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Evolution of *sfbI* Encoding Streptococcal Fibronectin-Binding Protein I: Horizontal Genetic Transfer and Gene Mosaic Structure

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Streptococcal fibronectin-binding protein is an important virulence factor involved in colonization and invasion of epithelial cells and tissues by *Streptococcus pyogenes*. In order to investigate the mechanisms involved in the evolution of *sfbI*, the *sfbI* genes from 54 strains were sequenced. Thirty-four distinct alleles were identified. Three principal mechanisms appear to have been involved in the evolution of *sfbI*. The amino-terminal aromatic amino acid-rich domain is the most variable region and is apparently generated by intergenic recombination of horizontally acquired DNA cassettes, resulting in a genetic mosaic in this region. Two distinct and divergent sequence types that shared only 61 to 70% identity were identified in the central proline-rich region, while variation at the 3' end of the gene is due to deletion or duplication of defined repeat units. Potential antigenic and functional variabilities in SfbI imply significant selective pressure in vivo with direct implications for the microbial pathogenesis of *S. pyogenes*.

Streptococcal fibronectin-binding protein I (SfbI), also known as protein F1, is probably the most extensively characterized fibronectin-binding protein in group A streptococcus (GAS; *Streptococcus pyogenes*). Identified simultaneously in two independent laboratories (15, 39), SfbI is a membrane-anchored surface protein shown to be a major adhesin of GAS (13, 15, 28, 29, 39, 42). In addition, SfbI binds to fibrinogen via a domain in the amino terminus (20) and human immunoglobulin (IgG) via the carboxy-terminal fibronectin-binding repeat domain (25). SfbI is protective in mice (14, 34) and a mucosal adjuvant (26). According to various reports, it is present in 64 to 82% of strains (13, 22, 28). Beside its ability to mediate adherence to host cells via binding to fibronectin as a bridging molecule, SfbI is a potent invasin of epithelial cells (16, 27). While efficient adhesion is mediated through the fibronectin-binding repeat region, internalization into host cells is governed by the fibronectin-binding spacer region in a cooperative manner (41). Recent work demonstrates that SfbI-induced integrin clustering initiates caveola recruitment and subsequent caveola-mediated uptake of the pathogen into epithelial and endothelial cells (33). The interaction between the fibronectin-binding repeat region and the N-terminal F1 modules of fibronectin has been elegantly resolved by defining the structure of the complex that is formed (35). Another important and recently identified function of SfbI is the ability to recruit collagen via prebound fibronectin. SfbI thereby mediates bacterial aggregation, colonization of the collagen matrix, and phagocyte evasion (9). SfbI, like many membrane-associated streptococcal proteins, has a repetitive structure and exhibits decreasing variability from the amino terminus to the carboxy

terminus (40). The protein is synthesized as a propeptide with a signal sequence at the amino terminus of the protein which is subsequently cleaved to form the mature protein. The amino terminus of the mature protein contains a domain rich in aromatic amino acids, adjacent to which is a proline-rich domain. The proline-rich domain is made up of two regions: the proline-rich repeats and the proline-rich upstream region encompassing the region previously described as proline-rich repeats 1 and 2 (40). Toward the carboxy terminus lie two functional fibronectin-binding domains: the fibronectin-binding repeat domain and a second fibronectin-binding domain which lies upstream of the fibronectin-binding repeats and which has been designated spacer 2 (40) or the upstream fibronectin-binding domain (36). These domains are highly conserved and display variability only in the number of fibronectin-binding repeats, which may range from one to six (28, 42). Variable numbers of proline-rich repeats have also been reported; however, no correlation between variation in this domain and the numbers of fibronectin-binding repeats has been found (42). The carboxy terminus consists of a wall-associated region, a membrane anchor motif (LPATGD), and a membrane-associated region rich in hydrophobic residues, all of which are typical of surface proteins of gram-positive cocci (11).

Several groups have described the arrangement of the genomic region containing the gene encoding SfbI (6, 30). This locus has been designated the fibronectin-collagen-T antigen (FCT) region (6) since it may contain genes encoding the fibronectin-binding protein SfbI and/or protein F2, a collagen-binding protein (Cpa), and the T antigen. Incorporation of sequences from the recently completed M3 (4), M5 (http://www.sanger.ac.uk/Projects/S_pyogenes), and M18 (37) genomes reveals a complex gene arrangement in this region which is indicative of a zone of frequent genetic rearrangement.

Mosaic alleles are made up of distinct DNA cassettes of different phylogenetic origins which have recombined to produce

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novel chimeric genes. While gene mosaic structures for genes encoding streptokinase (19), hyaluronidase (24), streptococcal superantigen (32), and a novel collagen-like protein called SclA (31) have been reported in GAS, the classic example in GAS are members of the M-protein family encoded by genes within the Vir (or *mga*) regulon (43). Horizontal transfer and intergenic recombination have played major roles in the evolution of the *emm*-like family, contributing to both sequence variation and variation in the overall architecture of the Vir regulon (44). Since different members of the M-protein family exhibit different protein-binding capabilities, this rearrangement may have direct ramifications for the virulence and tropisms of the various strains. This is supported by the observation that different architectural arrangements (*emm* pattern types) have been associated with preferences for specific tissue sites (5).

