>
</tr>
<tr>
<td>pJD377</td>
<td>pJY4164:: 1.2kb SacI-HindIII fragment of pJD364</td>
<td>This work</td>
</tr>
<tr>
<td>pJD380</td>
<td>pJY4164:: 0.36kb Sau3AI-SspI fragment of pJD330</td>
<td>This work</td>
</tr>
</tbody>
</table>

Em<sup>R</sup>, erythromycin resistant; Em<sup>S</sup>, erythromycin sensitive, Str<sup>R</sup>, streptomycin resistant; Rif<sup>R</sup>, rifampicin resistant; Nov<sup>R</sup>, novobiocin resistant.
2. General DNA techniques

Techniques for DNA fragment isolation, ligations, and plasmid isolation and purification were performed as previously described (Dillard and Yother, 1991; and as described by Sambrook et al., 1989, the relevant portions incorporated herein by reference). Plasmid screenings were done by scraping colonies from agar plates and incubating these in the lysis solution of Kado and Liu, 3% SDS, 50 mM Tris, pH 12.6 (Kado and Liu, 1981). The lysates were run on agarose gels to determine plasmid sizes.

3. Library construction

A plasmid library of random fragments was constructed by digesting chromosomal DNA from S. pneumoniae strain WU2 to completion with Sau3AI and ligating these fragments into the BamHI site of pJY4163. The resulting ligation mixture was electroporated into E. coli LE392, and transformants were selected on L agar plates containing 300 μg erythromycin/ml. Individual colonies were patched on erythromycin plates. Each plate contained 100 colonies and constituted a pool. Transformants were pooled by scraping the plates.

4. Transformations

Encapsulated strains of S. pneumoniae were induced to competence as has been described in Yother et al., 1986, incorporated herein by reference. Non-encapsulated strains were made competent for transformation by growth in Todd Hewitt broth (Difco, Detroit, MI) supplemented with 0.01% CaCl₂, 0.5% BSA, and 0.5% yeast extract. S. pneumoniae cells were allowed to express transforming DNA for 2 h. before plating on agar medium. Electroporation of H₂O washed E. coli LE392 cells resuspended in 10% glycerol was performed in a BTX Electro Cell Manipulator 600 according to the
instructions of the manufacturer (Biotechnologies and Experimental Research, Inc., San Diego, CA).

5. Preparation of *S. pneumoniae* chromosomal DNA

Cultures of *S. pneumoniae* (100 ml) were grown to stationary phase in the presence of 1% choline chloride to prevent autolysis (Briese and Hakenbeck, 1983). The bacteria were centrifuged at 5000 rpm for 10 min and resuspended in 2.5 ml TE buffer (10 mM TrisHCl, 1 mM EDTA, pH 8.0). SDS was added to 1%, and the cells were lysed at 65°C for 15 min. One fifth volume 5 M KOAc (pH 8) was added, and incubation was continued at 65°C 15 min, followed by incubation on ice for 60 min. Cell debris was removed by centrifugation at 10,000 rpm for 10 min, the supernatant was added to 2 volumes of ethanol, and the DNA was hooked out with a glass rod. The DNA was dried, resuspended in TE, and further purified by CsCl/ethidium bromide buoyant density gradient centrifugation (Radloff et al., 1967).

6. Recovery of plasmids resolved from *S. pneumoniae* chromosomes.

A 10 ml culture of late log phase *S. pneumoniae* was centrifuged at 5000 rpm for 10 min. The supernatant was removed, and the cells were resuspended in 100 µl lysis buffer (Saunders and Guild, 1980). Following a 5 min incubation at 37°C, 900 µl of Birnboim and Doly solution I was added, and the rest of the alkaline lysis procedure was carried out as for *E. coli* (Birnboim and Doly, 1979). The resulting preparation contained very little plasmid DNA and was therefore electroporated into *E. coli* where significant quantities of plasmid could be obtained and isolated as described (Birnboim and Doly, 1979).

7. Mapping by chromosomal transformation

The integration frequency was used to determine the linkage of spontaneous mutations. Chromosomal DNA from a
streptomycin resistant derivative of the capsule mutant was used to transform the other mutants. The number of encapsulated transformants obtained divided by the number of streptomycin resistant transformants obtained is defined as the integration frequency (Bernheimer and Wermundsen, 1972). Integration frequencies of 0.02 to 0.03 resulted from crosses between mutants carrying mutations in the same locus. Integration frequencies of about 0.3 indicated the mutations were in different loci.

Linkage of plasmid insertions to capsule mutations was determined by using chromosomal DNA from wild type strains carrying non-destructive plasmid insertions to transform capsule mutants. Transformants were selected on erythromycin and screened for encapsulation. The co-transformation frequency determined the degree of linkage.

8. Capsule detection by ELISA

To obtain a crude capsule extract, a 10 ml culture of log phase S. pneumoniae cells was centrifuged at 5000 rpm for 20 min, and the supernatant was removed. The pellet was resuspended in 1 ml PBS (50 mM sodium phosphate pH 7.4, 100 mM NaCl). Protein content was determined on 100 µl of washed cells using the Bio-Rad kit (Bio-Rad, Richmond, CA). The remaining cells were extracted with an equal volume of 24:1 chloroform/isooamyl alcohol. Following centrifugation, the aqueous phase was precipitated with two volumes of ethanol. The precipitate was resuspended in 1 ml PBS, then treated with RNase at 1 mg/ml and DNase at 0.2 mg/ml for 3 h at 37°C, and pronase at 1 mg/ml for 2 h at 37°C. The extract was then re-extracted and precipitated as before. The precipitate was resuspended in PBS and used to coat the wells of a microtiter plate for ELISA analysis. Values were normalized to protein content. ELISAs were
performed by the standard technique (Ausubel et al., 1987). Monoclonal antibody 16.3 was used to detect type 3 capsular polysaccharide (Briles et al., 1981a). ELISA plates were read at 405 nm in a Biotek model 320 plate reader (Bio-Tek Instruments, Winooski, VT).

9. Percoll gradient centrifugation

For density determinations, 10 ml of log phase cells were centrifuged at 4000 rpm for 10 min, washed once with water, and then resuspended in 1 ml water. A volume of 300 μl of cells was loaded on top of a 10 ml 0-100 or 25-100% continuous Percoll gradient. Gradient density marker beads were loaded on top of the gradients. The gradients were centrifuged at 10,000 rpm for 15 min with the brake off. Percoll and density marker beads were purchased from Pharmacia (Piscataway, NJ). Non-encapsulated strains of S. pneumoniae exhibit a higher density in Percoll gradients than encapsulated strains (Briles et al., 1992). Percoll gradients were also used to enrich for encapsulated cells expected to result from low frequency events. Percoll gradients were used to obtain binary capsule type transformants and to enrich for spontaneous revertants to capsule production.

B. Results

To identify the type 3 capsule region of S. pneumoniae, a Sau3AI library of fragments was cloned from the type 3 encapsulated strain WU2 to direct insertions into the chromosome of strain WU2. The library used in the insertion-duplication mutagenesis procedure was constructed by cloning random Sau3AI fragments in the plasmid pJY4163, which carries an erythromycin-resistance marker. Since this plasmid is unable to replicate in S. pneumoniae, all erythromycin-resistant transformants should contain insertions at the chromosomal site of the target Sau3AI fragment (Morrison et al., 1984). By
transforming the library of clones into strain WU2, the inventors obtained 5 non-mucoid isolates among 491 erythromycin-resistant transformants. However, further studies involving transformation of the parent strain with either chromosomal DNA from the mutants or plasmids recovered from the mutants showed that the plasmid insertions were neither linked to nor responsible for the capsule mutations.

To determine if the non-mucoid isolates were truly deficient in the production of type 3 capsular polysaccharide, several methods were employed. In slide-agglutination assays, none of the five mutants reacted with polyclonal antisera specific for type 3 polysaccharide. Centrifugation through Percoll density gradients revealed that the mutant strains were much denser than the encapsulated parent strain. WU2 cells had a density \(<1.01 \text{ g/ml}\), whereas all five mutants migrated at 1.09 to 1.10 \text{ g/ml}. These data suggested that complete capsules were not produced by the mutants. However, these tests might not reveal the presence of short or sparse polysaccharide chains on the cell surfaces or capsular material not translocated to the surfaces. To determine if such material was present, ELISA analysis was carried out on crude cell extracts. Capsular material in the extracts was detected using a monoclonal antibody directed against type 3 polysaccharide. The analysis indicated that mutants JD531 and JD541 (designated mutants A1 and A2 respectively) made no detectable capsular material, whereas mutants JD542, JD551, and JD571 (designated mutants B1, B2 and B3 respectively) made significant levels of reactive material. The common laboratory strain Rx1 was also found to produce significant levels of type 3 capsular material (FIG. 1). Although Rx1 is generally referred to as a non-encapsulated derivative of the type 2 strain D39, it was transformed three times
with chromosomal DNA from derivatives of the type 3 strain A66, chosen twice for type 3 encapsulation, and chosen finally for non-encapsulation (Ravin, 1959; Shoemaker and Guild, 1974).

No capsular material could be detected by ELISA in the culture supernatant fluids of mutants JD531 and JD541, indicating that these mutants were not merely defective in attachment of the polysaccharide. Only low levels of capsular material were detected in supernatants of Rx1, JD542, JD551, and JD571 cultures.

The five mutations resulting in the capsule-deficient phenotype were mapped to two loci by chromosomal transformation. Reciprocal crosses between the mutants yielded encapsulated transformants for each combination, but not for transformation of a mutant with DNA from the same strain. The mutations were thereby determined to be genotypically distinct. The transformations also revealed that the mutations in JD531 and JD541 were more closely linked to each other than to the mutations in JD542, JD551, and JD571. Likewise, the mutations in JD542, JD551, and JD571 were more closely linked to one another than to the other two mutations.

The integration frequencies for those mutations judged to be closely linked were similar (0.02 to 0.03) and were ten-fold lower than those judged to be not as closely linked. The genotypic data thus agreed with the phenotypic data, i.e., the two mutations leading to total loss of capsule synthesis mapped together, and the three mutations causing lack of proper capsular polysaccharide processing mapped together. The loci containing these mutations were temporarily designated cpsA and cpsB, respectively, and the mutations were named as indicated in FIG. 1.
Table 4 shows the transformation frequencies of capsule mutations. *S. pneumoniae* strains were transformed with chromosomal DNA from strain JD636, a streptomycin resistant WU2 (Table 3), and streptomycin-resistant transformants were selected. Transformation frequencies are calculated from cultures not induced to competence. With optimal induction, strain WU2 may exhibit a transformation frequency approaching that of the non-encapsulated mutants. However, during the mutagenesis procedure, sub-optimal transformation frequencies were observed (0.0003 to 0.006%). JD908 (Table 3) was also included, it contains an insertion mutation resulting in loss of capsule expression (described in FIG. 4). It would appear that the non-encapsulated mutants are highly transformable, suggesting that the reason for their over-representation in the original transformant population was because of their transformability. The mutagenesis procedure, by selecting for transformability, has enriched for mutants already deficient in capsule (Table 4).

### Table 4. Transformation frequencies of capsule mutants.

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Mutation</th>
<th>Total cfu</th>
<th>Str&lt;sup&gt;R&lt;/sup&gt; cfu&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Transformation frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JD531</td>
<td>cpsA1</td>
<td>2.0 x 10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>5.0 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.3</td>
</tr>
<tr>
<td>JD541</td>
<td>cpsA2</td>
<td>1.4 x 10&lt;sup&gt;8&lt;/sup&gt;</td>
<td>7.6 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.6</td>
</tr>
<tr>
<td>JD542</td>
<td>cpsB1</td>
<td>2.4 x 10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>20 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.8</td>
</tr>
<tr>
<td>JD551</td>
<td>cpsB2</td>
<td>2.0 x 10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>4.6 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.2</td>
</tr>
<tr>
<td>JD571</td>
<td>cpsB3</td>
<td>2.4 x 10&lt;sup&gt;8&lt;/sup&gt;</td>
<td>5.0 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.2</td>
</tr>
<tr>
<td>WU2</td>
<td>wt</td>
<td>0.0 x 10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>1</td>
<td>0.0000</td>
</tr>
<tr>
<td>JD908</td>
<td>cpsS</td>
<td>10 x 10&lt;sup&gt;5&lt;/sup&gt;</td>
<td>10 x 10&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.1</td>
</tr>
</tbody>
</table>

<sup>b</sup> Str<sup>R</sup> streptomycin resistant.
EXAMPLE 2

Identification of a Clone Containing a Capsule Gene

To identify DNA fragments capable of repairing the cpsA1 mutation, JD611, a derivative of JD531 lacking the pJY4163 insertion, was transformed with pools of pJY4163 clones containing Sau3AI fragments from strain WU2. Transformations and DNA manipulations were performed as described in Example 1. In this insertion-duplication restoration procedure, the plasmid clone is inserted into the mutant chromosome, leading to duplication of the homologous target fragment and restoration of one wild type copy of cpsA (FIG. 2). Erythromycin-resistant transformants were screened visually for the mucoid phenotype. One plasmid clone was identified which restored encapsulation in the cpsA1-containing mutant. Due to the duplication of the target fragment, the plasmid insertion could resolve out of the chromosome by homologous recombination at low frequency. Therefore, transformation of E. coli with DNA from the encapsulated transformant and selection for erythromycin-resistance allowed recovery of the plasmid, designated pJD330, that had repaired the cpsA1 defect.

Transformation of the capsule-deficient mutants with pJD330 suggested that the clone contained part of cpsA. When pJD330 was inserted into the chromosome of the cpsA1-containing mutant, 56% of the erythromycin-resistant transformants became encapsulated (Table 5). The failure of the remainder of the transformants to become encapsulated indicated that the cloned fragment contained only one end of the gene. The site of the crossover relative to the site of the mutation determines whether the mutation will be located in the incomplete copy of the gene or the full-length copy. If the recombination occurs on the left, as shown in FIG. 2, the full-length gene is wild type and capsule
is restored. However, if the crossover occurs on the right, the mutation is located in the full-length copy, and no capsule is obtained. This interpretation is supported by the observation that transformants of the cpsA1 mutant which incorporated the plasmid but did not become encapsulated, spontaneously gave rise to encapsulated, erythromycin-resistant colonies, either by excision and reinsertion of the plasmid or by gene conversion. The cpsA2 defect was not repaired by pJD330, suggesting that the site of this mutation was either not present on the plasmid clone or was located too near the end of the clone for crossover to repair the defect.

Table 5. Restoration of encapsulation with pJD330.a

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Mutation</th>
<th>Ery^R cfu^b</th>
<th>Cps^+ cfu</th>
<th>Cps^+ frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JD611</td>
<td>cpsA1</td>
<td>475</td>
<td>267</td>
<td>56</td>
</tr>
<tr>
<td>JD619</td>
<td>cpsA2</td>
<td>26</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>JD692</td>
<td>cpsB1</td>
<td>13</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>JD614</td>
<td>cpsB2</td>
<td>124</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Rx1</td>
<td>cps^-</td>
<td>158</td>
<td>49</td>
<td>31</td>
</tr>
<tr>
<td>661</td>
<td>capD6</td>
<td>56</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>A66R2</td>
<td>capD4</td>
<td>4</td>
<td>3</td>
<td>75</td>
</tr>
</tbody>
</table>

a Mutants deficient in capsule production were transformed with pJD330 DNA. Transformants were plated on erythromycin to select for those containing chromosomal insertions of pJD330. Erythromycin-resistant transformants were screened for mucoldy. Cps^+ frequency is the ratio of Cps^+/Ery^R cfu.

b Ery^R. Erythromycin resistant.

Transformation of pJD330 into the cpsB-containing mutants did not restore any of these to encapsulation,
suggesting that \textit{cpsB} is not present on pJD330. However, strain Rx1 was restored to type 3 encapsulation by pJD330 (Table 5).

