A Structural Model for Binding of the Serine-Rich Repeat Adhesin GspB to Host Carbohydrate Receptors

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Abstract

GspB is a serine-rich repeat (SRR) adhesin of Streptococcus gordonii that mediates binding of this organism to human platelets via its interaction with sialyl-T antigen on the receptor GPIbα. This interaction appears to be a major virulence determinant in the pathogenesis of infective endocarditis. To address the mechanism by which GspB recognizes its carbohydrate ligand, we determined the high-resolution x-ray crystal structure of the GspB binding region (GspBbr), both alone and in complex with a disaccharide precursor to sialyl-T antigen. Analysis of the GspBbr structure revealed that it is comprised of three independently folded subdomains or modules: 1) an Ig-fold resembling a CnA domain from prokaryotic pathogens; 2) a second Ig-fold resembling the binding region of mammalian Siglecs; 3) a subdomain of unique fold. The disaccharide was found to bind in a pocket within the Siglec subdomain, but at a site distinct from that observed in mammalian Siglecs. Confirming the biological relevance of this binding pocket, we produced three isogenic variants of S. gordonii, each containing a single point mutation of a residue lining this binding pocket. These variants have reduced binding to carbohydrates of GPIbα. Further examination of purified GspBbr-R484E showed reduced binding to sialyl-T antigen while S. gordonii harboring this mutation did not efficiently bind platelets and showed a significant reduction in virulence, as measured by an animal model of endocarditis. Analysis of other SRR proteins revealed that the predicted binding regions of these adhesins also had a modular organization, with those known to bind carbohydrate receptors having modules homologous to the Siglec and Unique subdomains of GspBbr. This suggests that the binding specificity of the SRR family of adhesins is determined by the type and organization of discrete modules within the binding domains, which may affect the tropism of organisms for different tissues.

Introduction

The serine-rich repeat (SRR) glycoproteins of Gram-positive bacteria are an expanding family of microbial adhesins and virulence factors [1–6]. These proteins consist of a distinctive signal sequence and export-targeting region at the N-terminus, a short SRR region (~50–170 amino acids), a ligand binding region, a second lengthy SRR region (~400–4000 amino acids), and a cell wall anchoring motif at the C-terminus (Fig. 1) [1,2,4,6–9]. The binding regions of the SRR glycoproteins contain significant sequence variation, which appears to account for their broad range of binding targets, including platelet membrane and salivary
Author Summary

The binding of bacteria to human platelets is thought to be important for development of infective endocarditis, a life-threatening infection of the cardiovascular system. *Streptococcus gordonii* is a leading cause of endocarditis. This pathogen uses a protein called GspB to attach to carbohydrates on human platelets. While this binding interaction appears to be mediated by a specific, contiguous domain within GspB, little is known about the molecular details of the interaction between GspB and the carbohydrate receptors on its human host. We therefore determined the crystal structure of the region of GspB that binds to platelet carbohydrates, both alone and in complex with a synthetic carbohydrate receptor. Using this structure as a guide, we were able to produce three strains of *S. gordonii* that lacked the ability to bind to platelet carbohydrates. One of these isogenic variants was studied more in-depth and lacked the ability to bind to human platelets in vitro and was reduced in virulence when tested in vivo. These studies provide the first structural information detailing the molecular interactions between any serine-rich repeat adhesin and its host receptor, and identify how different, related adhesins may have evolved different specificities for host receptors.

glycoproteins [1,4,9–11], endothelial cells [12], epithelial cells [13], erythrocytes [14,15], and keratins [16,17]. Expression of SRR proteins has been associated with increased virulence in several animal models of infection, including endocarditis [6,18], meningitis [12], pneumonia [5], and bacteremia [7,19]. A number of bacterial surface components have been shown to mediate platelet binding in vitro, either by interacting directly with receptors on the platelet membrane, or via bridging molecules, such as fibrinogen [20–29]. The contribution of these interactions to virulence, however, has been assessed for relatively few of these adhesins. Previous studies have focused on the molecular basis for the SRR adhesin mediated binding of gram-positive bacteria to human platelets, and the role of this process in the pathogenesis of infective endocarditis. This interaction appears to be important for the attachment of blood-borne bacteria to platelets on the surface of damaged cardiac valves, thereby initiating infection. In addition, the subsequent deposition of platelets onto the infected endocardium may be due in part to bacterium-platelet binding, resulting in the formation macroscopic vegetations, which are the hallmark lesions of this disease [30]. Three of the SRR proteins (GspB of *Streptococcus gordonii* strain M99, Hsa of *S. gordonii* strain Challis, and SrpA of *Streptococcus sanguinis* strain SK36) bind human platelets through their interaction with glycopaladin, which is the carbohydrate-rich extracellular portion of the platelet membrane glycoprotein GP Ibα [9,11]. While the specific carbohydrate receptor for SrpA has not yet been identified, GspB and Hsa recognize sialylated trisaccharides [1,11,31,32]. Dot blot assays using immobilized carbohydrates have demonstrated that GspB has high fidelity for sialyl-T antigen (i.e. NeuAcα2–3Galβ[1–3]GalNAc), one of the major carbohydrates on GP Ibα [11], while Hsa binds to glycopaladin via either sialyl-T antigen or sialylactose (Neu5Acα2–6Galβ[1–4]Glc) [31,32]. Binding of these SRR adhesins to platelets is a high affinity process, with the interaction between GspB and glycopaladin having a KD of 2.38 × 10⁻⁸ M [11] and appears to be a major factor in the pathogenesis of infective endocarditis, since the loss of GspB or Hsa expression results in a marked reduction in virulence [33].

Structural information can enhance the understanding of the determinants of binding specificity. Here, we report the high-resolution crystal structure of GspBBR, both alone and in complex with the disaccharide α-2,3-sialyl (1-thioethyl)galactose, a precursor to synthetically-produced sialyl-T antigen. From these structures, we identified that a subdomain of GspBBR resembling mammalian sialic acid binding proteins binds to α-2,3-sialyl (1-thioethyl)galactose. Site directed mutagenesis and in vivo studies in a rat model of infective endocarditis verified that this carbohydrate binding site within GspBBR mediates binding of *S. gordonii* strain M99 to sialyl-T antigen, the host carbohydrates of the platelet.
membrane receptor GPIbα, and intact platelets, and that this interaction is important for virulence. Our analysis of the structure further identified that GspBBR contains an unusually modular fold, prompting us to re-analyze the sequences of the SRR family of adhesins. We determined that other structurally uncharacterized SRR adhesins also contain modules within their binding regions, suggesting that particular subdomains may be included, removed, or interchanged, manifesting in the broad range of binding partners observed in the family.

Results

Overall structure of GspBBR

We determined the structure of GspBBR to 1.4 Å resolution using the method of Multiwavelength Anomalous Dispersion from a single Dy³⁺ derivative (Fig. 2, Table 1, Table 2). GspBBR folds into an elongated rod, with dimensions of ~130 Å × 30 Å × 30 Å. This rod is comprised of three apparently independently-folded subdomains arranged in a linear fashion, like beads on a string. The secondary structure of each of the three subdomains is predominated by β-strands. Interestingly, the first two subdomains are organized around core folds that resemble those found within the eukaryotic immunoglobulin (Ig) superfamily (Fig. 3). Ig-folds have previously been identified in prokaryotic proteins [34–41].