Although evaluation of the accumulation of mutations is a valuable tool for assessment of strain relatedness, transfer of whole cassettes of DNA between strains complicates such analyses. Multilocus sequence typing with several presumably selection-neutral housekeeping genes has been used to determine the extent of intergenic recombination in bacterial genomes. The lack of congruence between maximum-likelihood trees for each of the "neutral" genes in GAS was indicative of very high rates of recombination, and it was concluded that this was sufficient to "obliterate the phylogenetic signal in [individual] gene trees" (10).

Recombination at distal sites such as the Vir regulon and the FCT region is most likely independent. Evidence to support this theory includes the fact that T types do not always conform to M types, with several T types reported for isolates of the same M type and vice versa (2, 17). In addition, while the distribution of *sfbl* within M types (28) or Vir types (VTs) (13) is generally consistent, there is evidence of heterogeneity of *sfbl* species within M types, as the number of fibronectin-binding repeats detected in M8 and M28 serotype strains is variable (28). Significant heterogeneity in *sfbl* sequences encoding the amino-terminal half of the aromatic amino acid-rich domain from M4, M12, M15, and M18 strains has also been reported (20).

The aim of this study was to examine precisely the genetic variability in the entire *sfbl* in order to determine possible mechanisms involved in the evolution of this important streptococcal adhesin and invasin. Additionally, we have examined whether particular *sfbl* sequence types (STs) are associated with the clinical source of the isolates.

(Part of this work was presented at the XVth Lancefield International Symposium on Streptococci and Streptococcal Disease, October 2002, Goa, India [R. Towers, M. J. Walker, and G. S. Chhatwal, Lancefield Int. Symp. Streptococci Streptococcal Dis. abstr. O5.3, 2002].)

MATERIALS AND METHODS

Bacterial strains. Isolates were obtained from the Menzies School of Health Research streptococcal collection. This collection contains isolates from patients of the Royal Darwin Hospital and the surrounding Aboriginal communities. In addition, an M13 GAS reference strain from the United Kingdom (Cathy13) and the homologous *sfblII* strain (strain 75401) from Germany were included (22, 38). Background information regarding clinical details, VTs (12), and *emm* STs (3), when available, are included in Table 1. All streptococcal strains were grown on agar plates containing 5% sheep blood (Sigma) at 37°C overnight.

PCR. The template for PCR was prepared by using the InstaGene matrix (Bio-Rad) according to the instructions of the manufacturer, except that due to the small size of the streptococcal colonies, 10 to 20 streptococcal colonies were used. The sequencing template was amplified with primers *sfbl*-F1 and *sfbl*-R4. Each 50- μ l PCR mixture contained 5 μ l of InstaGene preparation. The PCR parameters used were an initial denaturing step at 94°C for 2 min and then 35 cycles of denaturation at 94°C for 1 min, annealing at 49°C for 1 min, and extension at 72°C for 2 min 30 s, with a final extension step of 72°C for 5 min.

Screening for *sfbl* and *rofA*. Many of the data on the distribution of *sfbl* in the GAS strains used in this study have been published elsewhere (8, 13). For the purposes of this study all strains were retested for the presence of *sfbl* by PCR, and the results were compared with previous results. PCR screening was performed with primers *sfbl*-F1 and *sfbl*-R3. The number of fibronectin-binding repeats was determined with primers *sfbl*-F4 and *sfbl*-R3. Isolates were also screened for the presence of *rofA* with primers *rofA*-F and *rofA*-R. The arrangement of *rofA*, relative to that of *sfbl*, was determined with primers *rofA*-R and *sfbl*-R6. All primer sequences are shown in Table 2, while their positions relative to those of the respective genes are shown in Fig. 1A.

DNA sequence analysis. The PCR product was used as the template for sequencing reactions. This template was generated with primers *sfbl*-F1 and *sfbl*-R4 or *sfbl*-R3. A total of 5 μ l of the PCR product was visualized in an ethidium bromide-stained 1% agarose gel to determine its purity and relative concentration. The remainder was purified with QIAquick columns (Qiagen), eluted in 30 μ l of sterile distilled water, and stored at -20°C. The primers used for the sequence reactions are shown in Table 2.

Sequence alignment was performed with a range of DNASTAR programs. Applied Biosystems electropherograms were aligned to generate DNA contigs by using SeqManII, and then DNA contigs were translated to amino acid sequences with the MegAlign program and were subsequently aligned by the Clustal V method. The alignment was then modified by eye before being converted back to the DNA sequence alignment.