\textbf{EXAMPLE 3}

cpsA Codes for UDP-Glucose Dehydrogenase

Transformation of two previously characterized mutants suggested that pJD330 contains part of the gene for UDP-glucose dehydrogenase. UDPG dehydrogenase is the enzyme which converts UDP-glucose (UDPG) into UDP-glucuronic acid (UDPGA). UDPG and UDPGA are the two nucleotide sugars which are required for type 3 capsule synthesis (Smith et al., 1960). The non-encapsulated mutants A66R$_2$ and 661 were previously shown to be deficient in the production of UDPG dehydrogenase due to mutations in the locus designated \textit{capD} (Bernheimer and Wermundsen, 1972). Transformation of A66R$_2$ (\textit{capD4}) with pJD330 restored encapsulation, whereas transformation of 661 (\textit{capD6}) did not (Table 5).

Transformations with chromosomal DNA (Table 6) confirmed that the other \textit{cpsA} and \textit{capD} mutations were closely linked to the region cloned in pJD330.

Transformations and other DNA manipulations were performed as described in Example 1. Strain JD770 was obtained by inserting pJD330 into the chromosome of the parental strain WU2. JD770 was found to produce wild type amounts of type 3 capsule. Chromosomal DNA from JD770 was used to transform those mutants which could not be restored to encapsulation by pJD330 (Table 6). The \textit{cpsA2} and UDPG dehydrogenase mutation \textit{capD6} were found to be >90\% linked to the plasmid insertion.
Table 6. Restoration of encapsulation with chromosomal DNA linked to pJD330 insertion.\textsuperscript{a}

<table>
<thead>
<tr>
<th>Recipient</th>
<th>Mutation</th>
<th>Ery\textsuperscript{R} cfu\textsuperscript{b}</th>
<th>Cps\textsuperscript{+} cfu</th>
<th>Cps\textsuperscript{+} frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JD692</td>
<td>cpsB1</td>
<td>17</td>
<td>13</td>
<td>70</td>
</tr>
<tr>
<td>JD614</td>
<td>cpsB2</td>
<td>79</td>
<td>58</td>
<td>73</td>
</tr>
<tr>
<td>JD619</td>
<td>cpsA2</td>
<td>42</td>
<td>39</td>
<td>93</td>
</tr>
<tr>
<td>661</td>
<td>capD6</td>
<td>6</td>
<td>6</td>
<td>100</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Transformants were plated on erythromycin to select for those which had incorporated the region containing the insertion. Erythromycin-resistant transformants were screened for mucoidy. Cps\textsuperscript{+} frequency is the ratio of Cps\textsuperscript{+}/Ery\textsuperscript{R} cfu.

\textsuperscript{b} Ery\textsuperscript{R} -- erythromycin resistant.

Deletion analysis was performed to more closely localize the sites of the mutations cpsA1, capD4, and the mutation in Rx1 (cps\textsuperscript{-}). By transforming with plasmid subclones and making no selection for insertion of the plasmids, the inventors were able to observe recombination events that occurred as a result of double crossovers between the cloned fragment and its homolog in the chromosome (FIG. 3A). Transformations with several subclones revealed that the sites of the mutations could all be localized to a 250 bp PvuII-SspI fragment common to pJD380 and pJD369 (FIG. 3B). The fact that the same fragment which restores encapsulation in a UDPG dehydrogenase mutant also restores encapsulation in the cpsA1-containing mutant suggests that cpsA encodes UDPG dehydrogenase. From here on, the cpsA loci is designated cps3D (see Example 9).

It is known that transformation of UDPG dehydrogenase mutants, including 661, with chromosomal
DNA from a type 1 strain restored type 3 encapsulation by incorporation of the type 1 specific genes at a site other than that occupied by the type 3 genes (Bernheimer et al., 1967, Bernheimer and Wermundsen, 1972). Type 1 capsular polysaccharide contains galacturonic acid; therefore, type 1 strains are expected to produce UDPG dehydrogenase (Austrian et al., 1959). When UDPG dehydrogenase mutants of a type 3 strain were transformed with DNA from type 1 strains, the UDPG dehydrogenase from type 1 complemented the type 3 mutation, allowing the production of both capsular polysaccharides. The UDPG dehydrogenase gene from the type 1 strain was never observed to repair the mutation in the type 3 gene (Bernheimer and Wermundsen, 1969). When the cpsA1 mutant JD611 was transformed with chromosomal DNA from a type 1 strain, type 3 encapsulated transformants were obtained at a frequency of $3 \times 10^{-6}$. This frequency is in agreement with that observed for transformation of mutant 661 (capD6) to binary encapsulation (Bernheimer and Wermundsen, 1972) and above the spontaneous reversion frequency ($< 8 \times 10^{-9}$).

**EXAMPLE 4**

**Genetic and Physical Map of the Type 3 Capsule Region**

**A. Methods**

1. Southern Blotting

   Southern blotting was performed using the vacuum blotter and chemiluminescent detection system purchased from Stratagene (La Jolla, CA). The PosiBlot® 30-30 pressure blotter is part of an integrated system designed to transfer DNA or RNA from agarose gels quickly and efficiently onto solid support matrices, such as Stratagene’s hybridization membranes including the Nitrocellulose membranes, Duralose-UV™ membranes (reinforced nitrocellulose), Duralon-UV™ membranes (nylon) or Illuminator™ nylon membranes.
Following electrophoresis the gels are stained in 5 μg/ml of ethidium bromide (EtBr) in water, destained in water and then photographed. Prior to Southern transfer, the gels are pretreated by depurination, denaturation and neutralization. Depurination entails treating the gels with 0.25 N HCl for 5-30 minutes with gentle shaking. Denaturation consists of pouring off the HCl and adding a 0.5 N NaOH and 1.5 M NaCl denaturation solution, enough to cover the gel. The gels are treated for 5 minutes to one hour with gentle shaking. Neutralization involves pouring off the denaturation solution and adding a 0.1 M Tris-HCl (pH 7.5) and 1.5 M NaCl neutralization solution, enough to cover the gel. They are then treated for 5 minutes to one hour with gentle shaking.

Gels are then ready for blotting, which is performed with gloved hands. The membrane is pre-wet in distilled water (dH₂O) and then in transfer buffer for 5 minutes. 10x SSC buffer-10x SSPE buffer or 25 mM sodium phosphate (pH 6.5) is the transfer buffer for nylon membranes. For nitrocellulose or Duralose-UV membranes, 20x SSC buffer should be used.

The membrane and gel are set up in the Posiblot 30-30 pressure blotter and pressure exerted and adjust to 75 mm Hg. Blotting times vary for different gels and depend on the amount and size of the nucleic acid; size, thickness and percentage of gel; and depth of gel wells and volume of sample loaded on the gel; which are routinely optimized. After the allotted blotting time, the position of the wells on the membrane is marked and the gel removed. The gel is generally stained and destained in ethidium bromide to check the efficiency of transfer. The membrane is removed from the device and placed on clean Whatman 3MM paper to allow the excess buffer to be absorbed. Once the membrane is free of standing liquid, but still damp, the membrane and the
Whatman 3MM paper is placed under a UV light and crosslinked. Alternatively, dry the membrane in a 80°C drying oven for 1-2 hours prior to crosslinking.

Boehringer Mannheim's Genius™ Nonradioactive Detection System, a chemiluminescence-based, nucleic acid detection kit, permits fast, safe and sensitive detection of DNA and RNA immobilized on nylon membranes. As little as 0.1 pg of target plasmid DNA can be detected in a 30-minute exposure of the processed blot to X-ray film or, in a similar exposure time, 1 pg of a single-copy gene can be detected in less than 1.0 µg of genomic DNA. The Nonradioactive Detection System can also be used for rapid Northern-blot analysis of RNA.

After transfer and crosslinking, the membrane is prehybridized for 1 hour at 42°C. The labeled probe is placed in a microfuge tube containing 100 µl of sonicated salmon sperm DNA (10 mg/ml) stock and heated in a boiling water bath for 5 minutes. This is pulse-spun to collect condensation and stored on ice until ready to add to hybridization. The probe is added to prehybridization solution and hybridized, with shaking, overnight at 42°C using standard hybridization solutions as described by the Genius® protocol. This is washed once for 15 minutes at room temperature in 0.1x SSC/0.1% SDS and then washed twice for 15 minutes at 60°C in 0.1x SSC/0.1% SDS for each wash. The probe is then ready for detection. The BRL 1 Kb DNA ladder was used as a molecular weight size standard (Bethesda Research Laboratories, Gaithersburg, MD). Biotin labeled probes for hybridization were prepared by nick-translation using the BRL BioNick kit (Bethesda Research Laboratories). High stringency conditions, as described above should result in the detection of sequences ≥95% homologous to the probe. Reduced stringency was achieved by lowering the wash, or hybridization and wash temperatures to room temperature.
At reduced stringency, sequences with 85% homology to the probe should have been detectable.

B. Results

Using pJD330 as a probe in Southern hybridizations, a physical map of the type 3 capsule region of strain WU2 was developed (FIG. 4). Using the information gained from the chromosomal mapping, the inventors identified and cloned the HindIII fragment located to the right of the pJD330 insert. HindIII fragments approximately 3 kb in size were cloned from the WU2 chromosome into pJY4164. By using pJD330 to screen for homology, a clone containing a 3.2 kb insert was identified. This clone, pJD366, was then used to determine the location of the cpsB mutants.

Transformation mapping using JD770 showed that the cpsB mutations were about 74% linked to the pJD330 insertion (Table 6). This high frequency indicated that cpsB might be adjacent to the fragment contained in pJD330. When pJD366 was used to transform strains containing the cpsB mutations, encapsulation was not restored. Insertion of pJD366 into the WU2 chromosome did not alter the production of type 3 capsule; therefore, the inventors were able to examine linkage of the insertion to the cpsB mutations and determine the relative location of cpsB. The pJD366 insertion was found to be only 25% linked to the cpsB mutations (as compared to 74% for the pJD330 insertion), suggesting that cpsB is located to the left of the pJD330 insert, as shown in FIG. 4.

In order to localize regions necessary for capsule production, insertion mutations using subclones of pJD330 and pJD366 were made. Transformation of strain WU2 with plasmids containing fragments internal to a gene or operon required for capsule production should result in
loss of encapsulation. Insertion of the plasmid containing the Sau3AI - XbaI fragment resulted in loss of encapsulation, indicating that this entire 1.6 kb fragment is within a single gene or operon required for capsule synthesis. Likewise, all insertions within this region eliminated capsule production (FIG. 4). Insertion of the plasmid containing the XbaI-PstI fragment did not disrupt capsule production, indicating that the end of the gene or operon is contained within this fragment.

None of the other insertions resulted in loss of capsule, indicating they were not internal to genes or operons required for capsule synthesis.

Since the plasmids used for the chromosomal insertions contain a promoterless cat gene, the inventors were able to establish the directions of transcription at the insertion sites. All insertions which contained the cat gene in the orientation to detect transcription proceeding to the right (as drawn in FIG. 4) resulted in chloramphenicol resistance. No transcription was detected in the opposite direction (FIG. 4).

EXAMPLE 5

Homology with Other Capsule Types

If the type-specific genes for capsule production are contained within a cassette, as has been proposed, these genes should show little homology to the type-specific genes from other capsule types. A high degree of homology should exist in the regions flanking the type-specific region (Austrian et al., 1959; Bernheimer and Wermundsen, 1972). The flanking regions may contain common genes necessary for production of capsule of any type or might not be involved in capsule production.
To determine if the regions cloned in pJD330 and pJD366 are specific to type 3 or are present in strains of other capsule types, HindIII digested chromosomal DNAs from strains of types 2, 3, 5, 6A, 8, 9, and 22 were Southern blotted and probed with these plasmids or fragments thereof. The fragment contained in pJD330 (the probe used was pJD351, containing 2.4 kb Sau3A1 fragment from pJD330) hybridized only with DNA from the type 3 strain, detecting the expected bands at 2.2 and 3.2 kb. No hybridization with the chromosomal DNA of the other six serotypes was detected, nor could the stringency be sufficiently lowered to detect homology in these strains.

When HindIII digests of chromosomal DNAs from these same strains were probed with the HindIII fragment from pJD366, the expected 3.2 kb band was observed in the type 3 strain, but a 1.1 kb band was found in every other capsule type. Probing with subclones of pJD366 containing the 2.1 kb HindIII-SacI fragment or the 1.2 kb SacI-HindIII fragment revealed that the homology resided in the more distal 1.2 kb fragment. Therefore, unlike the remaining 4.2 kb of DNA, which could be detected only in the type 3 strains, the 1.2 kb SacI-HindIII fragment (pJD377) showed a high degree of homology and could be detected at high stringency in all strains (2, 3, 5, 6A, 8, 9 and 22). This result suggests that this region may be the highly homologous flanking DNA predicted by the model to be adjacent to the type-specific genes.

**EXAMPLE 6**

**Transformation of Capsule Type**

To determine if all the type-specific genes necessary for the production of type 3 capsular polysaccharide were closely linked on the pneumococcal chromosome, strain JD770 was used as a donor in transformation of the type 2 strain D39. Laboratory
techniques were as described in Example 1. Seventy-three erythromycin-resistant transformants were obtained, and all 73 expressed type 3 capsule. No type 2 capsule could be detected by agglutination with type 2 specific antisera. Using chromosomal DNA from strain JD770, successful transformation of strains of type 5 and type 6B to type 3 encapsulation was also performed (Example 18).

By transforming the type 2 strain D39 with pJD366, isolates were obtained with the erythromycin-resistance marker closely linked to the type 2 capsule genes. These transformants were the result of recombination between the flanking regions of homologous non-type-specific DNA. Using DNA from one of these isolates, JD871, to transform the type 3 strain WU2 resulted in 95% co-transformation of type 2 encapsulation with erythromycin resistance. The remaining 5% were found to be type 3 encapsulated, indicating that only the flanking DNA or the plasmid alone was transferred to these isolates. Insertion of pJD366 into the type 5 strain DBL5 also resulted in a strain - JD875 - with the erythromycin resistance marker linked to the type-specific genes. This strain was successfully used to transform WU2 to type 5 encapsulation.

EXAMPLE 7
Direct Test of the Cassette Model

Transformations and other DNA manipulations were performed as described in Example 1. Southern Blotting was performed as described in Example 4.

If capsule type change involves a cassette-type recombination mechanism, then transformation of capsule type should result in replacement of the recipient's type-specific genes by those of the donor. In order to determine if such replacement does occur, DNA was used
from the type 3-specific region to probe *HindIII* digested chromosomal DNA from a strain which was originally type 2 and was transformed to type 3 (JD803), and a strain which was originally type 3 and was transformed to type 2 (JD872) (FIG. 5).  

*Hind III* digested chromosomal DNA from strain: 2 (D39 and JD871 from Example 6); 3 (WU2); 3/2 (JD872) and; 2/3 (JD803), were used in Southern blotting. First of all the Southern blot was probed with pJD343 and pJD368 Together these plasmids contain an 800 bp region (*HaeIII-MunI*) specific to type 3 and internal to *cpsS* (FIG. 5). The type 3 parent WU2 contained the expected 2.4 kb *HindIII* fragment specific to type 3, whereas neither the type 2 parent D39 nor its derivative JD871, which has pJD366 inserted into the chromosome, contained this fragment. When JD871 (type 2) was used to transform WU2, the resulting strain JD872 was type 2 encapsulated and had lost the 2.4 kb type 3-specific fragment. Similarly, when D39 was transformed with DNA from JD770 (type 3), the resulting strain JD803 was type 3 encapsulated and had gained the type 3-specific fragment.  

Reprobing of the same blot with the 1.2 kb Saci-*HindIII* fragment common to all capsule types (pJD377), revealed that the 1.1 kb *HindIII* fragment was present in each of the strains that now produced the type 2 capsule. Further, JD803 had also gained the 3.2 kb *HindIII* fragment present in WU2 (type 3), 2.1 kb of which is type 3-specific. This fragment was also present in JD871 and JD872 since it is contained in the plasmid insert (FIG. 5).  

The loss of type 3 genes by the strain converted to type 2 encapsulation indicated that capsule type change
does not occur by addition, but rather by replacement of the type-specific genes.

**EXAMPLE 8**

The DNA Sequence and Amino Acid Sequence of *cps3D*.