Several topology variants of Ig-folds have been characterized. A structural homology search using the EMBL Dalilite server [42] identified that the N-terminal subdomain of GspBBR contains a folding topology, strand inserts, and inter-sheet angle reminiscent of the DE-variant of the Ig-fold [36] (Fig. 3A–C). This Ig-fold topology is found in prokaryotic proteins and was first identified within the A-region of the Staphylococcus aureus CNA protein [35]. CNA belongs to the family of microbial surface components recognizing adhesive matrix molecules (MSCRAMMs) of Gram-positive pathogens [34–36]. Given its structural similarity, this N-terminal subdomain of GspBBR will be termed the CnaA subdomain. The highest structural similarity between the CnaA subdomain of GspBBR and any other structurally characterized protein is to the C-terminal subdomain of the binding region of SRR adhesin Fap1 from Streptococcus parasanguinis (Fap1NR-β) [43]. The RMS deviation of the structural alignment between the CnaA subdomain of GspBBR and the Fap1NR-β subdomain is 2.5 Å for 200 Cα atoms.
Surprisingly, a structural homology search using EMBL Dalilite [42] identified that the second subdomain contained a topology and strand insert reminiscent of the V-set Ig fold adopted by eukaryotic sialic acid binding immunoglobulin-like lectins [44–46] (Siglec; Fig. 3D–F). However, the C′ and C′′ strands normally inserted into the V-set Ig fold are instead replaced by a long loop inserted at the same location (Fig. 3D–F). The RMS deviation of the structural alignment between the second subdomain of GspB BB and Siglec-5 is 2.9 Å for 210 Cα atoms despite only 6% sequence identity. Like GspB, Siglec5 bind carbohydrate receptors. Accordingly, the second domain is termed the Siglec subdomain. This subdomain of GspB BB contained electron density consistent with a cation-binding site (see Supporting Text S1, Fig. S1, Supporting Protocol S1).

This was rather unexpected since Siglec5 themselves do not bind cations in the carbohydrate-binding domain. The seven-coordinate number suggests that under physiological conditions, this should be a Ca2+ binding site. We explored the role of Ca2+ and other cations for the binding of S. gordonii strain M99 to glyocalcin. However, metal depletion, metal substitution, and site directed mutagenesis of the residues coordinating the ion were all consistent with the cation not being essential for carbohydrate receptor recognition (data not shown).

While still predominated by β-strands, the C-terminal subdomain of GspB BB does not contain an Ig-fold. In fact, a structural homology search using DaliLite [42] did not identify any proteins of known structure with significant similarity to this C-terminal subdomain (Fig. 2D). As a result, it will be termed the Unique subdomain.

**Identification of the receptor binding site**

We sought to identify the details of the interaction between GspB BB and the host receptor sialyl-T antigen. Free sialyl-T antigen is a rare reagent that is not commercially available and is challenging to synthesize. Therefore, we developed a 4-step synthesis for α2,3-sialyl (1-thioethyl)galactose (NeuAcα2,3(1–3)Galβ) which is the disaccharide truncation of and a synthetic precursor to sialyl-T antigen (NeuAcα2–3Galβ1–3GalNAc) (see Supporting Protocol S1; Fig. S2) which is the disaccharide truncation of and a synthetic precursor to sialyl-T antigen (NeuAcα2–3Galβ1–3GalNAc). To confirm that α2,3-sialyl (1-thioethyl)galactose binds to GspB BB, we assessed the ability of this disaccharide to inhibit the binding of S. gordonii to glyocalcin. In the presence of 44 mM α2,3-sialyl (1-thioethyl)galactose, binding of S. gordonii to glyocalcin was reduced by 90% (Fig. 4A). Given the structural similarity to the native host receptor, this strongly suggests that α2,3-sialyl (1-thioethyl)galactose competes directly for the sialyl-T antigen binding site.

We determined the co-crystal structure of α2,3-sialyl (1-thioethyl)galactose in complex with GspB BB. New electron density consistent with bound disaccharide (Fig. 4B) was apparent within a defined pocket (Fig. 4C) located at the N-terminus of the C strand and C-terminus of the F strand in the Siglec domain (Fig. 3F) of the Siglec subdomain (Fig. 4D). While the similarity of the fold of the Siglec subdomain to mammalian Siglecs might predict a similar binding pocket, both the overall location of the α2,3-sialyl (1-thioethyl)galactose binding site on the domain and the specific contacts between GspB BB to the carbohydrate differ from that of structurally characterized mammalian Siglecs [44–46] (Fig. 5). In fact, in GspB BB a helix is positioned in the location where carbohydrates bind to Siglec5, precluding the use of a structurally similar binding site. Instead, the local secondary structure surrounding the α2,3-sialyl (1-thioethyl)galactose binding site resembles a β-grasp domain, which is a motif that commonly binds the α2,3-linkage of sialic acid based multivalent carbohydrates.

**Table 1.** X-ray data collection and refinement statistics.

| PDB Accession Number | 3QC5 | 3QC6 | 3QO1 |
|----------------------|------|------|------|
| Space group          | P2₁,2₁,2₁ | P2₁,2₁,2₁ | P2₁  |
| Wavelength (Å)       | 1.49756 | 1.03034 | 1.07806 |
| Beamline             | LS-CAT ID-21-F | LS-CAT ID-21-B | LS-CAT ID-21-D |
| Resolution (Å)       | 50.1-1.40 | 50.1-1.90 | 50.1-1.90 |
| Unit-cell a,b,c (Å)   | 33.7, 96.8, 100.2 | 33.3, 86.7, 117.9 | 69.9, 34.0, 83.4 |
| No. of measured reflections | 223296 | 84235 | 97430 |
| No. of unique reflections | 60816 | 26193 | 29986 |
| Multiplicity          | 3.7(2.0) | 3.2(2.9) | 3.2(2.6) |
| Imean/σ(I)           | 19.8(2.9) | 16.3(3.9) | 17.4(4.8) |
| Completeness (%)      | 92.7(54.1) | 93.8(92.4) | 96.3(93.0) |
| R_{work} (%)          | 4.1(24.4) | 6.8(35.3) | 8.9(36.5) |
| R_{free} (%)          | 14.2/17.2 | 21.5/26.1 | 17.8/21.4 |
| No. of reflections in test set | 2543 | 1265 | 1477 |
| No. of protein atoms  | 352 | 351 | 359 |
| No. of water atoms    | 400 | 231 | 212 |
| RMS deviation bonds (Å) | 0.012 | 0.013 | 0.12 |
| RMS deviation angles  | 1.38 | 1.36 | 1.30 |

1Numbers in parentheses indicate values for the highest resolution bin.
| Peak | Inflection | Remote |
|------|------------|--------|
| Space group | P2₁,2₁,2₁ | P2₁,2₁,2₁ | P2₁,2₁,2₁ |
| Wavelength (Å) | 1.59083 | 1.59122 | 1.02463 |
| Beamline | SSRL 9-2 | SSRL 9-2 | SSRL 9-2 |
| Resolution (Å) | 50.1-1.90 | 50.1-1.90 | 50.1-1.90 |
| Unit-cell a,b,c (Å) | 35.5, 98.5, 99.1 | 34.5, 98.5, 99.1 | 34.5, 98.5, 99.0 |
| No. of measured reflections | 203951 | 206064 | 431338 |
| No. of unique reflections | 23918 | 23929 | 48691 |
| Multiplicity | 8.5(8.1) | 8.6(8.1) | 8.9(8.3) |
| Imean/σ(I) | 31.2(19.3) | 31.2(18.4) | 27.7(16.2) |
| Completeness (%) | 98.0(96.2) | 98.0(95.6) | 97.2(99.6) |
| R_{work} (%) | 8.7(17.9) | 8.0(17.7) | 6.1(35.9) |
| Phasing Power² | 4.1 | 3.6 | 2.1 |

²Phasing power is defined as the mean value of the heavy atom structure factor divided by the lack of closure error.

The overall Figure of Merit was 0.83. doi:10.1371/journal.ppat.1002112.t002

**Table 2.** Derivative data collection and phasing statistics.
Carbohydrates [47–49]. In GspB BR, the backbone carboxyls of Ala506 and Arg449, the N\textsubscript{g}1 and N\textsubscript{g}2 of Arg484, the backbone amide nitrogen of Thr483 and the side chain hydroxyl of Tyr443 make direct contacts to the \(\alpha\)-2,3-sialyl (1-thioethyl)galactose disaccharide, while the side chain hydroxyl of Tyr485 makes water-mediated contacts (Fig. 4B, Table 3).

A comparison of the structure of GspB BR determined with and without the disaccharide did not show significant structural changes localized within the binding pocket, which suggests that it is pre-formed. Thus, while the physiologically-relevant receptor trisaccharide sialyl-T antigen is a rare reagent, we can qualitatively suggest its binding location by modeling the third carbohydrate onto the first two carbohydrate binding sites, with the third carbohydrate of sialyl-T antigen extending toward the Unique subdomain (Fig. S3).