Nucleotide sequence accession numbers. The nucleotide sequences for the 54 *sfbl* genes can be found in the GenBank database under accession numbers AJ347791 to AJ347844.

RESULTS

Screening for *sfbl* and *rofA*. The presence of *sfbl* was confirmed by PCR for 54 GAS isolates. Representative PCR results are shown in Fig. 1B. PCR was also used to determine the number of fibronectin-binding repeats as an initial indicator of variation (Fig. 1C). In addition, the relative distribution and arrangement of *rofA* with respect to those of *sfbl* were established by PCR. All strains were tested for the presence of *rofA* with primers *rofA*-F and *rofA*-R (30) (Fig. 1D), while primers *rofA*-R and *sfbl*-R6 were used to detect adjacent *rofA* and *sfbl* genes (Fig. 1E). *rofA* was present in all 54 strains and was situated immediately upstream of *sfbl*, but in an orientation opposite that shown in Fig. 1A.

Sequence analysis of *sfbl*. The entire *sfbl* gene was amplified by PCR for use as the sequencing template from most strains by using primers *sfbl*-F1 and *sfbl*-R4. Initially, the *sfbl* PCR products from 20 strains were sequenced in their entirety in both directions. It was determined from these data that the 3' end of *sfbl* was highly conserved, apart from the various numbers of proline-rich and fibronectin-binding repeats. As the number of fibronectin-binding repeats could be readily determined by PCR (Fig. 1C) and these results were consistent with preliminary DNA sequence data, no attempt was made to sequence the fibronectin-binding repeat domains for the additional 34 strains examined in this study, and conclusions as to the relatedness of the genes were drawn from the sequence and PCR results combined.

The nucleotide sequences were aligned with published *sfbl* sequences, including those encoding full-length SfbI (GenBank accession no. X67947) (2); protein F1 (GenBank acces-

TABLE 1. Summary of *sfbl* STs

<i>sfbl</i> ST	Strain	Clinical source	Disease ^a	VT	<i>emm</i> ST	Aro type	Pro type ^b	No. of Pro rpts ^c	No. of Fn rpts ^d
1	NS678	Throat	N	33.2	ND ^e	1	A1	2	1
1	NS704	Throat	N	33.2	ND	1	A1	2	1
2	NS1045	Skin	N	89	<i>emm60</i>	1	A1	1	2
2	NS1053	Blood	I	89	<i>emm60</i>	1	A1	1	2
3	NS1036	Skin	N	33.1	<i>emm110</i>	1	A1	1	4
4	BSB19	Skin	N	6	<i>stbsb19</i>	2	A1	5	3
5	NS473	Skin	N	37.1	<i>stns554</i>	2	A1	1	4
6	NS539	Skin	N	1	<i>emm22</i>	2	A1	3	4
7	NS35	Skin	N	22	<i>emm58</i>	2	A1	2	4
7	NS351	Skin	N	22	<i>emm58</i>	2	A1	2	4
7	NS474	Skin	N	22	<i>emm58</i>	2	A1	2	4
7	NS687	Blood	I	22	<i>emm58</i>	2	A1	2	4
8	BL16	Skin	N	25	<i>emm85</i>	2	A2	4	1
9^f	D471	Throat	N	ND	(M6)^g	3	A1	1	5
10	NS691	Skin	N	57	<i>emm69</i>	4	A1	1	4
10	NS930	Skin	I	57	<i>emm69</i>	4	A1	1	4
10	NS931	Blood	I	57	<i>emm69</i>	4	A1	1	4
11	NS16	Skin	N	37.1	<i>stns554</i>	5	B1	1	5
11	NS240	Blood	I	116	<i>st2904</i>	5	B1	1	5
12	NS190	Skin	I	19	<i>emm74</i>	5	B1	1	4
12	DRX5	Skin	N	30	ND	5	B1	1	4
13	NS210	Blood	I	34	<i>emm22</i>	5	B1	2	3
14	NS1120	Throat	N	105	ND	5	B3	5	3
15	NS179	Pus	I	7.2	<i>emm9.1</i>	6	B3	11	2
16	NS192	Blood	I	3.2	<i>emm100</i>	6	B2	4	4
17	CATHY13	Unknown	U	2.1	<i>emm13</i>	6	B1	4	1
17	NS297	Skin	N	3.1	<i>emm44/61</i>	6	B1	4	1
17	NS495	Skin	N	2.2	<i>emm90</i>	6	B1	4	1
17	NS730	Pus	I	2.2	<i>emm90</i>	6	B1	4	1
17	NS731	Pus	I	2.2	<i>emm90</i>	6	B1	4	1
17	NS732	Aspirate	I	2.2	<i>emm90</i>	6	B1	4	1
17	NS755	Blood	I	2.2	<i>emm90</i>	6	B1	4	1
17	NS989	Skin	N	2.2	<i>emm90</i>	6	B1	4	1
18	NS226	Blood	I	14.1	<i>emm42</i>	7	A1	5	4
18	NS244	Skin	N	14.1	<i>stns244</i>	7	A1	5	4
18	NS415	Skin	N	14.1	<i>stns244</i>	7	A1	5	4
19	NS14	Skin	N	96	<i>emm102</i>	7	A1	4	4
20	NS488	Throat	N	52	<i>emm12</i>	8	A2	5	4
20	NS880	Throat	N	52	<i>emm12</i>	8	A2	5	4
20	A735	Throat	N	ND	(M12)	8	A2	5	4
21	75401	Throat	N	ND	(M75)	9	B1	2	3
22	NS101	Blood	I	33.1	<i>emm110</i>	10	B1	1	3
22	NS516	Skin	N	33.1	<i>emm110</i>	10	B1	1	3
23	BSA39	Skin	N	7.1	<i>emm63</i>	10	B1	1	5
24	NS1132	Skin	N	104	<i>emm25</i>	11	B1	1	3
25	DRV1	Skin	N	18	<i>emm55</i>	12	B1	1	4
26	NS216	Blood	I	122	ND	12	B1	4	4
27	EF1949	Unknown	U	ND	(M15)	13	A2	5	4
28	NS424	Skin	N	61	<i>emm14</i>	13	A2	6	4
28	NS483	Skin	N	61	<i>emm14</i>	13	A2	6	4
28	NS501	Blood	I	61	<i>emm14</i>	13	A2	6	4
28	NS506	Skin	N	61	<i>emm14</i>	13	A2	6	4
29	NS534	Blood	I	37.1	<i>stns554</i>	13	A2	3	1
30	DRY5	Skin	N	37.1	<i>stns554</i>	13	A2	1	2
31	DSM2071	Unknown	N	ND	(M23)	14	A1	2	4
32	NS90	Blood	I	52	<i>stns90</i>	15	B1	1	4
33	NS195	Blood	I	120	ND	15	B1	1	5
34	NS1042	Skin	N	77	<i>stck401</i>	15	B1	1	2