**A. Methods**

1. DNA Sequencing

Templates for sequencing were prepared from double-stranded plasmid DNA by denaturing with NaOH (2 M) for 5 min at room temperature, and precipitating with 5 M NH₄OAc and ethanol. DNA was sequenced by the Sanger dideoxy method using the Sequenase 2.0 kit (US Biochemicals, Cleveland, Ohio) and ³⁵S-dATP. The oligonucleotide primers 5′-GCCACTATCGACTACGG-3′ (SEQ ID NO:17) and 5′-TCATTGTATGCCTCCG-3′ (SEQ ID NO:18), corresponding to bases 308 to 325 and 445 to 428 of the cloning vectors pJY4163 and pJY4164 (Yother, et al., 1992), respectively, were used routinely. Primers 5′-GTGAGATAAAATAGTGGCG-3′ (SEQ ID NO:19) and 5′-TCCAGCTCGTGCTATAATCT-3′ (SEQ ID NO:20), corresponding to bases 3474 to 3493 and 3596 to 3577, respectively, of the type 3 capsule locus (FIG. 6G) were also used. All oligonucleotide primers were purchased from Oligos, etc. (Wilsonville, OR). DNA sequencing of PCR products was performed using the US Biochem PCR product sequencing kit, according to the directions of the manufacturer. PCR products were sequenced at least twice, from separate amplification reactions. Greater than 97% of the sequence was obtained for each strand.

2. Sequence analysis

The University of Wisconsin Genetics Computer Group programs (Genetics Computer Group, 1991) were used in the analysis of the DNA sequence. Database searches were
performed using the TFASTA program to detect homology with the deduced amino acid sequences. Potential sigma-70 type promoter sequences were located by using the FIND program to search for sequences with six or less mismatches, as compared to the consensus sigma-70 promoter sequence (Mulligan and McClure, 1986), and having a spacing of 15 to 20 bp between the -35 and -10 hexamers. Potential promoter sequences were evaluated using the equations of Harr et al. (Harr, et al., 1983). The sequence presented in FIG. 6D through FIG. 6J (SEQ ID NO4, SEQ ID NO:5 and SEQ ID NO:6) has been submitted to GenBank for assignment of an accession number.

3. Chromosome crawling and inverse PCR

To isolate the 5' end of cps3D and upstream DNA, S. pneumoniae WU2 chromosomal DNA was first digested with Eco1136 II (an isoschizomer of SacI that results in blunt ends) and separated on a 0.6% agarose gel. Fragments ranging from 6 to 7 kb were excised and purified using GeneClean (Bio101, La Jolla, CA). A 35 bp XbaI UniAmp adaptor (Clontech, Palo Alto, CA) was ligated to the purified fragments. The fragment of interest was then amplified by using a primer for the adaptor and a primer corresponding to the predicted active site sequence (nt 1802 to 1781 of FIG. 6E) of cps3D. A 1.8 kb and a 3.8 kb PCR product was obtained. The 1.8 kb PCR product extended from the active site to the SacI site upstream of cps3D (FIG. 6E). The 3.8 kb PCR product extended from the active site to the second SacI site further upstream (FIG. 7). The polymerase chain reaction (PCR) was performed using AmpliTaq DNA polymerase (Perkin-Elmer Corp., Norwalk, CT) in a Perkin Elmer model 480 thermocycler according to the directions of the manufacturer. In a similar manner, the 0.9 kb fragment from the cps3D active site to the EcoRV site upstream of
cps3D was amplified from a 3.5 kb EcoRV fragment from the WU2 chromosome.

To isolate the region 5' of the repeat sequence, a SacI-MscI fragment internal to the repeat region (extending from nucleotide 1 to 257 of SEQ ID NO:4) was first cloned into the insertion vector pSF151 (kanamycin resistant, KmR), and used to direct an insertion-duplication event into the type 3 S. Pneumoniae WU2 chromosome. Chromosomal DNA from the resulting KmR strain, JD1008, was digested with HindIII, self-ligated, and transformed into the E. coli. The resulting KmR plasmid, pRS111, contained in the pSF151 vector and DNA flanking the insertion, i.e., DNA extending from the HindIII site in cps3B to the HindIII site in the repeat sequence (~2.3 kb of S. pneumoniae DNA).

B. Results

The cps3D nucleotide sequence is shown in FIG. 6E (SEQ ID NO:5). The Cps3D amino acid sequence : (SEQ ID NO:11) is highly homologous (56% identity, 73% similarity) to that of the UDP-glucose dehydrogenase (HasB) from Streptococcus pyogenes (Dougherty and van de Rijn, 1993). Two other sequences were detected in the GenBank which shared a high degree of homology with Cps3D. These open reading frames from the Escherichia coli and Salmonella enteritica rfb clusters have not been shown biochemically or genetically to be UDP-glucose dehydrogenases (Bastin, et al., 1993), but they share a high degree of homology with HasB and Cps3D.

Cps3D (SEQ ID NO:11) has several characteristics consistent with it being UDP-glucose dehydrogenase. The N-terminal amino acid residues 2 to 29 have all the characteristics of an NAD-binding site (Wierenga, et al., 1986), and this sequence is very homologous to regions from HasB, AlgD (the GDP-mannose dehydrogenase of
Pseudomonas aeruginosa [Deretic, et al., 1987]), and the two potential UDP-glucose dehydrogenases from E. coli and S. enteritica. The homology with AlgD was previously noted by García et al., in the deduced amino acid sequence of the S. pneumoniae gene cap3-1 (García, et al., 1993). They suggested that Cap3-1 was the type 3 UDP-glucose dehydrogenase. Sequence ID NO:1 and SEQ ID NO:5 is in complete agreement with that of García et al., from the EcoRV site to the ScaI site (nucleotide 883 to 1377 FIG. 6D and FIG. 6E, containing amino acids 1 to 117, SEQ ID NO:11). However, no other homology was seen, suggesting that these investigators had cloned only the 5' end of the gene.

The Cps3D sequence at amino acid residues 251 to 263 (SEQ ID NO:11) is consistent with this being the active site of the enzyme. This region is identical at the amino acid level with that of HasB and the putative E. coli and S. enteritica UDP-glucose dehydrogenases. The homology of the active site region of HasB with that of bovine UDP-glucose dehydrogenase and AlgD has been fully described (Dougherty and van de Rijn, 1993). The cysteine at residue 259 (SEQ ID NO:11) of Cps3D contains the essential thiol group of the reactive site (Ridley, et al., 1975). The predicted size of Cps3D (45 kDa) is also similar to the size of the E. coli enzyme (47 kDa) (Schiller, et al., 1976).

**EXAMPLE 9**

**Identification of Capsule Mutants**

DNA sequencing and manipulations were performed as described in Example 8.

To determine the nature of the two cpsA mutations, identified in Example 1, the regions were amplified from the chromosomes of the mutant strains and sequenced.
Each mutant (JD611 and JD619) contained a single base pair transversion resulting in a premature stop codon in the \textit{cps3D} sequence. The locations of the mutations are indicated in FIG. 6E.

To localize the three \textit{cpsB} mutations, also identified in Example 1, located upstream of the UDP-glucose dehydrogenase mutations (\textit{cpsA}), standard PCR or chromosome crawling was used to amplify fragments from the parent type 3 chromosome that contained either the 5' end coding sequence of \textit{cps3D} (nucleotide 1027 to 1802, FIG. 6E), the promoter and the 5' end of \textit{cps3D} (nucleotide 885 to 1802, FIG. 6D and FIG. 6E), or the 5' end of \textit{cps3D} plus approximately 1 kb of upstream DNA (nucleotide 1 to 1802). Each of these fragments was used to transform the capsule-deficient mutants JD614 and JD692. JD692 could be transformed to encapsulation using the 5' end coding sequence of \textit{cps3D}, whereas JD614 was not restored to encapsulation by this fragment but was restored by the fragment containing the 5' end plus 141 bp of upstream DNA, including the promoter. Both of the mutants were restored by the 1.8 kb fragment containing the 5' end of \textit{cps3D} and the upstream DNA, and neither was restored with a fragment containing the 3' end of \textit{cps3D} (nucleotide 1759 to 2385, FIG. 6E). Thus, these upstream mutations are not located in a separate gene but are in either the \textit{cps3D} structural gene or its promoter. Since some capsule material is produced by these mutants, a mutation within the coding region (as in JD692) must be a missense mutation or an in-frame deletion or insertion which reduces the activity of the enzyme. The mutation in JD614 may be in the promoter, and thus, a promoter down mutation, or it may be in the structural gene but too close to the beginning of the gene for recombination and repair to occur with the fragment used.
Amplification and sequencing of the 250 bp PvuII-SspI fragment from the mutant strains A66R2 and Rx1 showed that each contained a missense mutation in the cps3D coding sequence (FIG. 6E).

EXAMPLE 10

DNA Sequences of cps3S and cps3U

DNA sequencing and analysis was performed as described in Example 8.

The region just downstream of cps3D contains a second gene, cps3S, that is required for type 3 capsular polysaccharide biosynthesis. An open-reading frame, 1248 bp in length, is transcribed in the same direction as cps3D and is in the same reading frame (SEQ ID NO:5). The direction of transcription is in agreement with that determined using cat insertions as described in Example 4. Only 15 bp separate a potential start codon for cps3S from the stop codon of cps3D. The sequence AGGGG just upstream of the putative start codon may serve as a ribosome binding site (FIG. 6E), or due to the close proximity of cps3D, no ribosome binding site may be necessary. The deduced amino acid sequence of Cps3S predicts a protein of 48 kDa (SEQ ID NO:12), if the first start codon at nucleotide 1 (nucleotide 2227 in FIG. 6F) is utilized. Other potential start codons are located at nucleotide 1 plus 19 and +61 (nucleotide 2245 and 2287 FIG. 6F, respectively), however neither of these are positioned near a ribosome binding site.

A short region of dyad symmetry was detected downstream of cps3S at nucleotide 3718 to 3738 (FIG. 6F). The scores for primary and secondary structure yielded by the TERMINATOR program of the GCG sequence analysis package (p=3.95, s=22) suggest that this region could function as a weak rho-independent terminator. However,
this sequence is 241 bases past the *cps3S* stop codon and closer to the start of the next open reading frame, i.e., *cps3U*. Therefore, the sequence may have more to do with the expression of *cps3U* than of *cps3S*. In fact, a potential promoter sequence for *cps3U* was detected upstream of the region of dyad symmetry, suggesting the potential structure could serve as an attenuator of *cps3U* expression. The *cpsU* open reading frame (SEQ ID NO:5), 918 bp in length, is transcribed in the same direction as *cps3D* and *cps3S*, and is predicted to encode a protein of 34 kDa (SEQ ID NO:13).

**EXAMPLE 11**

*Cps3S is Homologous to Polysaccharide Synthases*

A search of the GenBank revealed that the predicted Cps3S protein (SEQ ID NO:12) is homologous to polysaccharide synthases. The greatest degree of homology was found with HasA, the hyaluronic acid synthase from *S. pyogenes* (23% identity, 50% similarity) (DeAngelis, et al., 1993b; Dougherty and van de Rijn, 1994). Hyaluronic acid consists of alternating N-acetyl glucosamine and glucuronic acid residues. Hyaluronic acid and the pneumococcal type 3 capsule are similar in structure in that both are composed of β(1-4) linked repeating disaccharide units containing glucuronic acid. Like pneumococcal type 3 capsule, hyaluronic acid capsule contains both β(1-3) and β(1-4) linkages, however the linkage to glucuronic acid is β(1-4) in hyaluronic acid but β(1-3) in type 3 capsule (Reeves and Goebel, 1941). Homology was also seen between Cps3S and NodC from *Rhizobium meliloti* (21% identity, 47% similarity). NodC is necessary for the synthesis of nodulation factor, a substituted oligosaccharide consisting of β(1-4) linked N-acetyl glucosamine residues (Lerouge, et al., 1990). It has previously been noted that HasA and NodC are homologous to polysaccharide synthases, including FBF15
of Stigmatella aurantiaca, pDG42 of Xenopus laevis, and chitin synthases from both *Saccharomyces cerevisiae* and *Candida albicans* (DeAngelis, et al., 1993b; Dougherty and van de Rijn, 1994; Atkinson and Long, 1992; Debellé, et al., 1992). Cps3S is also homologous to these proteins. These results suggest that Cps3S is the type 3 capsular polysaccharide synthase.

The PILEUP program was used to align the amino acid sequences of the bacterial polysaccharide synthases Cps3S (SEQ ID NO:12), HasA, NodC, and FBP15. Only a few clusters of amino acids are found to be conserved in all four proteins. A few of these, GKR (residues 131 to 133 SEQ ID NO:12), an acidic region VDSD (153 to 156 SEQ ID NO:12), DRXLT (256 to 260 SEQ ID NO:12), QQXRW (292 to 296 SEQ ID NO:12), and WXTR (418 to 421 SEQ ID NO:12), are also found in the eukaryotic polysaccharide synthases.

Since all four proteins contain highly hydrophobic stretches, hydrophobic amino acids are found conserved at several locations throughout the proteins. Four hydrophobic stretches identified in Cps3S are found in all four proteins. These regions may span the cell membrane. This hypothesis has been supported for NodC. Immunogold labeling revealed a surface location for NodC, and the C-terminal hydrophobic region was shown to direct the insertion of an alkaline phosphatase fusion protein to the cell membrane (Johnson, et al., 1989; John, et al., 1988). Earlier studies indicated that the type 3 capsule synthesizing activity also has a membrane location (Smith, et al., 1961). The last hydrophobic stretch may be required for the function of Cps3S since the insertion in JD897 which eliminated this region (the last 45 amino acids of the protein) resulted in loss of capsule production (FIG. 6F). Expression of Cps3S in *E. coli* was, like that of NodC, lethal to the host.
A. Method

1. Expression of Cps3S

A 2.1 kb Sau3AI-PstI fragment containing the 3' end of cps3D and the entire cps3S gene was cloned from pJD351 into the expression vector pKK223-3 (Brosius and Holy, 1984) at the polylinker BamHI-PstI sites to yield pJD424. Cultures of E. coli TG-1 (Sambrook, et al., 1989) or TG-1 transformants were grown to exponential phase, at which time isopropyl-b-D-thiogalactoside (IPTG) was added to a concentration of 1 mM to induce expression from the tac promoter of pKK223-3.

Transformations and other DNA manipulations were performed as described in Example 1.

B. Results

The sequence from residues 211 to 233 in NodC was noted for the large number of cysteine residues. It has been suggested that this region participates in the binding of divalent cations which are necessary for the production of chitin and chitin-like molecules (Atkinson and Long, 1992). Type 3 capsule synthesis requires Mg++ (Smith, et al., 1960). Although this region in Cps3S contains only one cysteine, the region is highly conserved between all four proteins.

The GenBank search also revealed that Cps3S has homology over short stretches to the rhamnosyl transferase RfbN from Salmonella enteritica, which is necessary for the production of O-antigen in type B strains. This enzyme creates an α(1-4) linkage to mannose in the O-antigen repeat unit. The homologous regions are a subset of those conserved regions common to HasA, NodC, and Cps3S, but the best homology is seen in the region 229 to 278 (SEQ ID NO:12).
In the production of Group B type III capsular polysaccharide, the galactosyl transferase CpsD transfers a galactose to a molecule located in the cell membrane. Rubens et al. (1993), suggested that the acceptor may be dolichol or a related molecule, and identified a region of CpsD with homology to putative dolichol binding regions of several proteins. Although it is not clear that such sequences are actually involved in dolichol recognition or binding (Schutzbach, et al., 1993), several similar regions (e.g., at residues 7 to 20, 21 to 38, and 388 to 401, as numbered in SEQ ID NO:12) are present in CpsS. Since the putative dolichol binding motif [FL(F/I)VXFXI(P/L)PFXY] (Albright, et al., 1989; Kelleher, et al., 1992) is a highly hydrophobic sequence that is rich in phenylalanines, the sites in Cps3S may actually reflect the hydrophobicity of the molecule and the A-T rich bias in the DNA sequence rather than indicating a specificity for dolichol-like molecules. It is not known whether S. pneumoniae utilizes an intermediate acceptor in capsule synthesis, however the capsular polysaccharides of several serotypes have been found to be covalently linked to the cell wall. Type 3 capsule, by contrast, is not covalently linked to the cell and is generally considered an exopolysaccharide (Sorensen, et al., 1990). Therefore, if Cps3S does use a membrane bound acceptor, it is likely not the final acceptor.