While disaccharide binding to GspB BR did not appear to induce obvious changes in conformations of side chains within the binding pocket, the interdomain angle between the CnaA and Siglec subdomains unexpectedly straightened by 40° (Fig. 6; Video S1). The change in orientation between the CnaA and Siglec subdomains involves rotation around a single hinge consisting of residues Lys398-Asp399-Thr400, where the side chain Asp399 is one of the ion coordinating residues.

**Functional relevance of the carbohydrate binding site**

To confirm that this crystallographically-identified \(\alpha\)-2,3-sialyl (1-thioethyl)galactose binding site is indeed important for GspB-

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Figure 3. Ig-fold topologies. A. Topology diagram of a C-set Ig-fold. B. Topology diagram of the DE-variant of a C-set Ig-fold identified in MSCRAMMs [36]. C. Topology diagram of the GspB BR CnaA subdomain. D. Topology diagram of a V-set Ig-fold. E. Topology diagram of a Siglec [45] showing the binding location of the sialylated carbohydrate in pink. F. Topology diagram of the GspB BR Siglec subdomain showing the binding location of the sialyl-T antigen in pink.

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mediated binding to GPIbα, we generated four isogenic variants of strain M99 with point mutations within the Siglec subdomain of GspB (Table 4). Three of these variants contained mutations within the crystallographically-identified binding site (Y443F, R484E and Y485F), while a control mutation (E401A) was located within the Siglec domain, but away from the receptor binding site. Importantly, none of the point mutations affected surface expression of GspB (Fig. 7A). Each of these isogenic variants had an 86% or greater decrease in binding to GPIbα in vitro, as compared with the parent strain ($p < 0.001$) (Fig. 7B). This decrease is comparable to that of the gspB null mutant. In contrast, binding by the E401A variant was not significantly different from that of M99 ($p = 0.44$).

We then selected the R484E mutation for more detailed study. The purified GST-GspBBR fusion protein harboring the R484E substitution exhibited a marked decrease in binding to biotinylated sialyl-T antigen (Fig. 7C). The R484E substitution also resulted in reduced binding to human platelets in vitro as assessed by quantifying the amount of input inoculum bound to fixed platelets (Fig. 7D). We next visualized platelets by Differential Interference Contrast (DIC) and fluorescence microscopy to quantify the number of platelets with surface-bound bacteria. In the presence of the DAPI-labeled PS846 (M99ΔgspB) or the PS2116 (M99 gspB<sup>−</sup>) strain, substantially fewer platelets had bacteria bound as compared to platelets in the presence of wild-type <i>S. gordonii</i> strain M99 (Fig. 8, Table 3). It should be noted that while the platelet numbers were normalized when placed onto the cover slip, during the experiment the wild type bacteria seemed to form microscopic aggregates with the platelets (data not shown) but this was not observed when platelets were mixed with strains PS2116 or PS846. This confirms the importance of this residue for carbohydrate binding.

Impact of Siglec-mediated binding on virulence

It has previously been demonstrated that loss of GspB expression results in a marked decrease in virulence, as measured by a rat model of infective endocarditis [18]. To assess whether binding via the Siglec subdomain to host carbohydrate receptors contributes to virulence, we examined the impact of the R484E substitution on the ability of <i>S. gordonii</i> strain M99 to produce endocardial infection. We first compared the relative virulence of M99 with strain PS2116, (M99 gspB<sup>−</sup>). Catheterized rats were simultaneously infected intravenously with $2 \times 10^5$ CFU of M99 and PS2116. After 72 h, the animals were sacrificed and the relative levels of bacteria within tissues determined. Animals co-infected with M99 and PS2116 (which carries the <i>spec</i> resistance marker just upstream of <i>gspB</i>) served as controls. When assessed at the above time-point, animals co-infected with M99 and PS2116 had significantly reduced densities of the mutant strain within vegetation ($\text{mean} \pm \text{S.D.} = 6.81 \pm 1.70 \log_{10} \text{CU/g. veg}$) as compared with parental strain M99 ($7.47 \pm 1.69 \log_{10} \text{CFU/g. veg}; P < 0.02$). Loss of sialyl-T antigen binding was also associated with
members adopt unrelated folds in their binding domains. Like GspBBR, Fap1NR appears to be composed of independently-folded subdomains; however its binding region only contains two modules whereas the binding region of GspBBR contains three. Intriguingly, Fap1NR contains a helical subdomain at its N-terminus (Fap1NR-α) that does not resemble any of the subdomains of GspBBR, and a CnaA subdomain at its C-terminus (Fap1NR-β) that resembles the N-terminal CnaA subdomain of GspBBR (GspBBR-C). This suggested to us that members of the SRR family could undergo reorganization of the modules within their host binding regions, with particular modules or combinations of modules conferring specific properties.

Accordingly, we re-analyzed the binding regions of selected members of the family using a new strategy, where we used BLAST [50] and ClustalW [51] to query the sequences of the binding regions of SRR adhesins with input sequences corresponding to subdomains of either GspBBR or Fap1NR, or of short regions (~200 amino acids) of sequences of structurally uncharacterized binding regions of SRR adhesins. These modified sequence comparisons strongly suggest that the binding regions of members of the SRR family have indeed evolved to contain modules (Fig. 9), and show several distinct groupings. For example, all three assessed SRR adhesins with carbohydrate binding partners contain the Siglec and Unique subdomains in tandem, strongly suggesting that the inclusion of these modules within the binding region confers lectin-like binding characteristics.

These focused sequence alignments additionally identify that a CnaA subdomain may be common in SRR adhesins, appearing in three of the eight binding regions that we assessed. As opposed to the combination of the Siglec and Unique subdomains, which are associated with carbohydrate binding, each adhesion containing a CnaA subdomain may have a different binding specificity (Fig. 9). Interestingly, in the SRR adhesins that we assessed, the CnaA subdomains were always found paired with sequence for another subdomain.

It is important to note that several of the binding regions of the SRR adhesins do not have significant sequence identity to any protein of known structure. For example, the binding region of \( S. agalactiae \) SRR1 (SRR1GBS) likely contains two subdomains, SRR1GBS-α at the N-terminus, which does not contain detectable sequence similarity to any protein of known fold, and SRR1GBS-β at the C-terminus, which has sequence similarity to MSCRAMMs and likely adopts a similar folding topology to CnaA. Likewise, no part of the binding regions of \( S. pneumoniae \) PsrP, \( S. aureus \) SraP, or \( S. epidermidis \) seSRR exhibited detectable sequence similarity to any known protein, but the latter two have high sequence similarity to each other.

**Discussion**

SRR adhesins have been identified in a variety of Gram-positive pathogens, and have been implicated as virulence factors in a wide spectrum of infections [5,6,12,18]. The diversity of these infections (e.g., endocarditis, meningitis, pneumonia) and the broad scope of

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**Table 3. Contacts at the receptor binding site.**

| Ligand Atom | Protein Atom | Distance (Å) |
|-------------|-------------|--------------|
| OAW         | Ala506 O    | 2.6          |
| NAO         | Ala506 O    | 2.5          |
| OAT         | HDH213*     | 2.5          |
| OAP         | HDH213*     | 3.1          |
| O2          | HDH199*     | 2.9          |
| O3          | Tyr443 OH   | 2.5          |
| SAC         | Thr483 N    | 2.6          |
| O5          | Arg484 N\_1| 3.1          |
| O6          | Arg484 N\_2| 2.7          |
|             | Arg449 O    | 2.9          |

*Water molecules make water mediated contacts to the protein and are not just solvating.

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their anatomic locations are consistent with the binding regions of the SRR proteins differing considerably in their selectivity. Indeed, although only a few ligands for this family of proteins have been identified, they range from carbohydrates (such as sialyl-T antigen) [32] to proteins (such as keratins) [16,17]. To date, however, the structural basis for this selectivity has been unknown.