^a Invasive or noninvasive disease: I, invasive; N, noninvasive; U, unknown.

^b Pro type, SfbI proline-rich region sequence subtype.

^c Pro rpts, "true" proline-rich repeats.

^d Fn rpts, fibronectin-binding repeats.

^e ND, not determined.

^f Boldface indicates information published previously.

^g M serotype, as stated in relevant publications.

TABLE 2. Primer sequences used in this study

Primer direction and primer	Sequence
Forward	
<i>sfbI</i> -F1	5'-GTCTTCTTGACAATAACGTGGTAAGCTC-3'
<i>sfbI</i> -F2	5'-GTAGCCTATGCTGCCGATGAGAAG-3'
<i>sfbI</i> -F3	5'-TTTGTACCAGAAAATCCCCCTAAACCTG-3'
<i>sfbI</i> -F4	5'-GACTTACCTATTGAAGATCCTCGTTATGAG-3'
<i>sfbI</i> -F5	5'-GCATGCGCGGGTGCTATCG-3'
<i>sfbI</i> -F7	5'-GTACAGAATATGTACAAGATAATCC-3'
<i>sfbI</i> -S1F	5'-GTGCTGAATATGTACCTGATAGTCC-3'
PCRPF1	5'-TATCAAATCTTCTAAGTGCTGAG-3'
<i>rofA</i> -F	5'-GCCAATAACTGAGGTAGC-3'
Reverse	
<i>sfbI</i> -R1	5'-CTCATAACGAGGATCTTCAATAGGTAAGTC-3'
<i>sfbI</i> -R2	5'-CACTCCTGGCTCTTTCGTATCTTCTG-3'
<i>sfbI</i> -R3	5'-GTATCTTCAACAATGGTCACTGTTTCACTG-3'
<i>sfbI</i> -R4	5'-CAGTTATCCACTATTACAGCATATTTGCGC-3'
<i>sfbI</i> -R5	5'-CGATAGCACCCGCGCATGC-3'
<i>sfbI</i> -R6	5'-GCAGCATAAGGCTACTTGCACAAAAC-3'
<i>sfbI</i> -R7	5'-GGATTATCTTGTACATATTTCTGTAC-3'
<i>sfbI</i> -R8	5'-CAGGTCTAGGGGATCTTCTGGTACAAA-3'
<i>sfbI</i> -S1R	5'-GGACTATCAGGTACATATTCAGCAC-3'
PCRPIR	5'-CTCAGCACTTAGAAGATTTTG-3'
<i>rofA</i> -R	5'-GGTTTGTCTTTTAGGT-3'

sion no. L10919) (20); protein F1.12 (GenBank accession no. AF447492) (6); protein F15 and 12 partial *sfbI* sequences (GenBank accession nos. AF009908 to AF009920) (20); and *gfbA*, an *sfbI* homolog from GGS (GenBank accession no. U131115) (21). The nucleotide sequence alignment can be found at GenBank alignment accession number ALIGN_000212 (data not shown).