EXAMPLE 12

Cps3U is Homologous to Glucose-1-Phosphate Uridylyltransferases and Cps3M is Homologous to Phosphomutases

The gene downstream of cps3S is designated as cps3U (SEQ ID NO:5) based on its probable function. The amino acid sequence of Cps3U (SEQ ID NO:13) showed a high degree of homology with glucose-1-phosphate
uridylyltransferases from several other bacterial species. The highest degree of homology was found with GtaB from Bacillus subtilis (55% identity, 73% similarity). The active site of glucose-1-phosphate uridylyltransferase has not been characterized from any of the bacterial enzymes, however, the active site in the enzyme from potato tuber (Solanum tuberosum) has been investigated. Kazuta et al., recognized 5 lysine residues present at the active site (Kazuta, et al., 1991), and by mutational studies Katsube et al., showed that one of these residues was important for function, and a second was absolutely required (Katsube, et al., 1991). Cps3U contains 24 lysines, six of which are absolutely conserved among the six bacterial glucose-1-phosphate uridylyltransferases in the database. Only one region from Cps3U containing a conserved lysine can be aligned well with the potato tuber enzyme sequence. It is homologous to the region containing the required lysine.

The final gene in SEQ ID NO:5, is cpsM with a deduced amino acid sequence (SEQ ID NO:14). The CpsM amino acid sequence revealed significant homology to both phosphoglucotransferases (PGM) and phosphomannomutases (PMM) from a diverse group of microorganisms. Contained with CpsM is a phosphoserine signature sequence (GIMVTASHTPAPFG) conserved within the reported active sites of both PGMs and PMMs. However, approximately 15% of the C terminus present in other phosphomutases, and apparently more important for their function, is absent from CpsM. Phosphomutase activity from a recombinant CpsM was not detected in E. coli, suggesting that cpsM may encode a non-functional protein.
EXAMPLE 13

cps3S and cps3D are Transcribed as an Operon

A. Methods

Southern blotting was performed as described in Example 4, all other DNA manipulations, including insertion deletion mutations, were performed as described in Example 1. The locations of mutations can be seen in FIG. 6E and FIG. 6F.

B. Results

Use of fragments subcloned from the cps3DSU region to direct insertion-duplication mutations in the parent type 3 chromosome resulted in several mutants that produced no detectable capsule (FIG. 9) and exhibited the extremely rough phenotype described by Taylor (1949). The colonies were very small and rough, and the cells clumped when grown in liquid culture. DNA sequencing revealed that the sites of the mutations are within cps3S (FIG. 6F). The lack of capsule production in these mutants must be due to loss of cps3S expression, rather than to a polar effect on downstream genes, since insertions within cps3U or cps3M, the next genes downstream, had no apparent effect on capsule production.

Molecular and genetic evidence suggest cps3S is in an operon with cps3D. Sequence analysis revealed no potential promoter sequences in the region upstream of cps3S (FIG. 6E, SEQ ID NO:5). The phenotypes of several insertion mutants also suggest that no promoter is located in the 3' end of cps3D and that cps3S is transcribed from the cps3D promoter. The sites of these insertions are shown in FIG. 6E, FIG. 6F and FIG. 7, however, the structures of the mutations are more fully illustrated in FIG. 9. To insure that the plasmids had inserted as expected for insertion-duplication mutations,
chromosomal DNA from the mutant strains was subjected to Southern blot analysis.

Insertion mutants were digested with MscI/FspI for JD982, MscI/SalI for JD983, and MscI/KpnI for JD908, JD902, and JD900 and run on agarose gels and blotted as described in Example 4. The blots were probed with vector pJY4164. Increasing distance from the MscI site to the end of the vector was demonstrated by an increase in the size of the upper band. A faint band in the JD982 lane was observed, likely a result of partial digestion. The 4.7 kb and 4.8 kb bands in JD982 and JD908, respectively, indicate that these mutants contain a duplication of the inserted plasmid.

Insertion of the plasmids results in a duplication of the cloned fragment. Therefore, mutant strains such as JD908, in which the duplicated fragment contains both the 5' end of cps3S and the 3' end of cps3D, have a full-length copy of cps3S downstream of the plasmid insertion. In addition, the full-length copy is contiguous with the 3' end of cps3D. Therefore, if cps3S had its own promoter, or if one were located in the 3' end of cps3D, these insertions should not result in a loss of cps3S expression. However, four such insertions have been made in the WU2 chromosome (JD846, JD897, JD898, and JD908), and even with a duplication of up to 450 bp of the 3' end of cps3D, a loss of capsule production was observed.

Two more internal insertions in cps3D were created. As expected these insertions eliminated capsule production (FIG. 9). However, since cps3D and cps3S are transcribed as an operon, this result does not prove that cps3D is required for capsule synthesis. That fact is demonstrated by the lack of capsule production seen in strains containing non-polar point mutations in cps3D (Example 14).
EXAMPLE 14

In Vitro Polymerization Assay

To evaluate the competence of the mutants to synthesize type 3 capsule, an in vitro polymerization assay was used.

A. Methods

1. In vitro polysaccharide synthesis

Type 3 capsular polysaccharide was synthesized and quantitated in vitro using a modification of the method of Smith, et al., 1961. Crude extracts containing cell membranes and cytoplasm were prepared from 200 ml of S. pneumoniae cultures harvested at an O.D. of 0.25 as described (Yother and White, 1994), except that cell material was concentrated 200-fold, and all steps were performed using a thioglycolate buffer (10 mM sodium thioglycolate, 5 mM MgSO₄, 100 mM Tris-HCl pH 8.3) to stabilize the enzymes (Smith, et al., 1960). The digestion of cell wall material by mutanolysin treatment was performed in this buffer and 20% sucrose. Protoplasts were sonicated three times for 15 s at 35% power at 0°C.

Polysaccharide synthesis was carried out at 34°C for 2 h in a 1 ml reaction containing 100 ml of extract, 5 mM UDP-glucose, 5 mM UDP-glucuronic acid (where indicated), and 1 mM NAD, in the thioglycolate buffer. The reaction was boiled 1 min then quickly cooled to 25°C in H₂O. Following centrifugation for 30 s at 8160 x g, the type 3 specific monoclonal antibody 16.3 (Briles, et al., 1981a) was added in excess to the supernatant and incubation was continued at 37°C for 30 min.

The specific antigen-antibody complexes were measured at 650 nm in a spectrophotometer, and the amount
of capsule was determined by comparison with a standard curve prepared using purified type 3 polysaccharide purchased from ATCC (Rockville, MD) (Bernheimer, 1953). Reactions were done in triplicate and were standardized to protein content of the crude extract, as determined in duplicate using the Bio-Rad Laboratories (Hercules, CA) protein assay kit.

B. Results

The spontaneous mutants JD611 and JD619 (cpsA1 and cpsA2), which contain stop mutations in cps3D, produce no detectable capsular material. However, both synthesized high molecular weight type 3 polysaccharide in a cell-free system in vitro when provided with the nucleotide sugar precursors, i.e., UDP-glucose and UDP-glucuronic acid (Table 7). No capsule was produced by these mutants when UDP-glucuronic acid was omitted from the reaction. These results indicate that these mutants produce no capsule due to the lack of UDP-glucuronic acid and support the conclusion that Cps3D is the UDP-glucose dehydrogenase. They also confirm that stop mutations in cps3D are not polar on cps3S. The increased amount of polysaccharide produced by the WU2 extract (as compared to that produced by that of JD611 or JD619) may be explained by the observation of Smith et al. (1961), that increased amounts of type 3 capsule are produced in vitro when a small amount of unpurified polysaccharide is already present in the reaction.

The mutants which contain insertions within cps3S (JD902), or between the full-length copies of cps3D and cps3S (JD908, JD897) were unable to synthesize significant amounts of capsule even with both precursors present. These results emphasize the role of Cps3S in capsule synthesis and support the conclusion that cps3D and cps3S are transcribed as an operon.
The capsule-deficient mutants JD614 and JD692 synthesized only small amounts of additional polysaccharide in the in vitro assay. This result is somewhat surprising since JD692, which was shown to contain a missense mutation within the cps3D coding region, should still make a functional Cps3S (i.e., the cps3D mutation must not be polar since the intact cells are able to synthesize some polysaccharide). The result may suggest that the defective UDP-glucose dehydrogenase in some way interferes with the ability to synthesize the normal polysaccharide. Alternatively, the stability of the cpsDS transcript may be altered by the mutation, resulting in a reduced amount of CpsS.
Table 7. *In vitro* capsule synthesis assay.

<table>
<thead>
<tr>
<th>strain</th>
<th>Cps phenotype</th>
<th>UDPGAb</th>
<th>CPS (μg/mg protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JD611</td>
<td>Cps3D⁻S⁺</td>
<td>+</td>
<td>9.8 ±0.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-</td>
<td>0.9 ±0.2</td>
</tr>
<tr>
<td>JD619</td>
<td>Cps3D⁻S⁺</td>
<td>+</td>
<td>5.7 ±0.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-</td>
<td>0.2 ±0.1</td>
</tr>
<tr>
<td>JD614</td>
<td>Cps3D⁺S⁺</td>
<td>NA⁺</td>
<td>5.4 ±0.4 (t₀)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>5.9 ±0.5 (0.5)</td>
</tr>
<tr>
<td>JD692</td>
<td>Cps3D⁺S⁺</td>
<td>NA</td>
<td>4.8 ±0.3 (t₀)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+</td>
<td>7.0 ±1.0 (2.2)</td>
</tr>
<tr>
<td>JD902</td>
<td>Cps3D⁺S⁻</td>
<td>+</td>
<td>1.7 ±0.3</td>
</tr>
<tr>
<td>JD908</td>
<td>Cps3D⁺S⁻</td>
<td>+</td>
<td>1.5 ±0.1</td>
</tr>
<tr>
<td>JD897</td>
<td>Cps3D⁺S⁻</td>
<td>+</td>
<td>1.1 ±0.1</td>
</tr>
<tr>
<td>WU2</td>
<td>Cps3D⁺S⁺</td>
<td>NA</td>
<td>3.8 ±0.2 (t₀)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>+</td>
<td>16.6 ±0.3 (12.8)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>-</td>
<td>16.3 ±0.8</td>
</tr>
<tr>
<td>D39</td>
<td>Cps2⁺</td>
<td>+</td>
<td>0.5 ±0.3</td>
</tr>
</tbody>
</table>

*a* Capsule phenotypes are based on the *cps3D* and *cps3S* genotypes. - indicates either a stop or insertion mutation (see FIG. 6E, FIG. 6F and FIG. 9 for locations of mutations). * indicates either a missense or in-frame deletion or insertion in *cps3D* that apparently also affects *cps3S*.

*b* NA, not applicable.

*c* For strains which produce capsule in vivo, the amount of polysaccharide present at the start of the assay (t₀) is given, and the amount of polysaccharide produced during the assay is indicated in parentheses.
EXAMPLE 15

Biochemical Pathway

Based on the genetic analysis, the homology of the amino acid sequences of the type-specific genes to the sequences of enzymes of known function, the behavior of the mutants in biochemical and immunochemical assays, and previous biochemical characterizations of type 3 strains (Austrian, et al., 1959; Dillard and Yother, 1994; Smith, et al., 1960; Smith, et al., 1961; Bernheimer, 1953), a pathway for the biosynthesis of type 3 capsular polysaccharide is proposed (FIG. 10). The last of the type-specific genes, \( \text{cps3M} \), is homologous with phosphoglucomutases from several bacterial species. Even though maintained in the type 3-specific region, \( \text{Cps3U} \) and \( \text{Cps3M} \) may not be required for capsule synthesis, since an insertion internal to \( \text{cps3U} \) (which has a polar effect on \( \text{cps3M} \)) does not result in loss of capsule production (FIG. 7 and FIG. 9), as judged by colony morphology on blood agar medium.

EXAMPLE 16

The Downstream Non Type-specific Flanking Region and Mapping Other Capsule Types

Southern blots of digested chromosomal DNA from strains 2, 3 and 6B and probed with pJD377 was performed as described in Example 4. Faint bands in addition to the band of interest was observed on Southern blots. This was probably due to the detection of fragments containing the amiA-like genes which have homology to \( \text{plpA} \). DNA was either digested with \( \text{BglII} \), \( \text{SacI} \) of \( \text{HindIII} \). Other laboratory techniques were as described in Example 1 or Example 8.

Sequence analysis of the 1.2 Kb \( \text{SacI-HindIII} \) fragment (from plasmid JD377) employed in Example 5
contained the 3' end of cps3M and the 5' half of a gene with 50% identity to the S. pneumoniae amiA (SEQ ID NO:6). The amiA-like sequence has recently also been identified by Pearce et al. and named exp1 (Pearce, et al., 1993), and subsequently renamed plpA (Pearce, et al., 1994). Further Southern hybridizations performed as described in Example 4 showed that the non-type-specific homologous DNA in the 1.2 kb SacI-HindIII fragment is plpA.

A partial copy of a transposase gene was also identified immediately adjacent to and between cpsM and plpA. Previous findings of repetitive elements linked to the capsule locus suggest that the deletions in this region may be the result of a transposition event, possibly one which introduced the type 3-specific cassette.

If, as in type 3, the homologous region is directly adjacent to the type-specific genes in other serotypes, it should be possible to map other type-specific genes using this fragment. This was found to be the case, and the chromosome maps of the capsule regions in strains of types 2, 3, and 6B, from Southern blots, are shown in FIG. 11. It can be seen in FIG. 11 that restriction sites located to the right of the plpA fragment are highly conserved in all three strains. The type 3 strain differs slightly in this region due to a deletion of the 5' end of plpA. The sites located to the left of plpA are divergent among the capsule types. The close linkage of the region to all the necessary type-specific genes for each type, combined with the different restriction maps and the fact that the type 3-specific genes are located directly adjacent to this fragment, suggests that this region contains the type-specific genes in all three capsule types.
EXAMPLE 17

The Upstream Non Type-specific Flanking Region

In order to isolate DNA 5’ of the biosynthetic genes, a 1.8 kb fragment extending from the upstream SacI sites to just before the PvuII site in cps3D (nucleotide 1 through 1802 of FIG. 6D and FIG. 6E, nucleotide 1 through 934 of SEQ ID NO:4 and nucleotide 1 through 868 of SEQ ID NO:5) was amplified from the type 3 WU2 chromosome using inverse PCR as described in Example 8. All other materials and methods were as described in Example 1, 4 and 8.

The 1.8 kb fragment was then used to probe HindIII-digested chromosomal DNA from seven S. pneumoniae serotypes (2, 3, 5, 6, 8, 9 and 22). The fragment hybridized strongly with the expected fragments at 2.2 and 2.3 kb in the type 3 strain. However, hybridization was also observed with fragments of 2.6 and 8 kb, along with weak hybridization with several other fragments (3.0, 3.1, and 4.4 kb). Likewise, each of the strains representing other capsule types contained two strongly homologous fragments (4.8 and 8.0 kb for types 2, 6B, 8, 9; 2.2 and 4.8 or 12 for types 5 and 22, respectively) and at least one weakly homologous fragment (4.4 kb). When chromosomal DNAs of types 2, 3, and 6B were digested with PstI, PvuII, or SacI/HindIII, and probed with the 604 bp SacI-HindIII fragment (pJD392) upstream of cps3D (within nucleotides 1 through 610; SEQ ID NO:4, FIG. 6D), 4 to 10 bands were detected in each.

Transformation studies were performed to examine linkage of the repeat upstream region to the type-specific capsule genes. The plasmid (pJD392) containing the 604 bp SacI-HindIII fragment was introduced into the chromosome of the type 3 strain. The insert, located in the 2.2 kb HindIII fragment (FIG. 7) adjacent to the type
3-specific genes, did not affect capsule production. When the resulting strain was used to transform recipients of types 2 and 6B, greater than 95% of the erythromycin-resistant transformants expressed type 3 capsule. However, when pJD392 was transformed into strains of types 2 and 6B, the plasmid inserted into the 8 kb HindIII fragment, and the type specific genes could not be moved to strains of heterologous types (i.e., 2, 3, or 6B) by transformation and selection for linkage to the erythromycin marker in the insertions. This suggests that this region may be found in more than one location and not necessarily located, adjacent to the cps locus in all other serotypes.