In several members of the family (e.g. the SRR adhesins of S. gordonii and S. pneumoniae) the domain between the two SRR regions mediates binding. However, for many of these SRR adhesins (i.e. SraP of S. aureus), the host receptor has yet to be identified [6,43], and the delineation of the binding region is assumed based upon sequence comparisons within the family. Among the best characterized of the SRR adhesins is GspB from S. gordonii, which has demonstrated binding affinity for the sialyl-T antigen carbohydrate decorating platelet glycoprotein GPIb. Our crystal structure of the binding region, GspB\_BR, identified a modular organization with three subdomains (Fig. 2A), two of which are organized around Ig-folds. Proteins containing Ig-folds are commonly found within the mammalian immune system, where they exhibit a variety of functions; however, Ig-folds are not uncommon within pathogens, where they act exclusively as virulence factors. The first characterized bacterial protein to contain an Ig-fold was PapD, a chaperone for the assembly of pili in E. coli [37]. The Ig-fold has been observed in several other types of virulence factors, including components of pili (carbohydrate-binding fimbriae) [41,52], and in adhesins of varying specificity for host receptors, including MSCRAMMs [34–36] and invasins [38–40].

Co-crystallization of GspB\_RR with α-2,3-sialyl [1-thioethyl]galactose identified a specific binding site for this disaccharide within the Siglec subdomain (Fig. 4B–D), and the introduction of single point mutations within this region significantly reduced levels of binding of S. gordonii to glyocalcicin. Moreover, the R484E substitution in GspB reduced platelet binding by M99 (Fig. 7D, Fig. 8, Table 5), and had a marked reduction in binding of the GST-GspB\_RR fusion protein to sialyl-T antigen (Fig. 7C), confirming the importance of the Siglec subdomain for carbohydrate binding. These results indicate that this binding pocket within the Siglec subdomain is required for binding to sialyl-T antigen, and that the interaction of this GspB subdomain with sialyl-T antigen is important for binding to platelets.

When examined in a co-infection model of infective endocarditis, the isogenic variant of M99 expressing GspB-R484E (strain PS2116) was also significantly reduced in virulence (most notably a 78% reduction in bacterial levels within vegetations as compared with the parent strain). When PS2116 was tested with PS846 (M99∆gspB) in this model, the strains had comparable but reduced densities within target tissues (as compared with WT M99), indicating that they were similarly reduced in virulence. These findings indicate that the predominant property of GspB contributing to the pathogenesis of endocarditis is its interaction with sialyl-T antigen. In previous in vivo studies, co-infection with M99 and PS846 yielded more pronounced differences between the strains, as measured by densities of organisms within target tissues [18]. This may reflect subtle experimental differences, such as inoculum size, but could also be due to residual binding activity of the mutated GspB. Of note, PS2116 had low but detectable levels of platelet binding in vitro. Similarly, the GST-GspB\_RR-R484E had measurable levels of binding to sialyl-T antigen, as compared to GST alone. This residual binding seen in vitro, presumably due to the other key residues identified by crystallography, may account for the residual virulence of the PS2116 strain. Alternatively, although ligands for GspB other than sialyl-T antigen have not been identified, it is conceivable that GspB has other interactions with host receptors.

| Table 4. Strain names with descriptions. |

| Strain name | Description |
|-------------|-------------|
| M99         | S. gordonii strain M99 wild-type |
| PS2116      | S. gordonii strain M99 gspB\_RM4E |
| PS846       | S. gordonii strain M99ΔgspB |
| PS2161      | S’gspB:spec control strain |

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control residue not located near the binding pocket. C. Binding of sialyl-T antigen to purified GST-GspBBR wild-type and variant (R484E) proteins were immobilized in microtiter wells, and the binding of biotinylated sialyl-T antigen to each protein was detected by using peroxidase-conjugated streptavidin along with a chromogenic peroxidase substrate. Binding is expressed as the mean ± standard deviation (n = 3). The O represents purified GST-GspBBR wild-type, the x represents purified GST-GspBBR-R484E, the u represents purified GST-GspBBR-R484E, and the o represents purified GST-GspBBR-R484E. D. Binding of M99 and derivative strains to human platelets. Binding is expressed as the percent of input bacteria that remained bound to platelets after repeated washing of the wells (mean ± standard deviation).

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Figure 7. Amino acid substitutions in the carbohydrate binding site of GspBBR. A. Expression of GspB and variant proteins on the S. gordonii cell surface. Peptidoglycan-linked proteins were extracted from the bacterial cell surface and analyzed by western blotting using a polyclonal anti-GspA serum (upper panel). The nitrocellulose membrane was then re-probed using a polyclonal anti-GspB serum (lower panel) in order to assess the relative efficiency of protein extraction and protein loading (GspA is another LPXTG-linked protein that is expressed by S. gordonii M99). B. Binding of S. gordonii strain M99 and derivative strains to glycoscalicin immobilized in microtiter wells. Binding is expressed as the percent of input bacteria that remained bound to glycoscalicin after repeated washing of the wells (mean ± standard deviation). The labels of each column indicate the sequence of the protein expressed in S. gordonii, with M99 indicating wild type, ΔgspB indicating a strain where the gene encoding GspB has been deleted, and remaining columns indicating the amino acid substitution present in the protein. E401 is a site of GspB BR.

Figure 6, Video S1}

between the CnaA and Siglec subdomains (Fig. 6, Video S1). The interdomain hinge is centered at the ion binding site, making the interdomain angle change reminiscent of calcium dependent interdomain angle changes observed in C-cadherin, which also contains modules of Ig-fold topology [39]. There are several possibilities for the origin and role of this interdomain reorganization in GspBBR. It is first possible that GspBBR exhibits natural flexibility between these two domains, and that the difference in crystal contacts artifactually resulted in the straighter form of the protein being trapped in the co-structure of GspBBR with α-2,3-sialyl (1-thioethyl)galactose. Intriguingly, however, an interdomain straightening is also observed in small angle x-ray scattering (SAXS) of the binding region of SRR protein Fap1NR upon lowering the pH to 5.0, where Fap1NR has highest affinity for its host receptor [43]. Those experiments, performed in solution, are not limited by crystal contacts and suggest that interdomain straightening upon ligand binding could be a conserved feature of members of the SRR family. Physiological roles of interdomain straightening upon ligand binding include improved hydrodynamics in the cardiovascular system or oral cavity. The elongation upon ligand binding is perhaps more compellingly reminiscent of the interdomain straightening that occurs upon the binding of P-selectin and β3-integrin to their respective ligands [54]. These proteins have been suggested to form “catch bonds” in order to have increased affinity to their ligand upon the application of tensile force. Each of these possibilities for the observed straightening of GspBBR upon ligand binding is currently under investigation.

While both SiglecS and GspBBR bind to carbohydrate receptors, the structural details of disaccharide binding to this novel bacterial Siglec bear little resemblance to the binding of sialic acid moieties to characterized mammalian Siglecs [44–46] (Fig. 5). Indeed, comparison of the binding sites reveals that the secondary structure of GspBBR has a helix that is found at the same location where 3’ sialyllactose binds to Siglec-5 (Fig. 5A), thus eliminating the possibility of analogous carbohydrate binding. Instead, GspBBR appears to use a β-grasp domain to form the specific host receptor binding site, a domain found in another group of sialic acid binding proteins, staphylococcal superantigen-like proteins (SSLs) that forms a V-like cleft for carbohydrate binding [47–49]. Nevertheless, both mammalian Siglecs and GspBBR have binding pockets predominated by tyrosines and arginines. With a closer look at each binding site, in Siglecs, the sialic acid moiety of the binding partner makes a salt bridge with an arginine residue. This salt bridge is not present in the binding site of GspBBR despite the presence of R484 and its key role in binding; this arginine instead binds to the G6 hydroxyl and the pyranose oxygen of the 1-thioethylgalactose (Table 3).

Carbohydrate Recognition by GspB
Importantly, the structure of GspB BR provides insight into the binding repertoire of other SRR proteins, and specifically highlights which modules mediate binding to host carbohydrate receptors. Homologues of GspB containing regions with high sequence similarity to the Siglec and Unique subdomains (such as S. gordonii strain Challis Hsa and S. sanguinis SrpA) are also demonstrated lectins. Supporting this sequence analysis, our co-crystal structure of GspB BR with 2,3-sialyl(1-thioethyl)galactose demonstrates that the first two carbohydrates of host receptor bind within a pre-formed pocket on the Siglec subdomain. Using this experimental co-structure as a starting point, we could model binding of the sialyl-T antigen. This model is consistent with the third carbohydrate of this trisaccharide binding to an extended lobe on this pocket. This region of the binding pocket is still within the Siglec subdomain but extends toward the Unique subdomain (Fig. S3).

