Spa Contributes to the Virulence of Type 18 Group A Streptococci

DUNCAN G. J. McLELLAN,1,2 EDNA Y. CHIANG,1 HARRY S. COURTNEY,1,2 DAVID L. HASTY,1,3 SHIRLEY C. WEI,1,2 MARY C. HU,4 MICHAEL A. WALLS,4 JACKIE J. BLOOM,4 AND JAMES B. DALE1,2,5

VA Medical Center1 and Departments of Medicine2 and Anatomy and Neurobiology,3 University of Tennessee, Memphis, Tennessee 38104, and ID Biomedical Corporation, Bothell, Washington 980114

Received 22 September 2000/Returned for modification 5 December 2000/Accepted 12 February 2001

Streptococcal protective antigen (Spa) is a newly described surface protein of group A streptococci that was recently shown to evoke protective antibodies (J. B. Dale, E. Y. Chiang, S. Liu, H. S. Courtney, and D. L. Hasty, J. Clin. Investig. 103:1261–1268, 1999). In this study, we have determined the complete sequence of the spa gene from type 18 streptococci. Purified, recombinant Spa protein evoked antibodies that were bactericidal against type 18 streptococci, confirming the presence of protective epitopes. Sera from patients with acute rheumatic fever contained antibodies against recombinant Spa, indicating that the Spa protein is expressed in vivo and is immunogenic in humans. To determine the role of Spa in the virulence of group A streptococci, we created a series of insertional mutants that were (i) Spa negative and M18 positive, (ii) Spa positive and M18 negative, and (iii) Spa negative and M18 negative. The mutants, the parent M18 strain (18-282) were used in LPS assays to determine resistance to phagocytosis, growth in human blood, and mouse virulence. The results show that Spa is a virulence determinant of group A streptococci and that expression of both Spa and M18 is required for optimal virulence of type 18 streptococci.

Group A streptococci (GAS) are major human pathogens that cause a wide variety of illnesses, ranging from uncomplicated pharyngitis and pyoderma to life-threatening infections such as necrotizing fasciitis and toxic shock syndrome (23). The virulence of GAS is determined in part by their ability to resist opsonization by complement and phagocytic killing by neutrophils (14). It has been known for many years that these virulence characteristics are mediated in large part by the M protein on the surface of the organisms (16). In addition, the M proteins contain protective epitopes that elicit opsonic antibodies that promote phagocytic killing in the immune host (16). Previous studies have shown that insertional inactivation of the emm gene in some GAS serotypes results in an avirulent phenotype (7, 19).

In a recent study of type 18 streptococci, we found that inactivation of the emm 18 gene had only a minor effect on the ability of the mutant to grow in human blood and on the 50% lethal dose (LD90) in mice compared to the M-positive parent strain (10). We used the M-negative mutant to identify a new surface protein, streptococcal protective antigen (Spa), that was distinct from M protein and contained protective epitopes which elicited bactericidal antibodies (10). The present study was undertaken to determine whether Spa functions as a virulence determinant of type 18 streptococci. Using a series of M-negative and Spa-negative mutants, we show that both of these surface proteins contribute to the virulence of type 18 streptococci. In addition, we have cloned and sequenced the complete spa gene, which allowed a direct comparison of its structure to that of emm genes.

MATERIALS AND METHODS

Bacterial strains. The parent type 18 streptococcal strain 87-282 (designated 18-282) and its M-negative mutant, M18Δf, have been described previously (10, 11). Escherichia coli DH5α was used for all molecular cloning experiments except with plasmid pQE-30, which was maintained in E. coli M15.

Cloning, expression, and purification of recombinant Spa (rSpa). The first 636 bp of spa18, encoding the NH2-terminal half of the mature Spa protein, were amplified by PCR and ligated into the expression vector pQE-30 (Qiagen Inc., Valencia, Calif.) containing a six-histidine tag. The tag was used to transform E. coli M15, and clones were selected for ampicillin resistance and screened for expression of Spa protein by a colony blot assay with anti-Spa rabbit serum (10). One colony was selected for high-level expression and purification of the six-histidine-tagged Spa protein as described by the manufacturer (The Qiagen Expressionist; Qiagen). Eluted proteins were dialyzed against phosphate-buffered saline (PBS; 0.05 M NaH2PO4, 0.15 M NaCl [pH 7.4]), and purity was assessed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The recombinant protein migrated as a single band with an apparent molecular mass of 24 kDa.