To isolate DNA 5' of the SacI site in the repeat region (SEQ ID NO:4), a 257 nucleotide SacI-MscI fragment internal to the repeat region was used to direct an insertion-duplication event in the type 3 S. pneumoniae WU2 chromosome. The flanking 2.3 kb HindIII fragment was then closed from the chromosome, as described in Example 8. The 1.4 kb SacI fragment from within this HindIII fragment was then used to probe HindIII-digested chromosomal DNA from three S. pneumoniae serotypes (2, 3, and 6B). The fragment hybridized strongly with the expected 2.3 kb band in the type 3 strain. Less intense bands were also observed at 0.8, 3.0 and 8.0 kb. The type 2 and 6B strains each contained a strongly homologous band at 0.8 kb and a strong but less intense band at 9.0 kb. Digestion of types 2, 3, and 6B with other enzymes followed by hybridization with the SacI fragment yielded the results shown in Table 8. The results show that this region is found in only one location in all serotypes examined. The weak bands observed for serotype 3 probably represent hybridization of the 5' end of the repeat region 132 nucleotides upstream of the SacI site (FIG. 7).
TABLE 8. BANDS DETECTED BY SOUTHERN BLOTTING

<table>
<thead>
<tr>
<th>Restriction Enzyme used to digest chromosomal DNA.</th>
<th>BglIII</th>
<th>SacI</th>
<th>SphI</th>
</tr>
</thead>
<tbody>
<tr>
<td>type 2</td>
<td>&gt;12*</td>
<td>4.0,&gt;12</td>
<td>12,&gt;12</td>
</tr>
<tr>
<td>type 3</td>
<td>&gt;12*</td>
<td>1.4, (weak at 3.5,8.5,&gt;12)</td>
<td>10 (weak at 12.5,13)</td>
</tr>
<tr>
<td>type 6B</td>
<td>&gt;12*</td>
<td>4.0,12</td>
<td>12,&gt;12</td>
</tr>
</tbody>
</table>

* The BglIII fragments were not identical in size.

Numbers represent size of bands in kb as observed on southern blots carried out as described in Example 4.

DNA in the 1.4 kb SacI fragment was sequenced using techniques as described in previous Examples, and can be seen, alongside the predicted amino acid sequences, in FIG. 6A, FIG. 6B and FIG. 6C (SEQ ID NO:1, SEQ ID NO:2, SEQ ID NO:3, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9 and SEQ ID NO:10).

EXAMPLE 18

Capsule Type Expression and Virulence in *S. pneumoniae*

In these studies, isogenic strains expressing the type 3 capsule were constructed and the effect on virulence was determined. Strains of types 2, 5, and 6B were used as recipients. The type 2 and 5 strains differ in virulence from the type 3 strain in terms of time required to cause death (shorter with type 2) and LD\textsubscript{50} (lower with type 5). The type 6B strain is of low virulence in mice. The results showed that expression of the type 3 capsule attenuated the virulence of the type 5 strain, caused the type 6B strain to become highly virulent, and had no effect on the type 2 strain. Thus, in general, the expression of virulence was correlated with the type of capsule expressed.
A. Methods

1. Transformations, serotyping, ELISAs and restriction enzyme fragment patterns.

Transformations ELISAs and DNA manipulations were performed as described previously in Example 1. All transformants and parental strains were serotyped with capsule type-specific antisera (Statens Seruminstitut, Copenhagen, Denmark) in slide agglutination assays. Genomic DNA, was digested with HindIII for 4 h at 37°C and electrophoresed overnight through 0.7% agarose in Tris-borate-EDTA buffer.

2. Analysis of PspA.

Bacteria were grown in CDM containing 2% choline, a condition that causes release of PspA into the culture medium. Filtered, unconcentrated supernatant fluids (20 μl) were electrophoresed in sodium dodecyl sulfate (SDS)-12% polyacrylamide gels. Western blotting (immunoblotting) was performed by using a semidry electroblotter (Bio-Rad Laboratories, Richmond, Calif.), and the blots were processed as described previously (Yother et al., 1992). The PspA-specific monoclonal antibodies XiR278, Xi126, and 2A4 were kindly provided by Larry McDaniel (University of Alabama at Birmingham). Silver staining was performed by using the Silver Stain Kit from Stratagene Cloning Systems, Inc. (La Jolla, Calif.).

3. Characterization of morphology and capsule production.

For average chain length determinations, bacteria were grown in THY to an optical density at 600 nm (OD_{600}) of ~0.3. Chain lengths were determined microscopically by using a Petroff-Hauser counting chamber (Auther C. Thomas Co., Philadelphia, PA). An average of five squares was counted for each strain. Comparisons of average chain lengths were determined by using the two-sample rank test (Zar, 1984).
The number of cells per colony was determined by using bacteria grown on blood agar medium for 18 h at 37°C in 5% CO₂. A plug containing a single colony was obtained with a sterile Pasteur pipette and then resuspended in 50 μl of THY. Tenfold serial dilutions were performed in THY and plated on blood agar medium. Plates were incubated overnight at 37°C in 5% CO₂, and the number of CFU per colony was calculated.

Buoyant density determinations were performed by using bacteria grown on blood agar medium or in THY. Bacteria grown on solid medium were harvested by washing each plate with water, centrifuging the suspension, and then resuspending the pellet to an OD₆₀₀ of ~0.4 with water. Ten-milliliter liquid cultures, grown to an OD₆₀₀ of ~0.5, were harvested by centrifugation for 10 to 15 min at 8,000 to 16,000 × g. Bacteria were washed twice with water prior to being loaded onto 10-ml, continuous, 0 to 50% Percoll (Pharmacia, Piscataway, N.J.) gradients. As standards, 5 μl of density marker beads (Pharmacia) ranging in size from 1.033 to 1.076 g/ml, were also loaded. Gradients were centrifuged for 30 min at 8,000 × g with the brake off. A standard curve based on the migration of the marker beads was generated, and the density of the bacteria was determined by extrapolation.

For determinations of total capsule content, 1.5-ml cultures grown in CDM containing 0.0005% choline were harvested by centrifugation at 8,000 to 16,000 × g for 10 min. Culture supernatant fluids were filtered and saved, and the cells were resuspended in 500 μl of protoplast buffer (20% sucrose, 0.005 M Tris [pH 7.4], 0.0025 M MgSO₄). Cell sonicates were produced by three cycles of a 10-s pulse, followed by a 10-s incubation on ice, with a Fisher Sonic Dismembrator model 300 (Fisher Biotechnology, St. Louis, Mo.) with the intensity control
set at 30. Culture supernatant fluids and cell sonicates were stored at ~20°C.

For surface localization assays, 1.5-ml cultures grown to an OD$_{600}$ of ~0.5 were heat killed by incubation at 65°C for 30 min. Bacteria were harvested by centrifugation, and culture supernatant fluids were filtered and saved. After the pellets were washed twice with phosphate-buffered saline (PBS; 137 mM NaCl, 2.7 mM KCl, 4.3 mM Na$_2$HPO$_4$·7H$_2$O, 1.4 mM KH$_2$PO$_4$), the pellets were resuspended in 1.5 ml of THY. Samples were stored at 4°C.

4. Virulence assays.

The virulence of the type 3 derivatives was compared with that of the parental strains in BALB/ByJ female mice (Jackson Laboratory, Bar Harbor, Maine). Bacteria were grown to the mid-log phase in THY. Samples were diluted serially in sterile lactated Ringer’s solution, and 0.2 ml was used to infect mice intraperitoneally (i.p.) or intravenously (i.v.), as indicated. Fifty percent lethal doses (LD$_{50}$s) were determined by the method of Reed and Muench (1938) and compared by Fisher’s exact test (Zar 1984). Median times to death were analyzed by using the two-sample rank test (Zar 1984). The $P$ values were determined by using a two-tailed table.

B. Results

As described in Table 3, strain JD770 contains a non-destructive insertion in the type 3 capsule locus. The amount and cellular localization of the capsular material produced by JD770 is identical to that of its parent strain WU2. Transformation of JD770 chromosomal DNA, and selection for erythromycin resistance, results in isolates that express the type 3 capsule of the donor but not the capsule of the recipient strain (Example 6). Based on this result, JD770 DNA was used to transform
type 2, 5, and 6B recipients and selected for erythromycin-resistant isolates (see Table 9). All of the type 2 Ery^T transformants expressed the type 3 capsule but not the type 2 capsule. >95% of the type 5 and type 6B Ery^T transformants expressed the type 3 capsule but not the capsule of the recipient parent. The remainder of the type 5 and 6B transformants expressed the capsular type of the recipient parent only.
TABLE 9. BACTERIAL STRAINS

<table>
<thead>
<tr>
<th>Strain</th>
<th>Derivation(^a)</th>
<th>Relevant phenotype</th>
<th>Chain length(^b)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>WU2</td>
<td>Clinical isolate</td>
<td>Encapsulated type 3</td>
<td>2.3±0.14</td>
<td>Briles et al. 1981b</td>
</tr>
<tr>
<td>JD770</td>
<td>pJD330(^c) x WU2</td>
<td>Encapsulated type 3; Em(^r) 6.6±0.76</td>
<td>2.4±0.19</td>
<td>This study</td>
</tr>
<tr>
<td>D39</td>
<td>Clinical isolate</td>
<td>Encapsulated type 2</td>
<td>6.2±0.69</td>
<td>Avery et al. 1944</td>
</tr>
<tr>
<td>JD803 (2/3) (^d)</td>
<td>JD770 x D39</td>
<td>Encapsulated type 3; Em(^r) 5.2±0.74</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>JD804 (2/3)</td>
<td>JD770 x D39</td>
<td>Encapsulated type 3; Em(^r) 9.2±0.80</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>DBL5</td>
<td>Clinical isolate</td>
<td>Encapsulated type 5</td>
<td>9.2±0.80</td>
<td>Yother et al. 1982</td>
</tr>
<tr>
<td>TK5010 (5/3)</td>
<td>JD770 x DBL5</td>
<td>Encapsulated type 3; Em(^r) 2.1±0.08</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>TK5011 (5/3)</td>
<td>JD770 x DBL5</td>
<td>Encapsulated type 3; Em(^r) 2.4±0.13</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>TK5012 (5/3)</td>
<td>JD770 x DBL5</td>
<td>Encapsulated type 3; Em(^r) ND(^f)</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>DBL1</td>
<td>Clinical isolate</td>
<td>Encapsulated type 6B</td>
<td>2.7±11</td>
<td>Briles et al. 1992</td>
</tr>
<tr>
<td>TK3026 (6B/3)</td>
<td>JD770 x DBL1</td>
<td>Encapsulated type 3; Em(^r) 3.1±30</td>
<td>This study</td>
<td></td>
</tr>
<tr>
<td>TK3028 (6B/3)</td>
<td>JD770 x DBL1</td>
<td>Encapsulated type 3; Em(^r) 2.4±15</td>
<td>This study</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Crosses were done by transformation. Each derivative is the result of at least three backcrosses with the parental recipient strain.

\(^b\) The numbers of cells per chain (means ± standard errors) were determined during this study by microscopic examination.
pJD330 contains a *Sau3AI* fragment that is part of the type 3 capsule locus. It was constructed in pJY4163 (Yother et al., 1992) which contains an *Em*<sup>r</sup> marker. Insertion of pJD330 into the *S. pneumoniae* chromosome places the *Em*<sup>r</sup> marker in the type 3 locus but does not affect capsule expression.

For clarity, strains will be listed by strain number, genetic background, and capsule type expressed. For example, JD803 (2/3) represents a strain with a type 2 genetic background expressing a type 3 capsule.

Significantly different from recipient parent strain (*P* < 0.001).

ND, not determined.
To produce essentially isogenic strains, two independent transformants from each cross were backcrossed at least three times to their respective parent recipient strains. The final isolates were examined for restriction enzyme fragment patterns, pneumococcal surface protein A (PspA) expression, capsule expression, and morphological characteristics prior to testing in a mouse virulence model.

1. Restriction enzyme fragment patterns.

The HindIII restriction patterns of the strains used in these studies can be easily distinguished. In all cases, the type 3 derivatives constructed here were found to have the HindIII pattern of the recipient strain, indicating that gross alterations in the genomic DNA content had not occurred and that the parent donor strain JD770 had not been inadvertently re-isolated.

2. PspA expression.

PspA varies with respect to molecular weight, antigenic determinants, and strain distribution. PspA serotypes and capsular serotypes do not correlate. The strains used in these studies expressed PspAs that had different molecular weights and reacted with different PspA-specific monoclonal antibodies. In all cases, the PspAs of the type 3 derivatives constructed here were found to have the molecular weight and antibody reactivities of the parent recipient strains.

3. Morphologic characterization and capsule production.

Microscopic examination revealed that alteration of capsular type had no effect on the chain length of the type 2 and type 6B derivative strains. However, the chain lengths of the type 5 derivatives differed significantly from that of the type 5 parental strain and
were almost identical to that of the type 3 parent (Table 9).

Morphologically, type 3 strains exhibit large mucoid capsules when grown on blood agar plates, whereas type 2, 5, and 6B strains have small mucoid capsules. The type 3 derivatives of the type 2, 5, and 6B strains had a similar appearance to the type 3 parent on blood agar plates. The increase in colony size compared with that of the recipient parents did not appear to be due to cell number since similar numbers of cells per colony were observed for all of the parent and derivative type 3 strains (data not shown). To examine capsule production, Percoll density gradients and ELISAs were performed. Percoll density gradient centrifugation has been shown previously to differentiate capsular serotypes and amounts by density (Briles et al., 1992). In this assay, all of the derivatives had densities similar to that of the parent type 3 strain and distinct from that of the recipient parent strains (FIG. 12). Thus, all of the derivatives produced cell-associated, surface-localized type 3 capsule in amounts similar to that of the type 3 parent. The total amounts of capsule material produced, i.e., both cell associated and released, were determined in ELISAs to be similar for both the type 3 parent and each of the derivatives (FIG. 18). ELISAs were also used to confirm that the amounts of surface-accessible capsule were similar in the type 3 parent and the derivatives.


To assess the effect of alteration of capsule type on virulence, BALB/ByJ female mice were infected i.p. or i.v. with the type 3 derivatives and parent strains. Strain JD770, which contains the nondestructive erythromycin resistance marker in the type 3 capsule locus, did not differ from its parent type 3 strain WU2
in terms of median time to death (52 versus 49.5 h, i.p.) or LD$_{50}$ (75 versus 50 CFU, i.p.; 1 x 10$^5$ versus 2 x 10$^5$ CFU, i.v.). Thus JD770 was used in subsequent studies for comparisons with the type 3 derivatives. As expected, the recipient parent strains were significantly different from JD770 with respect to time to death or LD$_{50}$s (FIG. 14). Expression of the type 3 capsule had no apparent effect on the virulence of the type 2 recipient strain; i.e., the time required to cause death was not significantly different from that of the type 2 parent but was significantly different from that of the type 3 parent (FIG. 14A). However, alteration of capsular type had dramatic effects on the virulence of the type 5 and 6B strains. In contrast to the highly virulent type 5 parental strain (LD$_{50}$, ~10 CFU) and the virulent type 3 parental strain (LD$_{50}$, ~10$^5$), the type 3 derivatives were not virulent even at doses of 10$^6$ CFU (FIG. 14B).

Switching of the type 6B capsule to type 3 resulted in a reduction of the LD$_{50}$ from >1 x 10$^6$ to ~6 x 10$^3$ CFU, a value that was similar to but still greater than the 7.5 x 10$^2$ value observed for the type 3 parent strain (FIG. 14C).