The structures of both GspB BR and Fap1 NR identified a CnaA-like subdomain within the binding region, and our sequence analysis (Fig. 9) additionally predicts a CnaA subdomain at the C-terminus of the binding region of S. agalactiae SRR1 (SRR1 GBS-B), an adhesin for keratin 4 [16]. As opposed to the Siglec subdomain, which contains an identifiable receptor binding site, the precise function of the CnaA subdomain is less clear. MSCRAMMs, such as CnaA from S. aureus, recognize peptides through the formation of a binding site between two Ig-like domains (Fig. S4). The bound peptide forms an additional strand that becomes a part of the Ig-fold. In contrast, GspB BR, Fap1 NR, and SRR1 BR each apparently contain a single CnaA subdomain amongst the modules in the binding region (Fig. 9). Our modeling suggests that binding of a peptide between two domains in a manner analogous to binding of a peptide to other MSCRAMMs is not feasible. Indeed, an additional strand found in the subdomain of GspB BR occupies the peptide binding site of MSCRAMMs. (Fig. S4, Fig. 3C strand A). Nevertheless, in SRR adhesins, the CnaA subdomain is always found together with at least one other subdomain, suggesting that the function may either require or be tuned by the presence of a second subdomain. Indeed, recent studies on the binding region of Fap1 support this hypothesis. The structures of Fap1 NR-α (helical subdomain) and Fap1 NR-β (CnaA subdomain) identified a region of surface-exposed hydrophobic residues on each domain that is predicted to be contiguous based upon SAXS. Fap1 normally mediates binding of this bacterium to the oral cavity, and mutagenesis of these hydrophobic residues abrogated binding to an in vitro tooth model consisting of saliva-coated hydroxylapatite [43].

Our sequence analysis additionally predicts that the binding regions of other SRR adhesins contain modules that are structurally distinct from those identified in either GspB BR or Fap1 NR. Two of these, S. aureus SraP BR and S. epidermidis seSRR BR, have high sequence identity to each other, but no detectable sequence similarity to any other SRR adhesin (Fig. 9). This strongly suggests that they bind a common, as yet unidentified, host receptor that is distinct from the carbohydrates and keratins recognized by other SRR adhesin family members. By comparison, neither S. pneumoniae PsrP BR, which has been demonstrated to bind keratin 10, nor the N-terminal subdomain of S. agalactiae SRR1 BR, which binds keratin 4, have detectable sequence similarity to any currently available sequence (Fig. 9). Given this analysis, it is clear that a more detailed understanding of the binding characteristics of the SRR adhesin family will require that for each module type, the binding partner should be identified, and the structure of the binding region should be determined both alone and in complex with the appropriate host receptor.
Table 5. Binding of three variants of *S. gordonii* to human platelets.

| Strain | % of Platelets with Bacteria Bound (Mean ± S.D.) |
|--------|-----------------------------------------------|
| M99    | 32.0±2.8                                       |
| PS2116 | 3.0±1.7*                                      |
| PS846  | 7.0±1.7*                                      |

*p = 0.01 as compared with M99.

Note: The difference in binding between the PS2116 and PS846 strains were not statistically significant.

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Materials and Methods

Ethics statement

All procedures involving rats were approved by the Los Angeles Biomedical Research Institute animal use and care committee, following the National Institutes of Health guidelines for animal housing and care. Platelets were collected using a protocol approved by the UCSF Committee on Human Research (H1193-25513-07) and by the VUMC Human Research Protection Program IRB Committee (110364).

Protein expression and purification

The DNA encoding residues 233–617 of GspB (GspBBR) was cloned into the pGEX vector containing an N-terminal glutathione S-transferase (GST) fusion tag as described [11]. This clone contains a single base pair change as compared to the deposited NCBI sequence that results in a serine at position 444 instead of an asparagine. The protein was expressed with the BL21 Gold (Stratagene) in Luria Broth Medium as described [55] and purified using a GST affinity column (GE Healthcare). The GST tag was removed from GspBBR using Factor Xa, and GspB BR was further purified using size exclusion chromatography on a Superdex 200 10/300 GL column (GE Healthcare) with buffer containing 20 mM Tris pH 7.4 [55].

Crystallization

All crystals of GspBBR were grown using the hanging drop vapor diffusion technique at 23°C [55] using 1 mM protein solution and 1 mM of reservoir solution equilibrated against 1 mM of the reservoir solution. Crystals of native GspBBR grew from two chemically distinct sets of conditions. The first set of conditions included GspBBR at a concentration of 6 mg/ml buffered in 20 mM Tris pH 7.4 and equilibrated against a reservoir solution containing 33% PEG 3350, 0.1 M HEPES pH 7.5 and 0.15 M KCl at 23°C. These crystals belonged to the orthorhombic space group *P2₁2₁2₁* with unit cell dimensions a = 69.9 Å, b = 94.0 Å, c = 100.2 Å with α = β = γ = 90°. Prior to data collection, crystals were cryo-protected in a solution containing all of the chemical components of each reservoir solution and 15% glycerol then flash-cooled in liquid nitrogen. The dataset used for refinement merged to 1.4 Å resolution.

The second type of native GspBBR crystals grew when the protein was equilibrated against a reservoir solution containing 25% polyethylene glycol 3350, 0.1 M HEPES pH 7.5, 0.15 M NH₄CH₃COO, and 10 mM spermidine and induced crystal growth from 10 mg/ml GspBBR buffered in 20 mM HEPES pH 7.5 at 18°C. These crystals also belonged to the orthorhombic space group *P2₁2₁2₁*, but had altered unit cell dimensions of a = 33.4 Å, b = 86.7 Å, c = 117.9 Å, α = β = γ = 90°. Prior to data collection, these crystals were flash cooled in liquid nitrogen without additional cryo-protectant. The best dataset from this crystal form merged to 2.0 Å resolution.

Cysts of GspBBR in complex with the α-2,3-sialyl (1-thioethyl)galactose disaccharide were grown using the hanging drop vapor diffusion method with 6 mg/ml GspBBR in buffer containing 1 mM α-2,3-sialyl (1-thioethyl)galactose and 20 mM Tris pH 7.4. The reservoir solution contained 8% PEG 3350, 7.5 mM CoCl₂, 7.5 mM NiCl₂, 7.5 mM MgCl₂, and 0.1 M HEPES pH 7.5. Crystals grew within two days and were cryo-protected in reservoir solution supplemented with 20% glycerol and flash-cooled in liquid nitrogen. The disaccharide co-crystals formed in the primitive monoclinic space group *P2₁* with unit cell dimensions a = 69.9 Å, b = 94.0 Å, c = 83.4 Å with α = γ = 90° and β = 99.5°. The best data from this crystal form merged to 1.9 Å resolution.

Data collection and processing

Crystals were assessed for diffraction quality at the Stanford Synchrotron Radiation Lightsource (SSRL) beamlines 9-2, 11-1, and 12-2 and the Life Sciences Collaborative Access Team (LS-CAT) beamlines ID-21-D/F/G. Datasets were collected using the beamlines, temperatures, wavelengths, and detectors listed in Table 1 and Table 2. All data were processed using the HKL2000 [56] and CCP4 [57] suites of programs.