Source of variation in *sfbI*. Figure 2 shows an overall schematic of the variation observed in the *sfbI* sequences. As noted previously (40), variation in *sfbI* decreases in the 5'-to-3' direction. The variation at the 3' end of the gene is due to intragenic recombination, i.e., rearrangement of repetitive elements within genes through homologous recombination. The deletion or duplication of repeat units (narrower bars in Fig. 2) has resulted in a variable number of proline-rich and fibronectin-binding repeats. There is no relationship between the numbers of proline-rich repeats and the numbers of fibronectin-binding repeats, nor is there an association between the numbers of repeats and the upstream sequence variation. Therefore, it can be assumed that intragenic recombination is a frequent phenomenon within these regions.

Two distinct and divergent STs which exhibited 61 to 70% identity in pairwise alignments of the DNA sequences were identified within the central proline-rich region. These have been designated ProA and ProB. In addition, defined internal sequences have been duplicated or deleted, thereby generating two to three subtypes within each proline-rich region type (Fig. 2 and 3). In the case of ST ProB, in which up to two copies of the internal sequences were present, these repetitive structures were designated the ProB repeats.

The 5' end of *sfbI*, which encodes the aromatic amino acid-rich domain, is by far the most variable region of the gene. Fifteen distinct aromatic amino acid-rich domain sequences types (Aro types), designated Aro1 to Aro15, were identified. In pairwise alignments, a minimum 98.8% identity was seen within Aro types, while 50.3 to 97.6% identity was observed between types. While some of this variation is due to point

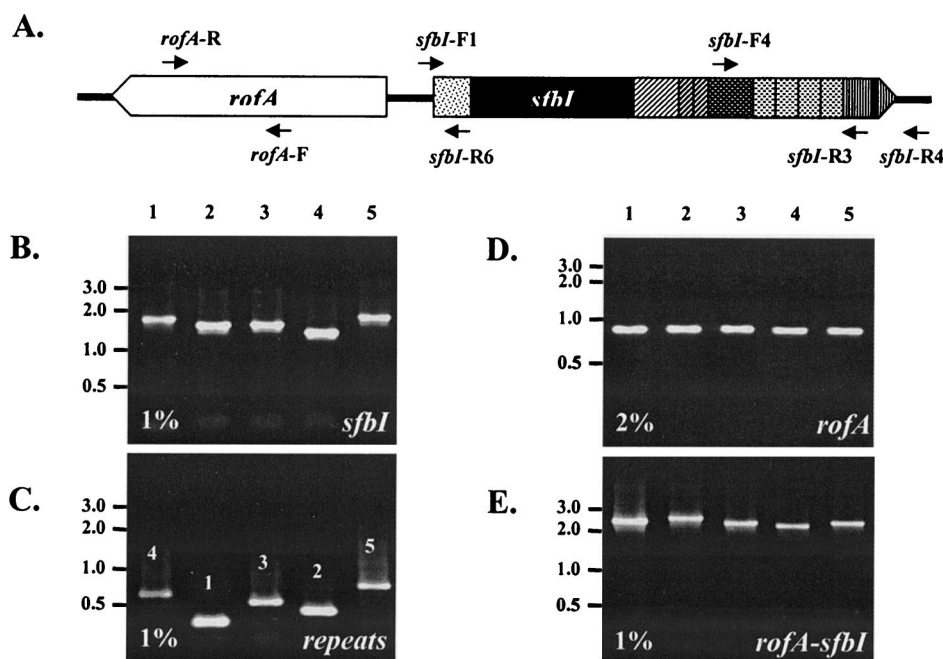


FIG. 1. (A) Positions of primers relative to the two genes. The shading of the SfbI domains is as described in the legend to Fig. 2. The schematic is not drawn to scale. (B to E) Ethidium bromide-stained 1 or 2% agarose gels showing the results of PCR screening for *sfbI* and *rofA* for five representative strains. (B) Screening for *sfbI* with primers *sfbI*-F1 and *sfbI*-R3; (C) determining the number of fibronectin-binding repeats in SfbI with primers *sfbI*-F4 and *sfbI*-R3; (D) screening for *rofA* with primers *rofA*-F and *rofA*-R; (E) determining arrangement of *rofA* relative to that of *sfbI* with primers *rofA*-R and *sfbI*-R6. DNA marker (GeneRuler DNA Ladder Mix, MBI Fermentas) sizes (in kilobases) are shown at the left. Lanes 1 to 5, GAS strains DSM2071, NS732, NS1132, DRY5, and BSA39, respectively.