These results may be indicative of the role other factors play in pneumococcal virulence. For example, the type 5 capsule may represent one that results in high virulence with few other factors required, whereas the type 3 capsule may require the presence of other factors to be highly virulent. The introduction of the type 3 capsule into the type 5 genetic background may thus result in the expression of a virulent capsule but, in the absence of other necessary factors, in an avirulent strain. The increase in virulence of the type 6B strain suggests that the type 3 capsule is probably more virulent than the type 6B. However, its failure to become as virulent as the type 3 parent is suggestive of
a lack of other virulence factors in the 6B background. The type 2 recipient was only slightly more virulent than the type 3 donor and no significant change was noted in the virulence of its derivatives (FIG. 12). This result may suggest that the type 2 and type 3 capsules are of equal virulence and that the "accessory factors" necessary for full virulence are present in both strains.

Whether the decrease in virulence of type 5 derivatives is related to the alteration of cell chain length is not known. Clearly, the parent type 3 strain is highly virulent with a similar chain length. The alteration in chain length may reflect a general change in the surface structure of the type 5 strains possibly resulting from the change in capsule expression. Because the strains constructed were transformed with chromosomal DNA the inventors cannot rule out the possibility that determinants closely linked to the capsule locus are affecting the outcome of these studies. However, because several backcrosses were performed, and because independent isolates exhibited identical characteristics, it is unlikely that unlinked determinants are responsible for the results.

**EXAMPLE 19**

**Increased Virulence of S. pneumoniae type 6B by Inactivation of plpA.**

In Example 18, the introduction of the type 3-specific cassette and linked genes into an avirulent type 6B strain resulted in expression of the type 3 capsule and an increase in virulence. To more clearly define the contribution of the capsular serotype to the virulence of S. pneumoniae, insertion-duplication mutagenesis was used to insert an erythromycin marker adjacent to the type 6B-specific capsule cassette in the 3' flanking region.
Surprisingly, introduction of this insertion resulted in an increase in virulence of the type 6B strain (derivative LD$_{50}$ of $10^3$ versus parental LD$_{50}$ of $>10^6$, intraperitoneal). This enhancement of virulence could be attributed to the 1.2 kb SacI-HindIII fragment from the type 3 strain WU2 (pJD377) that was used to direct the erythromycin marker into the type 6B chromosome. Transformation of the wild type 6B strain with the SacI-HindIII fragment alone, followed by intraperitoneal infection of the transformation mixture into mice resulted in death in less than 24 hours. Identical results were obtained using multiple smaller fragments of the SacI-HindIII fragment. JD377 (Example 5) comprises the 3' end of cps3M and part of the gene pI$pA$. The fragments used contained mutations that, like the original insertion, resulted in a defective pI$pA$ in the type 6B strain. These data suggest that avirulence of type 6B observed via the intraperitoneal route is due to expression of pI$pA$, and that the increase in virulence of the type 6B strain expressing the type 3 capsule is the result of inactivation of the linked pI$pA$ and not expression of the type 3 capsule.

* * *

All of the compositions and methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations may be applied to the composition, methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. More specifically, it will be apparent that certain agents which are both chemically
and physiologically related may be substituted for the agents described herein while the same or similar results would be achieved. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the invention as defined by the appended claims.

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The following references, to the extent that they provide exemplary procedural or other details supplementary to those set forth herein, are specifically incorporated herein by reference.


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(ii) INVENTORS: YOTHER, Janet
              DILLARD, Joseph P.

(iii) TITLE OF INVENTION: STREPTOCOCCUS PNEUMONIAE
              CAPSULAR POLYSACCHARIDE
              GENES AND PLANKING REGIONS

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   (A) MEDIUM TYPE: Floppy disk
   (B) COMPUTER: IBM PC compatible
   (C) OPERATING SYSTEM: PC-DOS/MS-DOS/ASCII
(D) SOFTWARE: PatentIn Release #1.0, Version #1.30

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(A) APPLICATION NUMBER: UNKNOWN
(B) FILING DATE: CONCURRENTLY HEREWITHE
(C) CLASSIFICATION: UNKNOWN

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(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 300 base pairs
(B) TYPE: nucleic acid

(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "DNA"
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GAGCTCGGTA TTTTTTGGAA CGTGATTTAG TCTATGTAGT TGCAAGTGAC ATGCACAATT 60
5
TAGACAGTAG ACCTCCATAT ATGCAACAGG CATATGATAT CATTGCTAAG AAATATAGAG 120
CGAAAAAGAC GAAAGAACTT TTTGTAGATA ATCCCAAGAA AATTATAATG GATCATTAAT 180
TAGGAGAAAA TATGAAGGAA CAAAACACTT TGGAATCGA TGTTATGCAG TATTCAGAGC 240
10
TTATTGGAAG AAGTGTCATT TTATTAGTG G CATTATACTT CCTCAGTTGC TTTTTCTAC 300

(2) INFORMATION FOR SEQ ID NO:2:

15

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 261 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

20

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "DNA"
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

CTTCCGTATG CCGCGTTCTG CCAAGACCCCA ATCGGCTGGC TCTTTGATAG AGTAGCTGCC 60
5
CAAAAAATTA TCAGTATTAC TCCTGCGTATG CTCGGCAACTT CTGGAGGCAA GACCGCCTCATGAT 120
ATCACTGTTC CGCCCAATAAA TAAACGCAAAT CACACTATTG GTTTTTTGCT AGTTTTATTT 180
10
GGAACCTAGTG TCACTAGTTCT TCTCTCTGAA CTTTTGGACA CTCATCTGAA ACGTCCGAAA 240
GATATCGAAG ATACACTGCA G 261

(2) INFORMATION FOR SEQ ID NO:3:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 262 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

5  CAGCACTTCT CATCTGGCAC AGCTGATTAA TCTCAGGGCT TATGTGATAC AAATATTGAA  60
  AATTATTTTG TAGTTCAATC GGGATCTGTA TCACCAAACC CTACAGCCTT GTCAACAAAGC  120
  AAAAAATTTTG TGGTTATGGT AAAGCTTTTT TCAAAAGAGG TCAGTATATT GAGTTGGTGG  180

10  AAACGATAAA ATACATAATA TTTATTCCT TTGGTTATCA AATTAGCCCC TCCTGAAGCT  240
    CCCAATTGA CGGCTTGAGC TC  262

(2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 934 base pairs
    (B) TYPE: nucleic acid
    (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: other nucleic acid
(A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

5
GAGCTCCAAAT CAAAGGTTGT TTGTACACTT TTGACAGAG GGAATACTGC AGAGGACAGC 60
AAACAGCCAT AGTACGTAAA AATCCAGACA CCTAAAGCAG ACAAAAGGTT TGCCATCGAG 120

10 TATAAAATCA TGTAGGGGAC GTTACCGTAG GTGCAGTTGCG GTAGAGAAT GTGTGTAGCC 180
AAAACATCAA GAGTACCGTT AGTTATCAGA AAGTTAATAA AAGAGAGAC GCTAAAAATG 240

GTAAAAGAG TGAGTGGGCC AAAATGAAGA AGTCTTTTGG GGTTAATCC CATGAGAGTG 300

15 GTGCGATGGA GGAAGAGAAA AGCAATAGCC AGCAGGTCAA TATGTATTTT GGTGCCGATAA 360
CCAATTCCAA TGGCTAGAGC AATGGCGCTA ATCATTATTA AATGAATCAT TGAATTGTCC 420

20 TTTAGTTAGA ATATAAGAAG AGGATAGATT GAAGTTGGAG AATACTGGGT GTCTTCTGAT 480
GTAAAGTGGT TGTTCAAAA CCACTCCCAA TGGCATAAAT TGCAATGGAA TTGAATGAC 540
-155-

5 TGTTATCATC ATGGGAGG ATGTTTAC AAGTCAAGC GTCTGATT TATCTTTA

600 CCTTGGCT ATGGTCCG CTTGTGCC ATGTGCGG GCCCGGGG CACCGAAG

660 TATCATGGG TTATTTGAGG TGGTTTNGG TGAAGGACT CATCAAGC CATACCC ACCGATACG

720 AGCTCGTTC AAGCTTATC TTAGGAAAA ATGGGTTTAT ATTCTTAA ATGGATGTA

780 TTTCTAAA AAGACCGAT CTTGACGTC AACAAGGCT ATAGTCTGATT ATACGTGA

840 GATTTTTG AAAAGACGTT TACGAGAATA ATCGTATAC GCCTTATCTT TCCAAGCTG

900 ATATAGGC ACAAAAAAC GTTGTATG C CCC

934

(1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 4951 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(2) INFORMATION FOR SEQ ID NO: 5:
(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = "DNA"

(x) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

5  TTGACAATAG ATAAATTAT TATATAATTA AACTATTGCT TTTAAATAA AGTGAGAATA  60
   TTAATAATGC AGAGAAAGAG GACTTGATGA AAATGAAAAAT TGCCATTGCA GGAAGTGGTT
10  ATGTAGGCTCT GTCTTTAGGC GTGGCTACTAG CTCAACATCA TGAAGTTAAG GTCATGGATG
   TTATAAAGGA TAAGGTAGAG TCGATAAAAAC ATAGAAAAATC TCCAAATAG GATGAGGCGA
15  TTGAGAAATA CTTAGTTGAA AAAGAGTGA ATCTTGAAAG CTAACTTAGT CCTGCACACG
   TTTATAAAGA CGTGGAGTAT GCTATTATGT CTACTGGCGA TAATTATGAT GTAGACTTAA
20  ATGCAGTTGA TACATCTCTC GGTGGAGCTG CTATCAGAGC TTGTATGGA TATAATGATA
   CTTGATCAAT CGTAATCAGA AGTACTATTC CTGAAGGTA TACTAAAGAA GTGAGGGAAA
25  AGTATAATAC AGATCGTATT ATTTTTTCTC CAGAGTTTCT ACGTGAATCC AAAGCTTTAT  540
ATGATAATTT GTATCCATCT AGAATTTGTG TAGGAACCTGA TTTGGATGAT TCTGAGTTAA 600
CAAAAAAGAGC ATGGCAGTTT GCAGATCTAC TTAAGGTTGG AGCTATTAGA GAAGAGGTTC 660
CGATACCTGGT TGTTGCTTTT AATGAAGCAG AGGTTGCAAA ATTTGTTTGT AACACTTACT 720
tGGCAACTCG CGTAGCTTAT TTTAATGAGA TAGATACATA TAGCGAGGTA AAAGGGCTTA 780
ATCCCAAGAC AATTATGGAT ATTTGGTTTT ATGGATCTAG AATTGGATCA GACTATAATA 840
ACCCTAGCTT TGTTACCGGA GGATATGCT TACCAAAGGA CACAAAGCAA TTGAAGCAGA 900
GTTTTAGGGA TGTTCTGAA AATCTGATTA CAGCTGTGGT GCAATCTAAT AAAACAAGAA 960
AAAGTTTATAT AGCTGGAGCT ATTCTTAGCT AACAACCTAG TGTTGTAGGT ATTTATAGAT 1020
TAATTATGAA ATCTGATTCT GATAATTTTC GTTCTAGTGC TGTTAAGGGA GTTATGGGAAC 1080
GTTTGGACAA TTATGGTTAA GAAATTGTTA TTTACGAACC TACTATTGAG TGTGATACCT 1140
TTATGGGATA CAGAGTAATTT AAATCTTATT ATGAATTTAA GAATATTTCT GACATTGTGG 1200
TAGCGAATCG TATGAACGAT GATTTAAGGG ATATAACAAGA AAAACTCTAT ACACCGCATT 1260
AACGTCATTA CAGGTTTCAA ATTACATGCT GAGAAAATTC AAGAATTGA AGAAAAACAT 4920

AACCTACATG TTTGTTTGTG AAGGAAGCTG A 4951

5

(2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1307 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

TGAGTATTATT GATTGCCCTCC AGCTTGGAGT TAAATAAGGG CATCTGGATG GCATTTGTCA 60

CGTATTTTCT GTAGGCGACC AGCGTGCTAA GAGCAGTTCT AAAGGCTTGA TTGAGTTGGG 120

GAAAAGCTC AGTTAACAAG TCAAAGAAAT GGTCGGCATG CTTTCTTGGC AGGTGAAAA 180
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<td>840</td>
<td>CAAAACCTTT GCGGATAG CCAAAGAGAAA ATGTGCACT TATGGGAGAT AATTGGAAG</td>
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<td>TGTGATTTTT GGGATTCTC AGGAAGTCG TCAATAGT GCAGCTTC TGAGAAGCT AAGTTTAT</td>
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GCCAGTTGAC CAGACAGCAA CTACAAAAGT TCAGCGCGTC CAATCTATGA AACAATCCTT 960
GGAAGTAACCT TTAGGAGCTG ATAATGTCAT TATGATATC CAACAACTAC AAAAGACGA 1020
AGTAAACAAT ATTACATATT TTGCTGAAAA TGCTGCTGGC GAAGACTGGG ATTTATCAGA 1080
TAATGTCGGT TGGGTCCAG ACTTTGCCGA TCCATCAACC TACCTTGATA TCATCAAACC 1140
ATCTGTAGGA GAAAGTACTA AAAACATATT AGGGTTTGAC TCAGGGGAAG ATAAATGAGC 1200
TGCTAAAAAA GTAGGCTAT ATGACTACGA AAAATTGTTT ACTGAGGCTG GTGATGAGGC 1260
TACAGATGTT CGTAAACGCT ATGATAAATA CGCTGCAGCC CAAGCTT 1307

(2) INFORMATION FOR SEQ ID NO: 7:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 58 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

```
Ala Arg Tyr Phe Leu Glu Arg Asp Leu Val His Val Val Ala Ser Asp
 1        5       10       15
Met His Asn Leu Asp Ser Arg Pro Pro Tyr Met Gln Gln Ala Tyr Asp
 20      25     30
Ile Ile Ala Lys Lys Tyr Arg Ala Lys Lys Ala Lys Glu Leu Phe Val
 35    40     45
Asp Asn Pro Arg Lys Ile Ile Met Asp His
 50   55
```

(2) INFORMATION FOR SEQ ID NO:8:

20  (i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 40 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Leu Gly Asn Met Lys Glu Glu Asn Thr Leu Glu Ile Asp Val Leu
1      5
Gln Tyr Ser Glu Leu Ile Gly Ser Val Ile Leu Leu Val Ala Leu
10     20
Tyr Phe Ser Cys Phe Phe
35     40

(2) INFORMATION FOR SEQ ID NO:9:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 87 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear
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<td>Arg Val Ala Gln Lys Ile Ile Ser Ile Thr Arg Ala Asp Val Ala</td>
<td>20 25 30</td>
</tr>
<tr>
<td>His Trp Arg Ser Lys Thr Ala Asp Ile Thr Ala Ser Pro Asn Lys</td>
<td>35 40 45</td>
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<tr>
<td>Arg Asn Thr Leu Ile Gly Phe Leu Ala Phe Ile Gly Thr Ser Val</td>
<td>50 60 65</td>
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<tr>
<td>Ile Val Leu Leu Leu Leu Leu Asp Thr His Val Lys Arg Pro Glu</td>
<td>70 75 80</td>
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(2) INFORMATION FOR SEQ ID NO:10:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 43 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS: single
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(x) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Gln His Phe Ser Ser Gly Thr Ala Asp Leu Ser His Gly Leu Cys Asp
  1  5 10  15

Thr Asn Ile Glu Asn Leu Phe Val Val Val Gln Ser Gly Ser Val Ser Pro
  20 25 30

Asn Pro Thr Ala Leu Ser Gln Ser Lys Asn Phe
  20 35 40

(2) INFORMATION FOR SEQ ID NO:11:
(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 394 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Met Lys Ile Ala Ile Ala Gly Ser Gly Tyr Val Gly Leu Ser Leu Ala
1  5  10  15

Val Leu Leu Ala Gln His His Glu Val Lys Val Ile Asp Val Ile Lys
15 20 25 30

Asp Lys Val Glu Ser Ile Asn Asn Arg Lys Ser Pro Ile Lys Asp Glu
35 40 45

Ala Ile Glu Lys Tyr Leu Val Glu Lys Glu Leu Asn Leu Glu Ala Ser
50 55 60
Leu Asp Pro Ala His Val Tyr Lys Asp Val Glu Tyr Ala Ile Ile Ala
65
70
75
80