Preparation of heavy atom derivatives and structure determination

A Dy³⁺ derivative was prepared by soaking pre-formed crystals of GspBBR in 1 mM DyCl₃ for three days. Data were collected at three wavelengths near the Dy³⁺ L₃ edge (Table 2). Dy³⁺ bound to a single site in the protein as determined using the SHELXD [58] subroutine in the program SHARP [59]. While HoCl₃ also successfully derivatized GspBBR, non-isomorphism between crys-

Table 6. Effect of Sialyl-T antigen binding on virulence in a rat model of infective endocarditis.

| Strain pairs | No. of animals | *S. gordonii* densities (Mean ± S.D. log₁₀ CFU/g tissue) |
|--------------|---------------|--------------------------------------------------------|
|              | Vegetations   | Kidneys       | Spleen       |
| M99 (wt)     | 7.47±1.69     | 3.72±0.72     | 3.77±0.66    |
| PS2116 (R484E) | 11            | 6.81±1.70*    | 3.40±0.86*   | 3.20±0.72** |
| PS846 (Agosp) | 5.76±2.15     | 2.72±1.32     | 3.32±1.10    |
| PS2116 (R484E) | 5.45±2.69     | 2.53±1.12     | 2.92±0.88    |

*p = 0.02;**p = 0.001 compared with M99.

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tals prevented the use of this second derivative in a traditional MIR calculation. The non-isomorphism was so severe that phases calculated in SHARP [59] only used data from a single DyCl$_3$ soaked crystal and did not include a native dataset for reference. This process resulted in reasonable phasing statistics and an overall figure of merit of 0.83 at 2.0 Å resolution (Table 2). Phases were improved by solvent flattening in DM [60] which produced electron density maps of high quality (Fig. 2B). Automated chain tracing was performed using PHENIX [61], which was able to trace residues 251–316 and 327–601, representing 94.7% of the model. This resulted in an initial R cryst of 23.8% and R free of 25.5%.

The lack of isomorphism between crystals prevented transfer of these initial coordinates to other data sets using a simple rigid body refinement. As a result, the model from the Dy$_3$ data set was transferred to the remaining data sets using the program PHASER [62] followed by rigid body refinement in CNS [63]. Each structure was subjected to alternate rounds of model building using the program COOT [64] and refinement using CNS [63] and REFMAC [65].

The coordinates for the α-2,3-sialyl (1-thioethyl)galactose were built using CCP4i Sketcher [66] and PRODRG [67]. Refinement statistics for all final models are listed in Table 1. Figures were created using PyMOL [68], and inter-domain rotations were determined using DynDom [69].

**Synthesis of the α-2,3-sialyl (1-thioethyl)galactose disaccharide**

Based upon work by Daníñéshey and coworkers [70], we developed a four-step synthesis for the sialyl-T antigen precursor, α-2,3-sialyl (1-thioethyl)galactose (see Supporting Protocol S1). The correct synthesis of the disaccharide was verified by NMR (Fig. S2). The α-2,3-sialyl (1-thioethyl)galactose was resuspended in water for all applications.

**Site-directed Mutagenesis**

Point mutations of gspB were introduced into the S. gordonii chromosome via a strategy that involved recombination by double cross-over between gspB codons 487–602 and a gene approxi-
mately 500 bp upstream. This approach ensured incorporation of only the intended mutation of *gspB* codons ranging from 399 to 485, and avoided possible imprecise recombination within the SSR regions. As a first step, the 5’ end of *gspB* (codons 1 to 486 along with 200 nts from a non-coding region upstream) was replaced with a chloramphenicol resistance cassette as follows. A 0.3 kb segment of a gene of unknown function upstream of *gspB* was amplified using PCR. The product was digested with XhoI and Clal and then cloned upstream of the *cat* gene in pC326 [71]. A segment spanning *gspB* codons 487 to 602 was then amplified using primers B487F and B602R, digested with SpeI and NotI, and cloned downstream of the *cat* gene. The resulting plasmid, pC326A5’S, B, was used to transform *S. gordonii* strain M99 as described [4]. One of the chloramphenicol-resistant transformants, (M99 Δ5’gspBR;cat), was selected for subsequent gene replacement.

A series of plasmids was then constructed to facilitate the replacement of the 5’ end of *gspB* in M99 Δ5’gspBR;cat. The XhoI-Clal fragment from pC326A5’S, B was cloned upstream of the *spec* gene in pS326 [72], and a 1 kb NsiI-SpeI fragment of *gspB* (spanning codons 1 to 296), was cloned downstream. A SpeI-NotI fragment spanning codons 296 to 602 was then cloned downstream of the NsiI-SpeI fragment. Point mutations in the resulting plasmid, pS326B602, were generated by a two-stage PCR procedure. In the first stage, primer 25F along with a reverse *gspB* primer, or the corresponding *gspB* forward primer along with primer B602R, were used to amplify the upstream or downstream segments, respectively. The two PCR products were combined for the second stage and then amplified using primers 25F and B602R. The PCR product was digested with SpeI and NotI and then used to replace the corresponding segment of pS326B602. The incorporation of only the intended change in any segment generated by PCR was confirmed by DNA sequence analysis. Plasmids were then used to transform M99 Δ5’gspBR;cat, resulting in a replacement of Δ5’gspBR;cat with a Δ5’gspBR::spec variant. As a control, the wild-type *gspB* sequence along with the *spec* cassette (pS326B602) was also crossed into the M99 Δ5’gspBR::spec chromosome (generating strain PS2161). Transformants were selected on spectinomycin and scored for the loss of chloramphenicol resistance. Expression of the variant GspB proteins on the bacterial cell surface was verified by western blotting as described [72].

**GspBBR** binding to biotinylated sialyl-T antigen

GST, GST-GspBRR and GST-GspBRR-R484E were purified from *E. coli* as described [55]. The purified proteins were diluted to 320 μM into DPBS, serial two-fold dilutions were made, and 50 μl of each dilution was added to wells of a 96-well microtiter plate. After incubating the plate overnight at 4°C, unbound proteins were removed by aspiration and wells were rinsed with 100 μl DPBS. Biotinylated sialyl-T antigen (sialyl-T antigen conjugated to biotin via a polycrylamide linker; GlycoTech Corporation) was diluted to 50 μg/ml in DPBS containing 1× Blocking Reagent (Roche), 50 μl was added to each well, and the plate was incubated for 2 h at RT with vigorous rocking. After removing unbound biotin-sialyl-T antigen, wells were rinsed three times with 100 μl DPBS, 50 μl of streptavidin-conjugated horseradish peroxidase (0.1 μg/ml in DPBS) was added to each well and the plate was incubated for 1 h at 23°C. The wells were washed twice with 100 μl DPBS, and then 200 μl of a solution of OPD (0.4 mg/ml citrate-phosphate buffer) was added to each well. The contents of the wells were mixed by gently vortexing the plate, and the absorbance at 450 nm was measured 20 min after the addition of the OPD substrate. Data were plotted as the means ± standard deviations, with n = 3.

**Binding of *S. gordonii* to glycolcalcin and platelet monolayers**

The binding of *S. gordonii* to immobilized glycolcalcin was performed as described previously [32]. In brief, strains were grown for 18 hr, washed twice with DPBS, sonicated briefly to disrupt aggregated cells, and then diluted to approximately 2×10^9 per ml. To determine whether Δ2,3 sialyl [1-thioethyl]galactose inhibited binding to glycolcalcin, the washed and sonicated bacteria were diluted into DPBS or DPBS containing 44 mM Δ2,3-sialyl [1-thioethyl]galactose, pH 7.5. The bacterial suspensions were then applied to wells of a microtiter plate that had been coated with glycolcalcin (1.25 μg/well). After 2 h at room temperature, the unbound bacteria were removed by aspiration. Wells were washed three times with DPBS, and the bound bacteria were released by trypsinization. The number of input and bound bacteria were determined by plating serial dilutions of the bacterial suspensions on sheep blood agar plates, and binding was expressed as the percent of the input bound to glycolcalcin. The binding of *S. gordonii* to immobilized human platelets was assessed as described previously [4]. Results of both assays are reported as the means ± standard deviations, with n = 6. Differences in binding were compared by the unpaired t-test.

**DIC and fluorescence microscopy**

*S. gordonii* strains M99 wild type, PS2116, and PS846 were grown in 5 mL of Todd Hewitt Broth at 37°C without shaking for 18 hours. Cells were then vortexed to resuspend bacteria and spun down at 4000× g for 10 min. The supernatant was removed and the bacteria washed twice with 5 ml of DPBS containing MgCl2 and CaCl2. The bacteria were then resuspended in 5 mL of DPBS and sonicated briefly to disrupt aggregated clumps. To 1 mL of each cell suspension, 500 nM 4',6-diamidino-2-phenylindole (DAPI) was added.