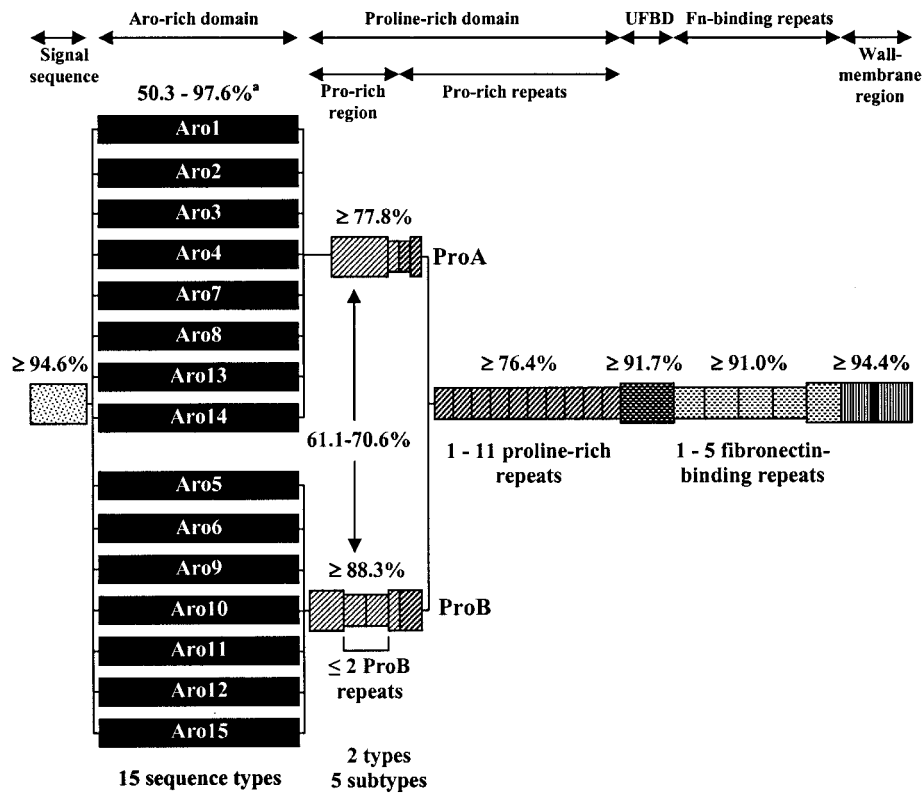


FIG. 2. Overall schematic of variation seen in SfbI. Decreasing variability of SfbI occurs from the amino-terminal aromatic amino acid-rich domain (solid black) through the proline-rich region (thin diagonal stripe) and to the proline-rich repeats (thick diagonal stripe) and the functional fibronectin (Fn)-binding domains (stippled). Narrower bars indicate the deletion or duplication of repeat units. The percent identities observed in pairwise DNA alignments is shown above each domain. Gaps introduced to aid alignment have been treated as single base changes. The diagram is not drawn to scale. UFBD, upstream fibronectin-binding domain.

mutations, which are easily identifiable as single nucleotide changes within otherwise conserved regions, the majority appear to be due to intergenic recombination. Similar to the *emm* gene, this intergenic DNA transfer has generated a gene mosaic structure in *sfbI*. The overall effects at the protein level can be seen in the amino acid sequence alignments of the various aromatic amino acid-rich domains (Fig. 4). Aro10 has been

used to illustrate how distinct DNA cassettes, flanked by highly conserved junction sequences, are shared between otherwise divergent STs. This type of mosaic allele can be generated only by horizontal gene transfer between strains.

Conserved structure of *sfbI*. Despite the high degree of variation observed, the overall structural properties of SfbI were conserved. Within the aromatic amino acid-rich domain, the

	Proline-rich region →		"True" proline-rich repeat →	
Consensus	<u>PDSPPKQGVFKESPKENPKIPELDQHEIPEVVPQEPSD---PVIPPVILDG</u>		<u>EEVPEVPSSELPALPPLMPELDGEVPE</u>	
Aro1 ProA1	.EN..R..T.....	.YG..T....SP...D.KH.D.NLE.TL...M.....Q....
Aro2 ProA1	.D.....DTS....	.H...T....SP...D.KH.D.NLE.TL...M.....S....
Aro2 ProA2	.D.....DTS....	.H...T....TP...D.KR.D.....S....
Aro3 ProA1	.D.....TS.....	.H...T....TP...D.KH.D.NLE.TL...M.....Q....
Aro4 ProA1	.KD.....T.....	.YG..T....SP...D.KH.DENLE.TL...M.....Q....
Aro7 ProA1	.D.....DTS....	.H...T....SP...D.KH.D.NLE.TL...M.....S....
Aro8 ProA2	.D.....DTS....	.H...T....TP...D.KR.D.....S....
Aro13 ProA2	.D.....DTS....	.H...T....TP...D.KR.D.....S....
Aro14 ProA1	.EN.....T.....	.HG..T....TP...G..R.NESLE.TL...M.....Q....
Aro5 ProB1	Q.N.....D.....
Aro5 ProB3	Q.N.....D.....	HEIPEVVPQEPSDPVIPAVIDG
Aro6 ProB1M.....D.....
Aro6 ProB2M.....D.....
Aro6 ProB3M.....V.D...R.....	EEVPEVVPQEPSNPVIPPVILDG
Aro9 ProB1M.....D.....Q....
Aro10 ProB1M.....D.....Q....
Aro11 ProB1M.....D...V.....Q....
Aro12 ProB1M.....D...R.....Q....
Aro15 ProB1M.....D.....Q....