Thr Pro Thr Asn Tyr Asp Val Asp Leu Asn Gln Phe Asp Thr Ser Ser
85
90
95

Val Glu Ala Ala Ile Lys Thr Cys Met Glu Tyr Asn Asp Thr Cys Thr
100
105
110

Ile Val Ile Lys Ser Thr Ile Pro Glu Gly Tyr Thr Lys Glu Val Arg
115
120
125

Glu Lys Phe Asn Thr Asp Arg Ile Ile Phe Ser Pro Glu Phe Leu Arg
130
135
140

Glu Ser Lys Ala Leu Tyr Asp Asn Leu Tyr Pro Ser Arg Ile Val Val
145
150
155
160

Gly Thr Asp Leu Asp Asp Ser Glu Leu Thr Lys Arg Ala Trp Gln Phe
165
170
175

Ala Asp Leu Leu Lys Gly Gly Ala Ile Lys Glu Glu Val Pro Ile Leu
180
185
190
Val Val Ala Phe Asn Glu Ala Glu Val Ala Lys Leu Phe Ser Asn Thr
195       200       205

Tyr Leu Ala Thr Arg Val Ala Tyr Phe Asn Glu Ile Asp Thr Tyr Ser
210       215       220

Glu Val Lys Gly Leu Asn Pro Lys Thr Ile Ile Asp Ile Val Cys Tyr
225       230       235       240

Asp Pro Arg Ile Gly Ser Asp Tyr Asn Asn Pro Ser Phe Gly Tyr Gly
245       250       255

Gly Tyr Cys Leu Pro Lys Asp Thr Lys Gln Leu Lys Ala Ser Phe Arg
260       265       270

Asp Val Pro Glu Asn Leu Ile Thr Ala Val Val Gln Ser Asn Lys Thr
275       280       285

Arg Lys Asp Tyr Ile Ala Gly Ala Ile Leu Ala Lys Gln Pro Ser Val
290       295       300

Val Gly Ile Tyr Arg Leu Ile Met Lys Ser Asp Ser Asp Asn Phe Arg
305       310       315       320
Ser Ser Ala Val Lys Gly Val Met Glu Arg Leu Asp Asn Tyr Gly Lys
   325  330  335

Glu Ile Val Ile Tyr Glu Pro Thr Ile Glu Cys Asp Thr Phe Met Gly
   340  345  350

Tyr Arg Val Ile Lys Ser Leu Asp Glu Phe Lys Asn Ile Ser Asp Ile
   355  360  365

Val Val Ala Asn Arg Met Asn Asp Asp Leu Arg Asp Ile Gln Glu Lys
   370  375  380

Leu Tyr Thr Arg Asp Leu Phe Gly Arg Glu
   385  390

(2) INFORMATION FOR SEQ ID NO:12:

(i) SEQUENCE CHARACTERISTICS:

   (A) LENGTH: 416 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS: single
   (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein

(xii) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Met Tyr Thr Phe Ile Leu Met Leu Leu Asp Phe Phe Gln Asn His Asp
1      5       10       15

Phe His Phe Phe Met Leu Phe Phe Val Phe Ile Leu Ile Arg Trp Ala
20     25      30

Val Ile Tyr Phe His Ala Val Arg Tyr Ser Tyr Ser Cys Ser Val
35     40      45

Ser Asp Glu Lys Leu Phe Ser Ser Ser Val Ile Ile Pro Val Val Asp Glu
50     55      60

Pro Leu Asn Leu Phe Glu Ser Val Leu Asn Arg Ile Ser Arg His Lys
65     70      75      80

Pro Ser Glu Ile Ile Val Val Ile Asn Gly Pro Lys Asn Glu Arg Leu
85     90      95
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Tyr Val Leu Leu Gly Met Ile Phe Ser Phe Gly Gly Arg Asn Phe Lys
355 360 365

Ala Met Ser Arg Met Lys Trp Tyr Tyr Val Phe Leu Ile Pro Val Phe
370 375 380

Ile Ile Val Leu Ser Ile Ile Met Cys Pro Ile Arg Leu Leu Gly Leu
385 390 395 400

Met Arg Cys Ser Asp Asp Leu Gly Trp Gly Thr Arg Asn Leu Thr Glu
405 410 415

(2) INFORMATION FOR SEQ ID NO:13:

15

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 306 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS: single
   (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met Lys Lys Val Lys Lys Ala Val Ile Pro Ala Ala Gly Leu Gly Thr
1       5       10       15

Arg Phe Leu Pro Ala Thr Lys Ala Leu Ala Lys Glu Met Leu Pro Ile
20      25      30

Val Asp Arg Pro Thr Ile His Phe Val Ile Glu Glu Ala Leu Arg Ser
10      35      40      45

Gly Ile Glu Asp Ile Leu Val Val Thr Gly Lys Ser Lys Arg Ser Ile
15      50      55      60

Glu Asp Tyr Phe Asp Ser Thr Phe Glu Leu Glu Tyr Ser Leu Arg Lys
20      65      70      75      80

Gln Gly Lys Met Glu Leu Leu Lys Ser Val Asn Glu Ser Thr Asp Ile
20      85      90      95

Lys Val His Phe Val Arg Gln Ser Ser Pro Arg Gly Leu Gly Asp Ala
100     105     110
Val Leu Gln Ala Lys Ser Phe Val Gly Asp Asp Pro Phe Val Val Met 115 120 125

Leu Gly Asp Asp Leu Met Asp Ile Thr Asp Ser Thr Ala Val Pro Leu 130 135 140

Thr Arg Gln Leu Met Asp Asp Tyr Asn Ala Thr Gln Ala Ser Thr Ile 145 150 155 160

Ala Val Met Pro Val Arg Tyr Glu Asp Val Ser Ser Tyr Gly Val Ile 165 170 175

Ser Pro Arg Leu Glu Ser Ser Asn Gly Leu Tyr Ser Val Asp Ala Phe 180 185 190

Val Glu Lys Pro Lys Pro Glu Glu Ala Pro Ser His Leu Ala Ile Ile 195 200 205

Gly Arg Tyr Leu Leu Thr Pro Glu Ile Phe Ser Ile Leu Glu Thr Gln 210 215 220

Lys Pro Gly Ala Gly Asn Glu Ile Gln Leu Thr Asp Ala Ile Asp Thr 225 230 235 240
Leu Asn Lys Thr Gln Ser Val Phe Ala Arg Glu Phe Val Gly Lys Arg
245  250  255