Preparation of antisera. The purified rSpa fragment (200 μg) was mixed with the adjuvant Rehydregal (aluminum hydroxide low-viscosity gel; Reheis, Inc., Berkeley Heights, N.J.) and injected intramuscularly into New Zealand White rabbits. Booster injections were given at 4 and 8 weeks, and sera were collected at 0, 4, 6, 8, 9, and 10 weeks. Methods for preparation of antisera against purified native Spa (anti-Spa), a synthetic peptide copying the NH2-terminal 23 amino acids of Spa (anti-Spa [I–23]), and recombinant M18 (anti-M18) have been described previously (10).

PCR, cloning, and sequencing of spa18. The sequence of a 636-bp fragment of spa18 had been determined as described previously (10). At the 3′ end of this fragment, a 21-bp sequence (TCTCTTTCAGAGTCAGCAACA) was later found to be an inverted repeat of an upstream sequence. When used as a single primer with type 18 chromosomal DNA, the result was a PCR product of 1,191 bp, which was ligated into pCR2.1-TOPO (Invitrogen Corp., San Diego, Calif.). DNA sequencing was performed by automated techniques at the University of Tennessee Molecular Resources Center, using primers from the 5′ and 3′ flanking ends of the plasmid. The ampiclon was found to contain the original 636-bp fragment as well as 550 bp of unique 5′ sequence. A search of the GenBank database showed considerable homology of this region with the signal sequence and 5′ noncoding region of the Streptococcus equi emm gene (GenBank accession no. UT73162). Hypothesizing that there would be homologies between the C termini, we constructed a reverse primer from the 3′ noncoding region of the S. equi emm gene (GCCCTAGTCTGAGGCCC) for use in combination with a...
TABLE 1. Association of 18-282, M18V, 18Spa−, and 18ISpa− with neutrophils and opsonization by anti-rSpa and anti-rM18

<table>
<thead>
<tr>
<th>Antiserum</th>
<th>18-282</th>
<th>M18V</th>
<th>18Spa−</th>
<th>18ISpa−</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preimmune</td>
<td>2</td>
<td>6</td>
<td>2</td>
<td>98</td>
</tr>
<tr>
<td>Anti-rSpa</td>
<td>96</td>
<td>98</td>
<td>0</td>
<td>94</td>
</tr>
<tr>
<td>Anti-rM18</td>
<td>100</td>
<td>10</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

* A standard inoculum of streptococci grown to early log phase were added to 0.05 ml of test antiserum and incubated for 15 min at 37°C; 0.2 ml of nonimmune, heparinized human blood was added and rotated end-over-end for 45 min at 37°C. Following rotation, smears were made on glass slides and stained with modified Wright’s stain. The percentage of neutrophils associated with streptococci was calculated by counting 50 consecutive neutrophils. Assays were performed at least three different times, and the results presented are representative of all experiments.

model was assessed after intraperitoneal challenge infections of Swiss white mice. Four groups of five mice each were challenged with 10-fold-increasing amounts of exponential growth-phase organisms of each strain. Deaths were recorded for 7 days after challenge infections, and the LD₅₀ was determined by the method of Reed and Muench (22).

ELISAs. Microtiter wells (Nunc-Immuno modules; Nalge Nunc International, Roskilde, Denmark) were coated with recombinant NH₂-terminal peptides of SpA, M10, M2, M3, M5, M6, M12, M18, M19, and M24 (5 μg/ml in 0.05 M carbonate [pH 9.6]). The recombinant M protein peptides contained the NH₂-terminal 30 to 50 amino acids of the mature M proteins that were repeated in the form of a dimer. Wells without peptide but containing all other reagents served as negative controls. The enzyme-linked immunosorbent assays (ELISAs) were performed using human sera from patients with acute rheumatic fever that were obtained as part of an epidemiological investigation of streptococcal sequelae conducted in Saudi Arabia in the 1980s. The sera were serially diluted in PBS (pH 7.4) with 0.05% Tween 20, added to the wells, and incubated at 37°C for 2 h. After washing, 5-aminosalicylic acid was added, and the A₅₇₀ was determined at 15 min by an MR 600 microplate reader (Dynatech Laboratories Inc., Chantilly, Va.). Positive controls were performed by a similar method, although rabbit sera raised against recombinant M proteins and SpA protein were used as the first antibody and peroxidase-conjugated goat anti-rabbit immunoglobulins (ICN Biomedicals, Aurora, Ohio) were added to the wells. The sections were then incubated with HRP-conjugated peroxidase color development reagent (Bio-Rad) used as the substrate.