Platelets were fixed using 1.6% paraformaldehyde. 500 μL of platelets were mounted on poly-L-lysine coated cover slips (in a 6-well tray) which was spun at 400× g for 10 min in order to promote platelet adherence to the cover slips. Excess platelets were removed by washing with Tris Buffered Saline (TBS) and 1 mL of TBS was added to each well. 500 μL of the bacterial suspension was added to the platelets and the samples were rocked vigorously for 30 minutes at 23°C. Excess bacteria were then removed by washing three times with TBS. Each sample was mounted onto a slide for microscopy. Slides were imaged using Nikon TiE Inverted light microscope equipped with a Photometrics Coolsnap HQ CCD camera. Platelets were imaged with a 100×1.49NA objective using DIC optics and a DAPI filter cube. Image-J software was used to create the contrast and composite images.

**Rat model of infective endocarditis**

A competition model of infective endocarditis was produced in Sprague-Dawley female rats (250–300 g) as described previously [18]. In brief, the animals were anesthetized with ketamine (35 mg/kg) and xylazine (10 mg/kg). A sterile polyethylene catheter was surgically placed across the aortic valve of each animal, such that the tip was positioned in the left ventricle. Catheters were left in place throughout the study. Catheterized animals were then infected intravenously (IV) with an inoculum containing 2×10^8 CFU of both strains (i.e., a 1:1 mixture of i) *S. gordonii* strain M99 and strain PS2116 (5′gspB::spec, ii) M99 and strain PS2161 (5′gspB::spec control strain), or iii) PS2116 and strain PS846 (M99 ΔgspB;EVP3; CmKK) [18,73]. At 72 h post-infection, animals were sacrificed with thiopental (100 mg, intraperitoneally). Animals were included in the final analysis only if the catheters were correctly positioned across
the aortic valve at the time of sacrifice, and if macroscopic vegetations were seen. All cardiac vegetations, as well as samples of the kidneys and spleens were harvested, weighed, homogenized in saline, serially diluted, and plated onto Todd Hewitt agar (THA) (for the parental S. gordonii strain M99) and THA containing 100 µg/ml of spectinomycin (for strains PS2116 and PS2161) or chloramphenicol 5 mg/ml (for strain PS846) for quantitative culture, to determine the number of CFU/g of S. gordonii strains within tissues. After 48 h of incubation at 37°C, bacterial colonies were counted.

The number of bacteria within tissues was expressed as the log10 CFU per gram of tissue. Differences between means were compared for statistical significance by the paired t-test, using p≤0.05 as the threshold for significance. The data were also analyzed by calculating a "competition index," which was defined as the ratio of organisms in the inoculum. The mean of the log 10 normalized ratios was tested against the hypothesized 'no effect' mean value of 0, as described previously, using a paired t-test.

Accesion numbers
The coordinates and structure factors for S. gordonii strain M99 GspBBr have been deposited in the Research Collaboratory for Structural Bioinformatics Protein Data Bank with accession codes 3QC5 (native 1), 3QC6 (native 2, crystal form 2), and 3QD1 (α-2,3-sialyl [1-thioethyl]galactose bound).

Supporting Information
Figure S1 The cation binding site. Stereo view of the cation binding site superpositioned with 2m|Fo|−d|Fc| electron density calculated in REFMAC5 [63] and contoured to 1.5 σ. Carbons are shown in grey, oxygens are shown in red, nitrogens are shown in blue and the cation is shown as a purple sphere. Bond distances are shown in A.

TI-F

Figure S2 1H NMR, 400 MHz (CDCl3) spectrum of α-2,3-sialyl (1-thioethyl)galactose.

TI-F

Figure S3 Predicted binding of sialyl-T antigen to the receptor binding pocket. Using the co-structure of GspBBr with α-2,3-sialyl (1-thioethyl)galactose as a starting point, we developed a model for binding of sialyl-T antigen to the Siglec subdomain. Coordinates for sialyl-T antigen were prepared in Coot [64] by manually linking the 3-position of N-acetylgalactosamine to the 1-position of the galactose in α-2,3-sialyl (1-thioethyl) galactose and removing the thioethyl group. The position of the α-2,3-sialyl (1-thioethyl) galactose group was fixed to its position in the experimentally-determined structure and the galactosamine manually optimized to avoid steric clashes. In this model, all three sugars bind within a contiguous pre-formed pocket on the surface of the protein between the Siglec and Unique subdomains. The Siglec subdomain is colored yellow, the Unique subdomain is colored blue, and the modeled sialyl-T antigen is colored with green carbons.

TI-F

Figure S4 Binding of MSCRAMMs to fibrinogen receptors. The interdomain peptide binding observed in SdrG (PDB entry 1R17) uses a "dock, lock, and latch" mechanism of fibrinogen recognition [34] is unlikely to be used by SRR adhesins. This interdomain mechanism requires two adjacent Ig-fold domains that create a binding site; the fibrinogen peptide binds between these two domains. Modeling additional interdomain angles in GspBBr suggests that the short linker region between the CnaA and Siglec subdomains will prevent the interdomain angle from becoming sufficiently acute to allow the Siglec subdomain to act as a surrogate second subdomain. In addition, GspBBr contains an additional strand at the N-terminus of the CnaA subdomain (Fig. 3C strand A’) that occupies the peptide binding site of MSCRAMMs. Unbound SdrG is colored gray and bound SdrG is colored red with the fibrinogen peptide colored cyan.

TI-F

Video S1 Range of interdomain angles observed in crystals of GspBBr. The movie morphs between states observed in the crystal structures and was produced using LSQM AN [75] and PyMOL [68]. The most extended conformation was observed in the costructure of GspBBr with the α-2,3-sialyl [1-thioethyl]galactose disaccharide.

MP4

Test S1 Results and discussion regarding the cation binding site.

DOCX

Protocol S1 Identification of the cation binding site and synthesis of α-2,3-sialyl (1-thioethyl)galactose.

DOC

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Author Contributions
Conceived and designed the experiments: GC GAS MJT PMS TMI. Performed the experiments: TMP BAB YQX BJM VY KMO MJT. Analyzed the data: TMP BAB YQX BJM TMI NJW MJT. Contributed reagents/materials/analysis tools: YY. Wrote the paper: TMP BAB YQX BJM GAS MJT PMS TMI.

References
1. Takahashi Y, Konishi K, Cisar JO, Yoshikawa M (2002) Identification and characterization of hsa, the gene encoding the sialic acid-binding adhesin of Streptococcus gordonii DLI. Infect Immun 70: 1209–1218.
2. HandleY PS, Correia FF, Russell K, Rosan B, DiRienzo JM (2005) Association of a novel high molecular weight, serine-rich protein (SrpA) with fibril-mediated adhesion of the oral biofilm bacterium Streptococcus cristatus. Oral Microbiol Immunol 20: 131–140.
3. Sanchez CJ, Shivshankar P, Stol K, Trakhtenbroit S, Sullam PM, et al. (2010) The pneumococcal serine-rich repeat protein is an intra-species bacterial adhesin that promotes bacterial aggregation in vivo and in biofilms. PLoS Pathog 6(6): e100044. doi:10.1371/journal.ppat.100044.
4. Siiboo IR, Chambers HF, Sullam PM (2005) Role of SrpA, a Serine-Rich Surface Protein of Staphylococcus aureus, in binding to human platelets. Infect Immun 73: 2273–2280.
5. Seifert KN, Addison EE, Whiting AA, Bohanac JK, Crowley PJ, et al. (2006) A unique serine-rich repeat protein (Str-2) and novel surface antigen (epiln) associated with a virulent lineage of serotype III Streptococcus agalactiae. Microbiology 152: 1029–1040.
6. Wu H, Fives-Taylor PM (1999) Identification of dippeptide repeats and a cell wall sorting signal in the fibrin-associate adhesin, Fap1, of Streptococcus parasanguis. Mol Microbiol 34: 1070–1081.
7. Plummer C, Wu H, Kerrigan SW, Meade G, Cox D, et al. (2005) A serine-rich glycoprotein of Streptococcus sanguis mediates adhesion to platelets via GPIb. Br J Haematol 129: 101–109.
10. Takamatsu D, Bensing RA, Prakash B, Fisher SJ, Sullam PM (2006) Binding of the streptococcal surface glycoproteins GspB and Hsa to human salivary proteins. Infect Immun 74: 1933-1940.