Tyr Asp Val Gly Asp Lys Phe Asn Phe Met Lys Thr Ser Ile Asp Tyr
260  265  270

Ala Leu Gln His Pro Gln Ile Lys Glu Ser Leu Lys Asn Tyr Val Ile
275  280  285

Ala Leu Gly Lys Gln Leu Glu Lys Leu Asp Asp Cys Ser Ser Ser Gly
290  295  300

His Leu
305

(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:
   (A) LENGTH: 397 amino acids
   (B) TYPE: amino acid
   (C) STRANDEDNESS: single
   (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

```plaintext
5  Met Asn Cys Ile Glu Ser Tyr Gln Lys Trp Leu Asn Val Pro Asp Leu
   1   5        10       15

   Pro Ala Tyr Leu Lys Asp Glu Leu Leu Ser Met Asp Asp Lys Thr Lys
   20   25   30

10  Glu Asp Ala Phe Tyr Thr Asn Leu Glu Phe Gly Thr Ala Gly Met Arg
    35   40    45

   Gly Tyr Ile Gly Ala Gly Thr Asn Arg Ile Asn Ile Tyr Val Val Arg
   50   55    60

15  Gln Ala His Arg Arg Pro Cys Lys Leu Val Glu Ser Lys Gly Glu Thr
    65   70    75    80

20  Ala Lys Lys Ala Gly Val Ala Ile Ala Tyr Asp Ser Arg His Phe Ser
    85   90    95
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Pro Glu Phe Ala Phe Glu Ser Ala Gln Val Leu Ala Ala His Gly Ile 100 105 110
Lys Ser Tyr Val Phe Glu Ser Leu Arg Pro Thr Pro Glu Leu Ser Phe 115 120 125
Ala Val Arg His Leu Gly Ala Phe Ala Gly Ile Met Val Thr Ala Ser 130 135 140
His Thr Pro Ala Pro Phe Asn Gly Tyr Lys Val Tyr Gly Ser Asp Gly 145 150 155 160
Gly Gln Met Leu Pro Ala Asp Tyr Ile Arg Ala Ile Asp Asn Pro Phe 165 170 175
Ala Val Val Leu Ala Asp Leu Glu Glu Ala Lys Ser Thr Gly Leu Ile 180 185 190
Glu Val Ile Gly Glu Thr Leu Asp Ala Ala Tyr Leu Glu Glu Val Lys 195 200 205
Ser Val Asn Ile Asn Gln Asp Leu Ile Asp Gln Tyr Gly Arg Asp Met 210 215 220
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The sequence continues beyond this point, but it is not fully visible in the image.
Thr Glu Leu Val Thr Lys Ile Ala Glu Ser Tyr Gly Glu Pro Cys Leu
355 360 365

Thr Ser Leu Gln Val Ser Asn Ser Ser Leu Arg Lys Phe Lys Asn Leu
370 375 380

Lys Lys Asn Ile Thr Tyr Met Phe Gly Phe Glu Glu Ser
385 390 395

10

(2) INFORMATION FOR SEQ ID NO:15:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 274 amino acids
(B) TYPE: amino acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xii) SEQUENCE DESCRIPTION: SEQ ID NO:15:
Gln Asp Ser Ile Thr Tyr Leu Val Gly Gln Ile Phe Asp Arg Gln Ser
1     5     10     15

Tyr Lys Tyr Thr Ser Lys Thr Arg Glu Glu Gln Lys Thr Ser Thr Lys
5     20    25     30

Lys Ala Leu Leu Asn Lys Asp Phe Arg Gln Ala Ile Ala Phe Gly Phe
35    40    45

Asp Arg Tyr Ala Ser Gln Leu Asn Gly Gln Thr Gly Ala Ser Lys Ile
10    50    55    60

Leu Arg Asn Leu Phe Val Pro Pro Thr Phe Val Gln Ala Asp Gly Lys
15    65    70    75    80

Asn Phe Gly Asp Met Ala Lys Glu Lys Leu Val Thr Tyr Gly Asp Glu
85    90    95

Trp Lys Asp Val Asn Leu Ala Asp Ser Gln Asp Gly Leu Tyr Asn Pro
20    100   105   110

Glu Lys Ala Arg Ala Glu Phe Arg Lys Ala Lys Leu Ala Leu Gln Ala
115   120   125
Asp Glu Ala Thr Asp Val Arg Lys Arg Tyr Asp Lys Tyr Ala Ala Ala
260 265 270

Gln Ala
5

(2) INFORMATION FOR SEQ ID NO:16:

10  (i) SEQUENCE CHARACTERISTICS:
     (A) LENGTH: 177 amino acids
     (B) TYPE: amino acid
     (C) STRANDEDNESS: single
     (D) TOPOLOGY: linear

15  (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

20  Leu Asn Ala Tyr Phe Gln Met His Lys Ser Ser Leu Ile Asp Phe Ile
1   5   10   15
Ile Gln Met Pro Tyr Ser Asn Ser Lys Leu Glu Ala Ile Asn Lys Leu
145 150 155 160

Ile Ser Phe Leu Gln Thr Gln Thr Cys Arg Leu Cys Phe Ser Ser Asn
5 165 170 175

Ser

10 (2) INFORMATION FOR SEQ ID NO:17:

(i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 18 base pairs
    (B) TYPE: nucleic acid
    (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
    (A) DESCRIPTION: /desc = "DNA"

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

GCCACTATCG ACTACGCG
- 190 -

(2) INFORMATION FOR SEQ ID NO:18:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 18 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

TCATTTGATA TGCCTCCG

15

(2) INFORMATION FOR SEQ ID NO:19:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 20 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid

(A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

GTGAGATAAA TAGTAGTGC

30

(2) INFORMATION FOR SEQ ID NO:20:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 20 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid
   (A) DESCRIPTION: /desc = "DNA"

(xii) SEQUENCE DESCRIPTION: SEQ ID NO:20:

10  TCCAGCTCGT GTCATAATCT  20
1. A nucleic acid segment less than about 10 kb in length that comprises a non-type specific \textit{S. pneumoniae} \textit{cps} gene flanking region of sufficient length to allow hybridization under standard hybridization conditions to a \textit{S. pneumoniae} \textit{cps} gene flanking region.

2. The nucleic acid segment of claim 1, wherein the segment comprises a non-type specific \textit{S. pneumoniae} \textit{cps} gene 5' flanking region.

3. The nucleic acid segment of claim 2, wherein the segment includes a non-type specific \textit{S. pneumoniae} \textit{cps} gene 5' flanking region encoding for a peptide comprising SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9 or SEQ ID NO:10.

4. The nucleic acid segment of claim 3, wherein the segment includes a non-type specific \textit{S. pneumoniae} \textit{cps} gene 5' flanking region encoding for a peptide comprising SEQ ID NO:7.

5. The nucleic acid segment of claim 3, wherein the segment includes a non-type specific \textit{S. pneumoniae} \textit{cps} gene 5' flanking region encoding for a peptide comprising SEQ ID NO:8.

6. The nucleic acid segment of claim 3, wherein the segment includes a non-type specific \textit{S. pneumoniae} \textit{cps}
gene 5' flanking region encoding for a peptide comprising SEQ ID NO:9.

7. The nucleic acid segment of claim 3, wherein the segment includes a non-type specific S. pneumoniae cps gene 5' flanking region encoding for a peptide comprising SEQ ID NO:10.

8. The nucleic acid segment of claim 2, wherein the segment comprises a non-type specific S. pneumoniae cps gene 5' flanking region having a sequence that corresponds to at least a 60 nucleotide long contiguous stretch of SEQ ID NO:4.

9. The nucleic acid segment of claim 8, wherein the segment comprises a non-type specific S. pneumoniae cps gene 5' flanking region having a sequence that corresponds to at least a 100 nucleotide long contiguous stretch of SEQ ID NO:4.

10. The nucleic acid segment of claim 9, wherein the segment comprises a non-type specific S. pneumoniae cps gene 5' flanking region having a sequence that corresponds to at least a 500 nucleotide long contiguous stretch of SEQ ID NO:4.

11. The nucleic acid segment of claim 10, wherein the segment comprises a non-type specific S. pneumoniae cps gene 5' flanking region having a sequence that corresponds to SEQ ID NO:4.
12. The nucleic acid segment of claim 2, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region.

13. The nucleic acid segment of claim 12, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region having a sequence that corresponds to at least a 15 nucleotide long contiguous stretch of SEQ ID NO:6.

14. The nucleic acid segment of claim 13, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region having a sequence that corresponds to at least a 30 nucleotide long contiguous stretch of SEQ ID NO:6.

15. The nucleic acid segment of claim 14, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region having a sequence that corresponds to at least a 60 nucleotide long contiguous stretch of SEQ ID NO:4.

16. The nucleic acid segment of claim 15, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region having a sequence that corresponds to at least a 100 nucleotide long contiguous stretch of SEQ ID NO:6.

17. The nucleic acid segment of claim 16, wherein the segment comprises a non-type specific *S. pneumoniae* cps gene 3' flanking region having a sequence that
corresponds to at least a 500 nucleotide long contiguous stretch of SEQ ID NO:6.

18. The nucleic acid segment of claim 17, wherein the segment comprises a non-type specific \textit{S. pneumoniae cps} gene 3' flanking region having a sequence that corresponds to SEQ ID NO:6.

19. The nucleic acid segment of claim 1, wherein the segment comprises a non-type specific \textit{S. pneumoniae cps} gene 5' flanking region and a non-type specific \textit{S. pneumoniae cps} gene 3' flanking region.

20. The nucleic acid segment of claim 19, wherein the segment comprises a 5' flanking region that encodes for a peptide comprising SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9 or SEQ ID NO:10 and a 3' flanking region sequence that corresponds to at least a 30 nucleotide long contiguous stretch of SEQ ID NO:6.

21. The nucleic acid segment of claim 20, wherein the segment comprises a 5' flanking region that encodes for a peptide comprising SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9 or SEQ ID NO:10 and a 3' flanking region sequence that corresponds to at least a 100 nucleotide long contiguous stretch of SEQ ID NO:6.

22. The nucleic acid segment of claim 21, wherein the segment comprises a 5' flanking region that encodes for a peptide comprising SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9
or SEQ ID NO:10 and a 3' flanking region sequence that corresponds to SEQ ID NO:6.

23. The nucleic acid segment of claim 19, wherein the segment comprises a 5' flanking region sequence that corresponds to at least a 60 nucleotide long contiguous stretch of SEQ ID NO:4 and a 3' flanking region sequence that corresponds to at least a 30 nucleotide long contiguous stretch of SEQ ID NO:6.

24. The nucleic acid segment of claim 23, wherein the segment comprises a 5' flanking region sequence that corresponds to at least a 100 nucleotide long contiguous stretch of SEQ ID NO:4 and a 3' flanking region sequence that corresponds to at least a 100 nucleotide long contiguous stretch of SEQ ID NO:6.

25. The nucleic acid segment of claim 24, wherein the segment comprises a 5' flanking region sequence that corresponds to SEQ ID NO:4 and a 3' flanking region sequence that corresponds to SEQ ID NO:6.

26. The nucleic acid segment of claim 1, further defined as including a type specific \textit{S. pneumoniae} \textit{cps} gene region of sufficient length to allow hybridization to a \textit{S. pneumoniae} \textit{cps} gene region under standard hybridization conditions.

27. The nucleic acid segment of claim 26, further defined as less than about 5,000 nucleotides in length.
28. The nucleic acid segment of claim 27, further defined as less than about 1,000 nucleotides in length.

29. The nucleic acid segment of claim 1, further defined as a recombinant vector.

30. A nucleic acid cassette less than about 20 kb in length that comprises a non-type specific S. pneumoniae cps gene 5' flanking region sequence and a non-type specific S. pneumoniae cps gene 3' flanking region sequence, the flanking region sequences being of sufficient length to allow hybridization under standard hybridization conditions to a S. pneumoniae cps gene flanking region.

31. A nucleic acid segment of up to about 20 kb in length, comprising a S. pneumoniae cps gene region of sufficient length to allow hybridization to a S. pneumoniae cps gene region under standard hybridization conditions.

32. The nucleic acid segment of claim 31, further defined as comprising a cpsB gene.

33. The nucleic acid segment of claim 31, further defined as comprising a cpsC gene.

34. The nucleic acid segment of claim 31, further defined as comprising a cpsE gene.
35. The nucleic acid segment of claim 31, further defined as comprising a \textit{cpsD} gene.

36. The nucleic acid segment of claim 31, further defined as comprising a \textit{cpsS} gene.

37. The nucleic acid segment of claim 31, further defined as comprising a \textit{cpsU} gene.

38. The nucleic acid segment of claim 31, further defined as comprising a \textit{cpsM} gene.

39. The nucleic acid segment of claim 31, further defined as comprising a \textit{plpA} gene.

40. The nucleic acid segment of claim 31, further defined as comprising a \textit{tnpA} gene.

41. The nucleic acid segment of claim 31, further defined as comprising a complete \textit{S. pneumoniae cps} gene region.

42. The nucleic acid segment of claim 31, wherein the \textit{S. pneumoniae cps} gene region is defined as a Type 3 \textit{cps} gene region.
43. The nucleic acid segment of claim 42, further defined as comprising a Type 3 \textit{cpsB}, \textit{cpsC}, \textit{cpsE}, \textit{cpsD}, \textit{cpsS}, \textit{cpsU}, \textit{cpsM}, \textit{tnpA} and \textit{lpA} gene.

44. The nucleic acid segment of claim 31, further defined as comprising a \textit{cps} gene flanking region, wherein the flanking region corresponds to any nucleic acid segment in accordance with the foregoing claims 1 through 25.

45. The nucleic acid segment of claim 44, further defined as less than about 10,000 nucleotides in length.

46. The nucleic acid segment of claim 45, further defined as less than about 5,000 nucleotides in length.

47. The nucleic acid segment of claim 44, further defined as a DNA cassette bounded at each terminus by a PCR primer of known sequence or a restriction enzyme recognition site.

48. The nucleic acid segment of claim 47, wherein the segment is bounded by an \textit{SphI} or \textit{SalI} site.

49. The nucleic acid segment of claim 44, further defined as a recombinant vector.

50. The nucleic acid segment of claim 49, further defined as recombinant vector comprising at least one
S. pneumoniae cps gene and sufficient flanking region to allow homologous recombination of the fragment in a S. pneumoniae host cell.

51. The nucleic acid segment of claim 50, further defined as comprising a complete S. pneumoniae cps gene region.

52. A recombinant host cell comprising a recombinant vector comprising a nucleic acid segment in accordance with claim 44.

53. The recombinant host cell of claim 52, further defined as a recombinant E. coli host cell.

54. The recombinant host cell of claim 52, further defined as a recombinant gram positive host cell.

55. The recombinant host cell of claim 54, further defined as a Bacillus, Staphylococcus, or Streptococcus host cell.

56. The recombinant host cell of claim 55, further defined as a recombinant S. pneumoniae host cell.

57. A recombinant host cell in accordance with claim 52, further defined as including an engineered resistance gene.
58. A recombinant *S. pneumoniae* cell of a selected serotype, the cell expressing a *cps* gene of another *S. pneumoniae* serotype.


60. A method for preparing a recombinant host cell, comprising preparing a *S. pneumoniae cps* gene and transforming a host cell with said gene.

61. The method of claim 60, wherein the host cell is defined as a *S. pneumoniae* host cell, and the *cps* gene is introduced by a method comprising the steps of:

(a) preparing a DNA segment that includes a selected *S. pneumoniae cps* gene flanked by sufficient *S. pneumoniae* flanking regions to allow homologous recombination in the *S. pneumoniae* host;

(b) transforming the *S. pneumoniae* host with the DNA segment; and

(c) selecting a recombinant host that expresses the *S. pneumoniae cps* gene.

62. The method of claim 61, wherein the DNA segment is a plasmid.
63. The method of claim 61, wherein the host, prior to transformation, is a high producer of the capsular polysaccharides.

64. The method of claim 63, wherein corresponding cps gene of the host has been replaced by homologous recombination with the recombinant cps gene.

65. The method of claim 61, wherein the cell is selected by means of a resistance gene.

66. The method of claim 65, wherein the resistance gene is positioned in the non type specific cps region.

67. The method of claim 66, wherein the resistance gene is an erythromycin resistance gene.

68. A method for detecting \textit{S. pneumoniae} in a sample, comprising the steps of:

\begin{enumerate}
\item[(a)] obtaining nucleic acids from a sample suspected of containing \textit{S. pneumoniae};
\item[(b)] subjecting said nucleic acids to hybridization with a \textit{S. pneumoniae} cps nucleic acid segment comprising a cps gene flanking region or a cps gene coding region of sufficient length to allow hybridization to \textit{S. pneumoniae} cps nucleic acids under standard hybridization conditions; and
\end{enumerate}
(c) detecting the hybridized nucleic acids.

69. The method of claim 68, wherein said *S. pneumoniae* cps nucleic acid segment comprises a non-type specific *S. pneumoniae* cps gene flanking region of sufficient length to allow hybridization under standard hybridization conditions to a *S. pneumoniae* cps gene flanking region.

70. The method of claim 68, wherein the nucleic acids from said sample are subjected to restriction enzyme digestion and size separation prior to hybridization with said *S. pneumoniae* cps nucleic acid segment.

71. The method of claim 70, wherein the nucleic acids are subjected to SphI digestion.

72. The method of claim 68, wherein said detection of hybridized nucleic acid involves PCR.

73. A method for determining the capsule type of an unknown *S. pneumoniae* strain, comprising obtaining nucleic acids from the strain and hybridizing said nucleic acids with a *S. pneumoniae* cps DNA segment comprising either:

(a) a non-type specific *S. pneumoniae* cps gene flanking region of sufficient length to allow hybridization under standard hybridization conditions to a
S. pneumoniae cps gene flanking region; or

(b) a type specific S. pneumoniae cps gene region of sufficient length to allow hybridization to a S. pneumoniae cps gene under standard hybridization conditions.

74. A method of generating an antibody response, comprising administering to an animal an immunologically effective amount of a Cps peptide or protein.

75. The method of claim 74, wherein the Cps peptide or protein is encoded for by any one of the nucleic acid sequences in the foregoing claims 32 through 40.

76. A method for detecting S. pneumoniae in a sample, comprising the steps of:

(a) obtaining proteins from a sample suspected of containing S. pneumoniae;

(b) binding said proteins with an antibody;

(c) detecting the bound proteins.
77. The method of claim 76, wherein said antibody corresponds to an antibody directed against a Cps protein or peptide.

78. The method of claim 77, wherein said antibody is labeled.

79. The method of claim 75, wherein said proteins are separated by electrophoresis.

80. A method for preventing infection of a subject with \textit{S. pneumoniae} by administering a composition comprising an antibody directed against a Cps protein or peptide.
FIG. 1
FIG. 3A

FIG. 3B
FIG. 4
FIG. 5
\textbf{FIG. 6A}

\begin{verbatim}
\texttt{Cps3B}
GAG CT CCG TAT TTT TTG GAA CGT GAT TTA GTT CAT GTA GTT GCA AGT GAC ATG CAC AAT TTA 
ala arg tyr phel glul arg asp leu val his val val ala ser asp met his asn leu

GAC AGT AGA CCT CCA TAT ATG CAA CAG GCA TAT GAT ATC ATT GCT AAG AAA TAT AGA GCG
asp ser arg pro pro tyr met glin glin ala tyr asp ile ile ala lys lys tyr arg ala
stop Cps3B

AAA AAA GCG AAA GAA CTT TTT GTA GAT ATT CCC AGA AAA ATT ATA ATG GAT CAT TAA TTA
lys lys ala lys gluel phe val asp asp pro arg lys ile ile ile met asp his *** leu
\texttt{start Cps3C}
GGA GAA AAT ATG AAG GAA CAA AAG ACT TTT CTT GTA GAT GTA TTT CAG TAT TCA GAG CTT
gly glu asp met lys gluel glun asn thr leu glu ile asp leu glun tyr ser glu leu

ATT GGA aGa agt gtc att tta tta gTG gCA tta tAC ttc TTT aGC GAA CTA C C ile gly arg ser val ile leu val ala leu tyr phe ser cys phe phe leu

-200 nt gap in Cps3C

\texttt{Cps3C (continued to PstI site. -45 nt expected beyond PstI to Cps3C stop)}

CTT CCG TAT gcc gcc TCC TCG CAA gcc CAC AAT ATC GGC TGG CTC TTT CTT TTT GTA CTA GCG
leu pro tyr ala ala phe leu glul asp pro ile gly thr leu phe asp arg val ala ala

CAA AAA ATG AGT ATT ACT GCT GAT GTA GCA GCG TGG AGG AAG ACG AAG ACC GCC GAT

\texttt{PstI}

GAT ATG GAA GAT ACA TGG CAG
asp ile glul asp thr leu glin

-360 nt gap

\texttt{Cps3B}
CAC CAC ttc tca TCT GGC ACA GCT GAT tta TCT CAC GCC tta TGT GAT ACA AAT ATT gaa
Gln his phe ser ser gly thr ala asp leu ser his gly leu cys asp thr asn ile glu

AAA TTA TTT GTA GAT TAA CCA GCC TGG CCT GTA CCA AAG CCT ACA GCC TGG TCA CAA AGC
ala leu phe val val glul ser gly ser val ser pro asp pro thr ala leu ser ser glu

End homology

with Cps159d Repeat

AAA AAA TTT TGT GAT GTA AAG CTT TTT TCA AAA GAG GTC AGT ATT GTA TTT GAG GCG TGG
lys asn phe

AAA CAA TAA AAT ACA TAA TTT ATT CCT TGG GAT ATT CAA TAA CCA GCC CCT GAA GCT

\texttt{SacI}

CCC CAA TGG ACG GCT TGA GCT C

\textbf{FIG. 6B}

\textbf{FIG. 6C}
FIG. 9
glc-6-P \[\rightarrow\] phosphoglucomutase \[\rightarrow\] Cps3M

glc-1-P \[\rightarrow\] glucose-1-phosphate uridylyltransferase \[\rightarrow\] Cps3U

UTP \[\rightarrow\] UDP-gluc + PPI

NAD\(^+\) \[\rightarrow\] UDPG dehydrogenase \[\rightarrow\] Cps3D

NADH \[\rightarrow\] UDP-glucA

UDP-gluc \[\rightarrow\] type 3 capsular polysaccharide synthase \[\rightarrow\] Cps3S

(-glcA-glc-)\(_n\)
FIG. 11
FIG. 12
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C12N15/31 C07K14/315 C12N15/52 C12N15/53 C12N15/60
C12N15/54 C12N15/61 C12Q1/68 C12P19/04 A61K39/09

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 6 C07K C12N C12P C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>MOL GEN GENET 239 (1-2). 1993. 188-195, GARCIA, E. ET AL. 'CLONING AND SEQUENCING</td>
<td>1-11</td>
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<tr>
<td>X</td>
<td>OF A GENE INVOLVED IN THE SYNTHESIS OF THE CAPSULAR POLYSACCHARIDE OF STREPTOCOCCUS</td>
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<tr>
<td>X</td>
<td>-PNEUMONIAE TYPE 3.' cited in the application see figures 2,4</td>
<td></td>
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<td>A</td>
<td>MOLECULAR MICROBIOLOGY, vol. 9, no. 5, 1993 pages 1037-1050, PEARCE, B. ET AL.</td>
<td>12-14</td>
</tr>
<tr>
<td>A</td>
<td>'Genetic identification of exported proteins in Streptococcus pneumoniae' cited</td>
<td></td>
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<tr>
<td>A</td>
<td>in the application see figure 4, Expl</td>
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X Further documents are listed in the continuation of box C.

X Patent family members are listed in annex.

* Special categories of cited documents:

' A' document defining the general state of the art which is not considered to be of particular relevance

'B' earlier document but published on or after the international filing date

' L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

'O' document referring to an oral disclosure, use, exhibition or other means

' P' document published prior to the international filing date but later than the priority date claimed

' T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

' X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

' Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

' A' document member of the same patent family

Date of the actual completion of the international search: 11 September 1995

Date of mailing of the international search report: 05. 10. 95

Name and mailing address of the ISA
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Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl
Fax (+ 31-70) 340-3016

Authorized officer: Andres, S
<table>
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<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td>P,X</td>
<td>JOURNAL OF EXPERIMENTAL MEDICINE 181 (3). 973-983, March 1995. DILLARD, J. ET AL. 'Characterization of the Cassette Containing Genes for Type 3 Capsular Polysaccharide Biosynthesis in Streptococcus pneumoniae.' see the whole document</td>
<td>1-52</td>
</tr>
<tr>
<td>P,X</td>
<td>JOURNAL OF BACTERIOLOGY 176 (20). 6375-6383, October 1994. ARRECUBIETA, C. ET AL. 'Molecular characterization of cap3A, a gene from the operon required for the synthesis of the capsule of Streptococcus pneumoniae type 3: Sequencing of mutations responsible for the unencapsulated phenotype and localization of the capsular cluster on the pneumococcal chromosome.' cited in the application see figures 1,2</td>
<td>1-11</td>
</tr>
<tr>
<td>P,X</td>
<td>MOLECULAR MICROBIOLOGY 12 (6). 1994. 959-972, DILLARD, J. ET AL. 'Genetic and molecular characterization of capsular polysaccharide biosynthesis in Streptococcus pneumoniae type 3.' cited in the application see figure 4</td>
<td>1-52</td>
</tr>
<tr>
<td>P,X</td>
<td>INFECTION AND IMMUNITY 62 (5). 1813-1819, December 1994. KELLY, T. ET AL. 'Effect of genetic switching of capsular type on virulence of Streptococcus pneumoniae.' cited in the application see figures 1,2; table 3</td>
<td>1,2</td>
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<td>P,X</td>
<td>WO-A-95 06732 (UNIV ROCKEFELLER ; MASURE H ROBERT (US); PEARCE BARBARA J (US); TUO) 9 March 1995 see example 3 see SEQ IDs 1 and 46</td>
<td>12-14, 31,39</td>
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