Oxidation and bactericidal assays. In vitro oxidation and bactericidal assays were performed as previously described (2). Briefly, oxidation assays were performed with whole, heparinized (10 U/ml) nonimmune human blood. A standard inoculum of streptococci, grown to log phase, was added to 0.05 ml of test antiserum and 0.2 ml of blood. For oxidation assays, the number of viable bacteria was calculated by counting 50 consecutive neutrophils. Assays were performed at least three different times, and the results presented are representative of all experiments.

Determination of LD₅₀. Lethality of GAS strains 18-282, M18V, 18Spa− (M positive, SpA negative), and 18ISpa− (M negative, SpA negative) in the murine model was assessed after intraperitoneal challenge infections of Swiss white mice. Four groups of five mice each were challenged with 10-fold-increasing amounts of exponential growth-phase organisms of each strain. Deaths were recorded for 7 days after challenge infections, and the LD₅₀ was determined by the method of Reed and Muench (22).

ELISAs. Microtiter wells (Nunc-Immuno modules; Nalge Nunc International, Roskilde, Denmark) were coated with recombinant NH₂-terminal peptides of SpA, M10, M2, M3, M5, M6, M12, M18, M19, and M24 (5 μg/ml in 0.05 M carbonate [pH 9.6]). The recombinant M protein peptides contained the NH₂-terminal 30 to 50 amino acids of the mature M proteins that were repeated in the form of a dimer. Wells without peptide but containing all other reagents served as negative controls. The enzyme-linked immunosorbent assays (ELISAs) were performed using human sera from patients with acute rheumatic fever that were obtained as part of an epidemiological investigation of streptococcal sequelae conducted in Saudi Arabia in the 1980s. The sera were serially diluted in PBS (pH 7.4) with 0.05% Tween 20, added to the wells, and incubated at 37°C for 2 h. After washing, 5-aminosalicylic acid was added, and the A₅₇₀ was determined at 15 min by an MR 600 microplate reader (Dynatech Laboratories Inc., Chantilly, Va.). Positive controls were performed by a similar method, although rabbit sera raised against recombinant M proteins and SpA protein were used as the first antibody and peroxidase-conjugated goat anti-rabbit immunoglobulins (ICN Biomedicals) were used as the second antibody. Antibody levels were graded as specified in the footnote to Table 3.

Assays for tissue cross-reactive antibodies. Rabbit antisera raised against the recombinant NH₂-terminal half of SpA (anti-rSpA) was tested for the presence of tissue cross-reactive antibodies by indirect immunofluorescence tests (9) using frozen sections (4 μm) of human myocardium, kidney, basal ganglia, and cerebral cortex. The sections were placed on gelatin-coated slides and fixed with 1% paraformaldehyde for 10 min. The slides were washed with PBS, incubated with anti-rSpA diluted 1:5 in PBS for 30 min at room temperature, and washed thoroughly in PBS. The sections were then incubated with fluorescein-conjugated goat IgG to rabbit IgG (Cappel, West Chester, Pa.) at a dilution of 1:40 in PBS for 30 min at room temperature. After washing, the slides were mounted in Gelvatol and examined in a fluorescence microscope (Carl Zeiss Inc., New York, Germany), where it was immobilized by UV irradiation. A fragment of pUC19 was ligated into pCR2.1-TOPO. This amplicon was sequenced as described above, using primers from flanking regions of known spa sequence.