FIG. 3. Alignment of amino acid sequences corresponding to the proline-rich domain of SfbI. The consensus sequence is shown at the top in boldface with underlining. The proline-rich region has two STs, ProA and ProB. Deletion or duplication events within these STs define the proline-rich region subtypes. Dots represent identical nucleotides relative to the consensus sequence, and hyphens indicate the positions of insertions found in particular isolates. The ProB repeats are indicated, and the first true proline-rich repeat is boxed.

distribution of aromatic amino acids remained relatively unchanged, and similarly, the distribution of proline residues within the proline-rich domain remained relatively unchanged. Over the entire length of the deduced amino acid sequence, hydrophobicity, average charge, and surface probability plots were generally conserved among the SfbI proteins from different strains. By restriction of comparisons to the aromatic amino acid- and proline-rich domains, the positively charged regions remained clustered in the aromatic amino acid-rich domain, despite sequence variation, with a few peaks in the proline-rich region before the negatively charged proline-rich repeats (Fig. 4). The periodicity of the repeat sequences is apparent in hydrophobicity, charge, and surface probability plots (data not shown). The theoretical pI of the putative SfbI proteins ranged between 4.11 and 4.81 (mean \pm standard deviation, 4.51 ± 0.13), indicating that SfbI would have a net negative charge at physiological pH.

Summary of *sfbl* STs. Thirty-five individual *sfbl* STs were identified on the basis of combinations of aromatic amino acid- and proline-rich domain STs and the numbers of proline-rich and fibronectin-binding repeats (Table 1). The strains within some VTs (VT2.2, VT14.1, VT22, VT33.1, VT33.2, VT52, VT57, and VT61) appeared to be clonal and were of the same *emm* and *sfbl* STs. M12 strain A735 (6) is also likely a clone of VT52, since all three strains are M12 and have identical *sfbl* genes. By contrast, VT37.1 was highly heterogeneous and included strains exhibiting *sfbl* STs 5, 11, 30, and 31. Conversely, other strains had virtually identical *sfbl* genes but belonged to different VTs; for example, *sfbl* ST18 included strains of VT2.1, VT2.2, and VT3.1. Some *sfbl* STs were identical except in the variable numbers of repeats; for example, *sfbl* ST30 and *sfbl* ST31 were identical except that the former has three proline-rich repeats and one fibronectin-binding repeat, while the latter has one and two repeats, respectively. This demonstrates that while the presence of *sfbl* is associated with VT, the species of the *sfbl* allele may be independent.

No correlation between the relative numbers of proline-rich and fibronectin-binding repeats was observed. There was also no apparent relationship between the numbers of repeats and aromatic amino acid- or proline-rich STs; for example, compare *sfbl* STs 4 to 8 within Aro2 (Table 1). Proline-rich domain STs were associated with particular aromatic amino acid-rich domain STs; however, several proline-rich domain sequence subtypes could be seen to be associated with a single aromatic amino acid-rich domain ST. For example, within Aro7, *sfbl* ST16, *sfbl* ST17, and *sfbl* ST18 exhibit proline-rich subtypes ProB3, ProB2, and ProB1, respectively.

Phylogenetic analysis of *sfbl*. The aromatic amino acid-rich domain was considered to be the greater indicator of relatedness, since the high degree of variability of this domain appeared to have been generated by many individual mutations, while the variability in the numbers of repeats was due to a smaller number of mutation events that occurred at a relatively high frequency but that involved large fragments of highly conserved DNA. This is probably best illustrated by the Aro6 group, the sequences of which are highly conserved in the aromatic amino acid-rich domain but which contain 0, 1, or 2 internal ProB repeats; 4 or 11 proline-rich repeats; and 1, 2, or 4 fibronectin-binding repeats. This suggests that the repetitive units are frequently shuffled and that this has little or no true

bearing on the phylogeny of the strain. Variation in *sfbl* has arisen through genetic exchange evolution of different alleles and is not independent. Therefore, a phylogenetic tree was not generated, since the principal assumption of such analysis is that the alleles have evolved independently.

Is the *sfbl* ST associated with the clinical source of the isolate? Information on the clinical sources of the isolates was available. Clonal distribution indicated that the skin was the principal reservoir for invasive disease isolates in the Northern Territory of Australia. Some *sfbl* STs were exclusively throat isolates (1, 9, 14, 21, 22); however, this association did not extend to Aro or proline-rich types. Further investigation revealed that all throat isolates were cultured from samples from nonindigenous people; therefore, this observation was probably due to the differences in the strains circulating in the two populations rather than to the tissue tropism that might be attributed to the SfbI variants. Overall, there was no common trend to indicate that particular or related STs were associated with the clinical sources of the isolates.