11. Takamatsu D, Bensing BA, Cheng H, Jarvis GA, Siboo IR, et al. (2006) Binding of the Streptococcus gordonii surface glycoprotein GspB and Hsa to specific carbohydrate structures on platelet membrane glycosylprotein Iib alpha. Mol Microbiol 58: 389–392.

12. van Sorge NM, Quach D, Gurney MA, Sullam PM, Nizet V, et al. (2009) The group B streptococcal serine-repeat 1 glycoprotein mediates penetration of the blood-brown barrier. J Infect Dis 199: 1479–1487.

13. Mistou MY, Dramsi S, Brega SA, Poyart C, Trieu-Cuot P (2008) Structural organization of the Streptococcus gordonii DL1 sialic acid-binding adhesin. Microbiol Immunol 52: 37–49.

14. Takahashi Y, Sandberg AL, Ruhl S, Muller J, Cisar JO (1997) A specific cell surface antigen of Streptococcus gordonii is associated with bacteriophage hemagglutination and adhesion to alpha-2,3-linked sialic acid containing receptors. Infect Immum 65: 5042-5051.

15. Sameu E, Eikmans BJ, Reinscheid DJ, Borge F (2007) The surface protein Sra-1 of Streptococcus agalactiae binds human keratin 4 and promotes collagen fibrillogenesis to epitheial HEP-2 cells. Infect Immum 75: 3245–3251.

16. Shvihshanuk P, Sanchez C, Rose LF, Ohrulina CJ (2009) The Streptococcus pneumoniae adhesin PefP binds to Keratin 10 on lung epithelium. Mol Microbiol 73: 663-679.

17. Xiong YQ, Bensing BA, Bayer AS, Chambers HF, Sullam PM (2008) Role of the serine-rich surface glycoprotein GspB of Streptococcus gordonii in the pathogenesis of infective endocarditis. Microbiol Path 45: 279–301.

18. Obert C, Sublett J, Kausch D, Hinojosa E, Barton T, et al. (2006) Identification of a Candidate Streptococcus pneumoniae core genome and regions of diversity correlated with invasive pneumococcal disease. Infect Immum 74: 4776-4787.

19. Petersen HJ, Keane C, Jenkinson HF, Vickerman MM, Jesionowski A, et al. (2006) Identification of Streptococcus gordonii surface proteins SspA/SspB and Hsa in platelet activation. Infect Immun 74: 6707-6718.

20. Takamatsu D, Bensing BA, Prakobphol A, Fisher SJ, Sullam PM (2006) Binding of the Streptococcus gordonii surface glycoprotein GspB and Hsa to specific carbohydrate structures on platelet membrane glycosylprotein Iib alpha. Mol Microbiol 58: 389–392.

21. Takahashi Y, Takashima E, Shimazu K, Yagishita H, Aoba T, et al. (2006) Contribution of sialic acid-binding adhesin to pathogenesis of experimental endocarditis caused by Streptococcus gordonii. Infect Immum 74: 740–743.

22. Symersky J, Patti MJ, Carson M, House-Pompeo K, Teale M, et al. (1997) Structure of the collagen-binding domain from a Staphylococcus aureus adhesin. Nat Struct Biol 4: 833-838.

23. Deivanayagam CC, Wann ER, Chen W, Carson M, Rajashankar KR, et al. (2002) A novel variant of the immunoglobulin fold in surface adhesins of Staphylococcus aureus: crystal structure of the fibrinogen-binding MSCR4, clumping factor A. EMBO J 21: 6660–6672.

24. Holmgren A, Branden CI (1989) Crystal structure of chaferpeptide protein PapD from Pseudomonas fluorescens: an immunoglobulin fold in an alpha/beta sandwich. J Mol Biol 212: 249–251.

25. Cota E, Jones C, Simpson P, Alfred H, Anderson KL, et al. (2006) The solution structure of the invasive tip complex from Afa/Dr fibrils. Mol Microbiol 62: 356-366.

26. Hamborszina ZA, Brown MS, Isberg RR, Bjorkman PJ (1999) Crystal structure of invasion: a bacterial integrin-binding protein. Science 286: 291-295.

27. Jakubovics NS, Kerrigan SW, Nobbs AH, Stromberg N, van Dolleweerd CJ, et al. (2005) Functions of cell surface-anchored antigen I/II family and Hsa proteins. Infect Immun 73: 1933-1940.

28. Zhubravleva MA, Trandem K, Sun PD (2008) Structural implications of Siglec-5-mediated sialylglycans recognition. J Biol Chem 283: 4729-4736.

29. Atwill RJ, Innamurra A, Sharma RS, Kiso M, Crocker PR, et al. (2006) Siglec-7 undergoes a major conformational change when complexed with the alpha(2,8)- disialylganglioside GT1b. J Biol Chem 281: 32774–32783.

30. Ramboarina S, Garnett JA, Zhou M, Li Y, Peng Z, et al. (2010) Structural insights into sialic-acid binding of gramin-positive bacteria. J Biol Chem 285: 32496-32457.

31. May AP, Robinson RG, Vinson M, Crocker PR, Jones EY (1998) Crystal structure of the N-terminal domain of sialadhesin in complex with single sialylactose at 1.83 A resolution. Mol Cell 1: 719-728.

32. Symersky J, Patti MJ, Carson M, House-Pompeo K, Teale M, et al. (1997) Structure of the collagen-binding domain from a Staphylococcus aureus adhesin. Nat Struct Biol 4: 833-838.

33. Sempersky J, Patti MJ, Carson M, House-Pompeo K, Teale M, et al. (1997) Structure of the collagen-binding domain from a Staphylococcus aureus adhesin. Nat Struct Biol 4: 833-838.

34. Symersky J, Patti MJ, Carson M, House-Pompeo K, Teale M, et al. (1997) Structure of the collagen-binding domain from a Staphylococcus aureus adhesin. Nat Struct Biol 4: 833-838.
66. Potterton E, Briggs P, Turkenburg M, Dodson E (2003) A graphical user interface to the CCP4 program suite. Acta Crystallogr D Biol Crystallogr 59: 1131–1137.
67. Schuttelkopf AW, van Aalten DM (2004) PRODRG: a tool for high-throughput crystallography of protein-ligand complexes. Acta Crystallogr D Biol Crystallogr 60: 1355–1363.
68. DeLano WL (2002) The PyMOL Molecular Graphics System.
69. Hayward S, Berendsen HJ (1998) Systematic analysis of domain motions in proteins from conformational change: new results on citrate synthase and T4 lysozyme. Proteins 30: 144–154.
70. Schwarz JB, Kuduk SD, Chen XT, Sames D, Glunz PW, et al. (1999) A broadly applicable method for the efficient synthesis of alpha-O-linked glycopeptides and clustered sialic acid residues. J Am Chem Soc 121: 2662–2673.
71. Mitchell J, Siboo IR, Takamatsu D, Chambers HF, Sullam PM (2007) Mechanism of cell surface expression of the Streptococcus mitis platelet binding proteins PblA and PblB. Mol Microbiol 64: 844–857.
72. Takamatsu D, Bensing BA, Sullam PM (2004) Genes in the accessory sec locus of Streptococcus gordonii have three functionally distinct effects on the expression of the platelet-binding protein GspB. Mol Microbiol 52: 189–203.
73. Bensing BA, Takamatsu D, Sullam PM (2005) Determinants of the streptococcal surface glycoprotein GspB that facilitate export by the accessory Sec system. Mol Microbiol 55: 1408–1411.
74. Alphey MS, Attrill H, Crocker PR, van Aalten DM (2003) High resolution crystal structures of Siglec-7. Insights into ligand specificity in the Siglec family. J Biol Chem 278: 3372–3377.
75. Kleywegt GJ (1996) Use of non-crystallographic symmetry in protein structure refinement. Acta Crystallogr D Biol Crystallogr 52: 842–857.