- Anti-rM18 100 10 100 100
- Anti-rSpa 96 98 0 94
- Preimmune 2 6 2 98
- SpA + 18Spa2 M18V 100 100 100 100
- SpA+ 18Spa- 100 100 100 100
- SpA− 18Spa+ 100 100 100 100
- Spa negative 100 100 100 100
- Spa positive 100 100 100 100
- Spa negative 100 100 100 100

FIG. 1. Alignment of the complete sequences of spa18 and the emm gene of S. equi. The arrow indicates the end of the leader peptide and the start of the mature protein, which was previously determined by amino acid sequence analysis (10). The LPSTGE motif is underlined. The GenBank accession number for the S. equi emm gene is U73162, and that for spa is AF086813.
Rabbit antisera known to cross-react with human myocardium, kidney, and brain were used as positive controls (3, 9, 15).

Indirect immunofluorescence assays of C3 binding to streptococci. Patterns of C3 binding to the surface of type 18 streptococci and the mutant strains were determined by confocal immunofluorescence microscopy by methods previously described (11). Briefly, bacteria grown to mid-log phase were incubated in either fresh human serum or plasma, washed, and then incubated with fluorescein-labeled goat anti-human C3 (Cappel). Photographs were made using a Nikon PCM2000 confocal laser scanning microscope in the Integrated Microscopy Center at the University of Memphis.

GenBank accession number. The previous GenBank entry (accession no. AFO86813) containing the partial sequence of spa has been updated to include the complete sequence.

RESULTS

DNA and amino acid sequences of Spa. No homology was found between the spa18 sequence and current entries in the type 1 streptococcal genome sequence database (http://www .genome.ou.edu/strep.html). The predicted amino acid sequence of Spa demonstrates features similar to other streptococcal surface proteins, including a 37-residue signal peptide, an LPSTGE anchor motif (13), and a hydrophilic tail. The calculated molecular mass of the native protein is 58.9 kDa, and analysis of the secondary structure showed a high alphahelical potential. Contrary to the amino acid sequence of other M-like proteins, Spa does not contain a proline-rich C terminus or internal tandem repeats.

A search of GenBank entries revealed considerable homology between Spa and the M protein of S. equi (Fig. 1). The homology is most apparent in the leader peptides and the C-terminal halves of the proteins. There is no significant sequence homology in the NH2-terminal halves of the mature proteins.

Characterization of the Spa-negative mutants of strains 18-282 and M18Ω. Inactivation of the spa gene in strain 18-282 and its M-negative mutant M18Ω was achieved by integrational plasmid mutagenesis using pUC19ermspa. Interruption of spa18 was confirmed by PCR analysis of chromosomal DNA obtained from the Spa-negative mutants created from 18-282 and M18Ω, designated 18Spa18 and 18M18Vspa18. Three sets of primer pairs were used, with one primer in each pair annealing to spa18 and the other primer in the pair annealing to pUC19 and the erm resistance gene.

Indirect immunofluorescence assays of C3 binding to streptococci. Patterns of C3 binding to the surface of type 18 streptococci and the mutant strains were determined by confocal immunofluorescence microscopy by methods previously described (11). Briefly, bacteria grown to mid-log phase were incubated in either fresh human serum or plasma, washed, and then incubated with fluorescein-labeled goat anti-human C3 (Cappel). Photographs were made using a Nikon PCM2000 confocal laser scanning microscope in the Integrated Microscopy Center at the University of Memphis.

GenBank accession number. The previous GenBank entry (accession no. AFO86813) containing the partial sequence of spa has been updated to include the complete sequence.

FIG. 2. PCR analysis confirming insertional inactivation of spa by pUC19ermspa. Chromosomal DNA extracted from 18-282 (lane a), M18Ω (lane b), 18Spa− (lane c), and 18M18Vspa− (lane d) was subjected to PCR analysis. The annealing locations of primers F1, F2, F3, R1, R2, and R3 are shown (diagram is not to scale). Primer pair F1/R1 amplified a single product from the uninterrupted 3’ end of spa in all four strains. Primer pairs F2/R2 and F3/R3 annealed to both spa and pUC19, resulting in PCR products of appropriate size only with 18Spa− and 18M18Vspa−. ss, signal sequence; erm, erythromycin resistance gene.
tion process. There was no reaction of anti-rM18 with the extracts from M18\(\Omega\) and M18Spa\(\Omega\) (lanes b and d). Anti-rSpa reacted with two closely related proteins extracted from 18-282 and M18\(\Omega\) (lanes e and f). The largest of these two proteins had a molecular mass of \(\approx 59\) kDa, which is consistent with the mass of the native Spa protein predicted from the amino acid sequence. There was no reaction of anti-rSpa with the extracts from 18Spa\(\Omega\) and M18Spa\(\Omega\) (lanes c and g), and 18Spa\(\Omega\) reacted with antisera against rM18 (lanes a to d) and rSpa (lanes e to h).