DISCUSSION

Components of both the Vir regulon and the FCT region have undergone frequent genetic rearrangement. These regions encode important virulence factors, principally adhesins with affinities for different human proteins such as fibronectin, plasminogen, collagen, fibrinogen, and immunoglobulins. Horizontal gene transfer has played a major role in the generation of variability within both regions, resulting in the mosaic alleles seen in *emm/enn* and *sfbl*.

The advantages of increased variability through horizontal gene transfer include the generation of antigenic variation and variable binding capabilities. As a mechanism of generating genetic diversity, horizontal gene transfer has the unique advantage of allowing incorporation of cassettes or domains which have already been checked for fitness in another genetic context. This bypasses the less efficient trial-and-error strategy and is particularly useful in altering the functional capabilities of a protein in a relatively short time frame.

The mosaic structure of the genes encoding the M-protein family has resulted in a divergent group of proteins exhibiting variable binding abilities and serotype-specific epitopes. The role that such variation plays in the pathogenesis of GAS is purely speculative; however, clear links have been shown between Vir regulon arrangements (*emm* patterns) and tissue tropism (5). We have shown that *sfbl* also has a gene mosaic structure especially apparent in the amino-terminal aromatic amino acid-rich domain and have identified 35 distinct *sfbl* STs. The biological function of the aromatic amino acid-rich domain is not known. To date, the reported binding functions of this domain include fibrinogen binding (20) and a putative carbohydrate-binding motif (40) which, from our data, appears to be highly conserved (Fig. 3A). The conserved distribution of aromatic amino acids within this domain, as well as the apparent intragenomic translocation of a highly homologous domain into Cpa (30), implies that the domain may have a significant function. In addition, the fact that the entire domain is present as a defined unit in Cpa implies that the function(s) and/or properties of the aromatic amino acid-rich domain require the whole domain. Despite the high degree of amino acid se-

quence variability, this domain remains positively charged. Since the proline-rich domain is predominantly negatively charged, we speculate that interaction between the two domains may be pivotal to the overall structure or folding of the native protein.

The other possibility is that variability has arisen in response to selective pressure from the immune system. Anti-M-protein antibodies are protective yet are serotype specific (23), as the protective epitopes reside in the variable amino terminus of the M protein (1, 7, 18). SfbI is a protective antigen when used in animal infection models (14); however, unlike M protein the protective epitope(s) resides in the conserved carboxy terminus and can provide protection from heterologous challenge (34). Natural infection in individuals from populations in which streptococcal infections are endemic also elicits significant levels of anti-SfbI IgG antibodies (13). Since these individuals are still infected, despite the presence of anti-SfbI IgG antibodies, it appears that anti-SfbI antibodies either are not protective in humans or are not present in significantly high enough concentrations to be effective. While the fibronectin-binding domain is protective and conserved, variability in the amino terminus may represent a mechanism of immune evasion, thereby directing the immune response away from the carboxy terminus.

Recombination within selection-neutral genes in GAS is common to the point that it masks phylogenetic signals (10). As a consequence, phylogenetic analyses of *sfbI* do not clarify evolutionary relationships between strains but clarify only the similarities between *sfbI* alleles. In looking at possible correlations between the *sfbI* STs and the clinical sources of the isolates, we did not find any associations. As with T-agglutination patterns, specific *sfbI* STs were consistently associated with *emm* STs; however, like T types, they were not restricted to singular *emm* STs. This seems to indicate that *emm* and *sfbI* are not closely linked. Given that the *tee* and *sfbI* genes both reside in the FCT region, it would be interesting to determine if T types and *sfbI* STs are more closely associated with each other than with *emm* STs. Such an analysis would require determination of the T types of the 54 GAS strains examined in this study.

The 34 individual *sfbI* STs identified in this study seem to have evolved through a combination of different evolutionary mechanisms working independently on distinct regions of the *sfbI* gene. The majority of divergence in the Aro domain appears to be due to intergenic recombination of horizontally transferred DNA cassettes, resulting in a genetic mosaic; two distinct and divergent STs are apparent in the central proline-rich region of *sfbI*; and the deletion or duplication of repeat units has resulted in a variable number of ProB, proline-rich, and fibronectin-binding repeats.

SfbI is an important virulence factor involved in colonization and internalization in fibronectin-rich epithelial tissues such as the upper respiratory mucosa and injured skin epithelium. Without clear associations between SfbI variants and function or tissue tropism, it is impossible to draw conclusions regarding the selective factors influencing these evolutionary mechanisms; however, the variability observed implies significant selective pressure on this locus in vivo. In light of the role played by intergenic transfer in the generation of variability in ligand-binding capacities in members of the M-protein family, the

implications for microbial pathogenesis of possible functional and antigenic variability in SfbI warrant further investigation.

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