Relative contributions of Spa and M18 in resistance to opsonization and phagocytosis. To determine the role of Spa in resistance to phagocytosis, we performed in vitro opsonophagocytosis assays using 18-282, M18\(\Omega\), 18Spa\(\Omega\), and the double mutant 18Spa\(\Omega\) (Table 1). In nonimmune blood containing preimmune rabbit serum, the parent 18-282, M18\(\Omega\), and 18Spa\(\Omega\) were equally resistant to phagocytosis, whereas the double mutant 18Spa\(\Omega\) was not resistant to phagocytosis. The organisms were also rotated in blood in the presence of either anti-rSpa or anti-rM18 to determine whether each strain was opsonized by the appropriate antiserum predicted by the phenotype. Parent strain 18-282 and mutant M18\(\Omega\) were opsonized by anti-rSpa, while mutant 18Spa\(\Omega\) was not (Table 1). Anti-rM18 opsonized only the parent and 18Spa\(\Omega\) strains, not the M18\(\Omega\) strain.

These results were confirmed by quantitating the growth of the organisms after rotation in whole, human blood containing either preimmune or immune rabbit serum (Table 2). After a 3-h rotation, the parent strain grew to 9.1 generations, while the M18\(\Omega\) and 18Spa\(\Omega\) mutants grew to 9.8 and 7.4 generations, respectively. The double mutant grew to only 4.5 generations. Again the bactericidal effect of anti-rM18 or anti-rSpa was consistent with the surface phenotype expected for each strain (Table 2). Taken together, these results show that Spa and M18 contribute to the ability of type 18 streptococci to resist phagocytosis in human blood. In addition, antibodies against either of these surface proteins are opsonic and able to promote phagocytic killing. When both Spa and M18 are inactivated, the organisms grow considerably less well in nonimmune blood, demonstrating that both surface proteins play a role in survival in blood in vitro of type 18 streptococci.

Mouse virulence studies. To further define the role of Spa as a virulence determinant, mouse virulence studies were performed by determining the \(LD_{50}\) following intraperitoneal injection of 10-fold-increasing inocula of each strain. The \(LD_{50}\) of 18-282 was \(2.8 \times 10^5\), that of M18\(\Omega\) was \(1.1 \times 10^6\), that of M18Spa\(\Omega\) was \(4.4 \times 10^5\), and that of M18Spa\(\Omega\) was \(1.1 \times 10^5\). Organisms were recovered from the spleens of mice that did not survive the challenge infection. M18\(\Omega\) retained its resistance to kanamycin, the selective marker associated with insertional inactivation of the \(emm\) gene (10), while 18Spa\(\Omega\) remained resistant to erythromycin and 18Spa\(\Omega\) continued to be resistant to both antibiotics. This demonstrated that the mutant phenotypes were stable in vivo. These results, taken together with those of the opsonization and bactericidal assays, indicate that the expression of Spa contributes to the virulence of type 18 streptococci.

C3 binding to the surface of the M18 parent and mutant strains. Indirect immunofluorescence assays were performed to assess the patterns of C3 deposition on the M18 parent and the M18-negative and Spa-negative mutants (Fig. 4). As shown in a previous study (11), the parent strain 18-282 incubated in nonimmune human plasma bound C3 mostly in a linear pattern (Fig. 4A), suggesting that the complement was binding to new cross walls. The M-negative (Fig. 4B) and Spa-negative (Fig. 4C) mutants similarly bound C3 as linear deposits. C3 deposition on the double mutant, however, was observed mostly in a circumferential pattern (Fig. 4D). Identical results were obtained when the bacterial strains were incubated in nonimmune human serum (data not shown). These data, taken together with the results of in vitro phagocytosis assays, indicate that both Spa and M protein confer upon type 18 streptococci the ability to resist opsonization by complement in nonimmune blood.

Human immune response to Spa. The human immune response to Spa was assessed using sera from patients known to have an antecedent streptococcal infection based on a documented history of acute rheumatic fever (Table 3). ELISAs were performed using a recombinant 212-residue NH\(_2\)-terminal fragment as the solid-phase antigen. Of the 20 different sera examined, 10 demonstrated measurable levels of antibody directed against Spa. Antibodies against various recombinant

![Image](http://iai.asm.org/Downloaded from http://iai.asm.org/)

**FIG. 3.** Western blot analysis of stationary-phase whole cell extracts of 18-282 (lanes a and c), M18\(\Omega\) (lanes b and f), 18Spa\(\Omega\) (lanes c and g), and 18Spa\(\Omega\) (lanes d and h) reacted with antisera against rM18 (lanes a to d) and rSpa (lanes e to h).

**TABLE 2.** Bactericidal activity of anti-rM18 and anti-rSpa against 18-282, M18\(\Omega\), and Spa-negative mutants

<table>
<thead>
<tr>
<th>Antiserum</th>
<th>CFU surviving 3-h rotation*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preimmune</td>
<td>5,280 (9.1)</td>
</tr>
<tr>
<td>Anti-rM18</td>
<td>10</td>
</tr>
<tr>
<td>Anti-rSpa</td>
<td>700</td>
</tr>
</tbody>
</table>

*Indirect bactericidal assays were performed by adding a standard inoculum of streptococci to 0.1 ml of test antiserum and incubating the mixture for 15 min at 37°C; 0.35 ml of whole, heparinized, nonimmune human blood was then added and rotated end-over-end for 3 h. Aliquots of 0.1 ml were then added to melted sheep’s blood agar, and pour plates were prepared. Viable organisms were quantitated after overnight incubation at 37°C. Numbers in brackets are inocula (CFU); number in parentheses are generations of growth following the 3-h rotation. The experiment was performed three times with similar results. The data presented are from one representative experiment.
N-terminal M protein fragments were assayed in parallel, and there was no obvious correlation between the presence of Spa antibodies and antibodies against particular type-specific M peptides (Table 3).

**Tissue cross-reactivity of anti-rSpa.** Studies have shown that several streptococcal antigens, including M protein, contain epitopes that evoke antibodies that cross-react with myocardium (8), renal glomeruli (15), cartilage (1), and brain (3). To assess whether anti-rSpa cross-reacted with human tissues, indirect immunofluorescence assays were performed on frozen sections of human myocardium, kidney, basal ganglia, and cerebral cortex. None of these assays demonstrated evidence of cross-reactivity between anti-rSpa and human tissues.

**DISCUSSION**

The central role of M protein in the pathogenesis of GAS infection is well established. In most cases, bacteria expressing M protein are able to resist opsonization and phagocytosis when incubated in nonimmune blood, while M-negative strains are readily killed (16). Several groups have investigated the properties of M protein that confer resistance to phagocytosis. These properties include the ability of M proteins to bind plasma fibrinogen, which coats the bacterial surface and blocks activation of the alternate complement pathway (25). In addition, M protein and fibrinogen bind factor H, a potent regulator of the complement cascade, which prevents the generation of C3b (12).

The ability to confer resistance to phagocytosis has become part of the functional definition of an M protein. However, recent studies have demonstrated that GAS have evolved other antiphagocytic properties that vary from one serotype to another and augment the ability of the bacteria to evade the host immune response. These other determinants of resistance to phagocytosis include surface proteins encoded by the *mrp* and *enn* genes that bind IgG and IgA, which may mask the surface of the organism and prevent nonspecific immune recognition (17). Studies in which both the *mrp* and *enn* genes were insertionally inactivated clearly demonstrated that the expression of both proteins contributes to resistance to phagocytosis (21). All group A streptococci also express C5a peptidase, which specifically inactivates the potent chemoattractant C5a and reduces the influx of polymorphonuclear leukocytes (6). The gene encoding for C5a peptidase (*scpA*) as well as *enn, mrp*, and *enn* are all members of a genomic regulon under the influence of the upstream positive regulator *mga* (5). Another important virulence factor is the hyaluronate capsule, which for some serotypes appears to be the major determinant of resistance to phagocytosis (11, 18).

In studies comparing the mechanisms of resistance to phagocytosis of type 24 and type 18 streptococci, it was clearly shown that the type 24 organisms were almost completely dependent on M protein expression and its binding of fibrinogen for growth in blood, while type 18 streptococci bound fibrinogen in much lower quantities and were more dependent on capsule expression for complete resistance to phagocytosis (11). The role of M protein in type 18 streptococcal infections was further examined in experiments where the *ennm18* gene was
insertionally inactivated, rendering the resultant mutant, M18I1, phenotypically M protein negative (10). Interestingly, the parent strain and the M18I1 mutant were equally resistant to phagocytic killing in nonimmune blood, and both strains demonstrated comparable virulence in mice after intraperitoneal challenge infections. Treatment of the organisms with hyaluronidase to remove the capsule resulted in a significant but incomplete reduction in growth of both organisms in whole blood. Thus, type 18 streptococci express a virulence determinant in addition to M protein and hyaluronate capsule.

In this study, we have shown that the presence of Spa on the surface of type 18 streptococci contributes to virulence and resistance to opsonophagocytosis. This was achieved by insertionally inactivating the native spa gene in both the parent M18 and mutant M18I1 strains. The expression of M protein and/or Spa was associated with an antiphagocytic phenotype, while the double mutant lacking both determinants was readily phagocytosed. Patterns of C3 opsonization of each strain were consistent with the results of phagocytosis assays, indicating that M protein and Spa both prevent deposition of C3 on the mature cell wall in either nonimmune human plasma or serum. The results of the phagocytosis and opsonization assays were confirmed by showing that the double mutant grew less well in nonimmune blood and demonstrated reduced virulence in a mouse model of infection compared to that of the parent or the single mutant expressing either M protein or Spa. Interestingly, the double mutant was not totally avirulent in mice, the single mutant expressing either M protein or Spa. Alternatively, the ancestral gene may have been acquired by horizontal gene transfer from another organism or from the environment.

Spa fulfills the functional definition of an M protein in that it is both a virulence determinant and protective antigen. Structurally, it shares some features with M proteins, including a signal peptide, a conserved gram-positive cell wall spanning sequence with an LPSTGE motif, and a predicted alpha-helical conformation throughout most of its length. Unlike M protein, however, Spa does not contain a proline-rich C terminus or N-terminal tandem repeats. The similarity between spa and the emm gene of S. equi was a very interesting finding. This group C streptococcus is an equine pathogen that causes a highly contagious upper respiratory tract infection known as streptococcal pharyngitis. The emm gene of S. equi encodes an M-like protein that is both a major virulence determinant and protective antigen of the organism (24). Although the C-terminal half of Spa is almost identical to the same region of the S. equi M protein, it is important to note that there is little similarity between the NH2-terminal halves of the mature proteins. This invites the speculation that the spa gene could have been acquired from S. equi (or vice versa) and has evolved under immunological and other environmental pressures into a unique surface protein. Alternatively, the ancestral gene may have been acquired by S. equi and group A streptococci from another organism and mutations or recombination occurred under the influence of different environmental pressures.

A critical question to address regarding the potential suitability of Spa as a vaccine candidate is its immunogenic potential and cross-reactivities. In this study, high-titer antibodies were generated in rabbits against a recombinant peptide copying the NH2-terminal half of Spa. In addition, this recombinant peptide was used as a solid-phase antigen to screen mice sera tested, indicating that the protein is expressed during infection and is immunogenic. Importantly, antibodies against the recombinant NH2-terminal half of Spa do not cross-react.
with human tissues. Since our previous studies have shown that antibodies against this fragment of Spa opsonize some heterologous serotypes of group A streptococci (10), studies are in progress to determine which serotypes may express conserved epitopes of this potentially important surface antigen.

ACKNOWLEDGMENTS

This work was supported by Public Health Service grant AI-10085 from the National Institute of Allergy and Infectious Diseases (J.B.D.), Merit Review funds from the Department of Veterans Affairs (J.B.D., D.L.H., and H.S.C.), and research funds from ID Biomedical Corporation. We appreciate the expert technical assistance of Sharon Frase, University of Memphis Integrated Microscopy Center, in obtaining images of C3 binding to streptococci using the confocal laser scanning microscope.

REFERENCES


Editor: V. J. DiRita