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Authors

Loukachevitch, Lioudmila V Bensing, Barbara A Yu, Hai [et al.](https://escholarship.org/uc/item/6ss5p5hw#author)

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Structures of the Streptococcus sanguinis SrpA Binding Region with Human Sialoglycans Suggest Features of the Physiological Ligand

Lioudmila V. Loukachevitch\$, **Barbara A. Bensing**†, **Hai Yu**#, **Jie Zeng**#,^ , **Xi Chen**#, **Paul M. Sullam**†, and **T M Iverson**\$,%,&,@,*

\$Department of Pharmacology, Vanderbilt University, Nashville, Tennessee 37232, USA

%Department of Biochemistry, Vanderbilt University, Nashville, Tennessee 37232, USA

&Center for Structural Biology, Vanderbilt University, Nashville, Tennessee 37232, USA

@Vanderbilt Institute of Chemical Biology, Vanderbilt University, Nashville, Tennessee 37232, USA

†Division of Infectious Diseases, Veterans Affairs Medical Center, University of California at San Francisco and the Northern California Institute for Research and Education, San Francisco, California 94121, USA

#Department of Chemistry, University of California, Davis, CA 95616, USA

^School of Food Science, Henan Institute of Science and Technology, Xinxiang, 453003, China

Abstract

Streptococcus sanguinis is a leading cause of bacterial infective endocarditis, a life threatening infection of heart valves. S. sanguinis binds to human platelets with high avidity, and this adherence is likely to enhance virulence. Previous studies suggest that a serine-rich repeat adhesin termed SrpA mediates the binding of S. sanguinis to human platelets via its interaction with sialoglycans on the receptor GPIba. However, in vitro binding assays between SrpA and defined sialoglycans failed to identify specific high-affinity ligands. To better understand the interaction between SrpA and human platelets, we determined cocrystal structures of the SrpA sialoglycan binding region (Sr_{PRR}) with five low-affinity ligands: three sialylated trisaccharides (sialyl-T antigen, 3'-sialyllactose, and 3'-sialyl-N-acetyllactosamine), a sialylated tetrasaccharide (sialyl-Lewis^X) and a sialyl galactose disaccharide component common to these sialoglyans. We then combined structural analysis with mutagenesis to further determine whether our observed interactions between $SrpA_{BR}$ and glycans are important for binding to platelets and to better map the binding site for the physiological receptor. We found that the sialoglycan binding site of

^{*}**Corresponding Author** To whom correspondence should be addressed: tina.iverson@vanderbilt.edu.

Author Contributions LVL performed crystallization and collected diffraction data; BAB performed site-directed mutagenesis, binding assays, and data analysis; HY and JZ synthesized sialyl T-antigen and the Neu5Ac-galactose disaccharide under the direction of XC; PS analyzed data; TMI determined the structures, performed crystallographic refinement, and analyzed the data. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. PDB coordinates and structure factors have been deposited with the RCSB protein data bank (www.rcsb.org/pdb) with accession codes 5IIY, 5IJ1, 5IJ2, 5IJ3, and 5KIQ.

 $SrpA_{BR}$ is significantly larger than the sialoglycans cocrystallized in this study, which suggests that SrpA binding to platelets is either multivalent or occurs via a larger, disialylated glycan.

INTRODUCTION

Serine-rich repeat (SRR) adhesins are cell wall anchored glycoproteins of streptococci and staphylococci that mediate adherence of the organism to proteins or glycans¹. These interactions may be critical for commensal or pathogenic bacteria to attach to host tissues, and have been shown to enhance virulence 2^{-7} . The SRR adhesins contain ligand-binding domains that can fold independently and are highly variable in both their sequences and ligand repertoires. The differing specificities of the SRR adhesins for host receptors could explain the tropisms of the associated pathogens for specific tissues or anatomical sites.

Viridans group streptococci are a major cause of bacterial infective endocarditis^{8, 9}. Attachment of the organism to platelets is a required step in the pathogenesis of this infection^{10, 11}, and many streptococcal strains express SRR adhesins that mediate binding to sialoglycans on platelet glycoprotein GPIb^{2, 4, 12–14}. Among the best-studied sialoglycanbinding adhesins are SrpA from *Streptococcus sanguinis* strain SK36, GspB from Streptococcus gordonii strain M99, and Hsa from S. gordonii strain Challis (Fig. 1). SrpA, GspB, and Hsa can each mediate streptococcal adherence to carbohydrates on human platelets via a conserved sialoglycan binding region (BR) termed $SrpA_{BR}$, $GspB_{BR}$, and Hsa_{BR}, respectively^{2, 4, 15}. The BRs of these three adhesins are clear homologs, with SrpA_{BR} 32% identical to the corresponding region of GspB_{BR}^{15} and 51% identical to $H\text{sa}_{BR}^3$. However, there are major differences in the carbohydrate-binding spectrum of SrpA, GspB, and Hsa¹⁶. GspB, which is expressed by a human endocarditis isolate and contributes to virulence in animal models of this disease, has nanomolar affinity for platelets⁴ and binds a very narrow range of sialoglycans¹⁶. Hsa was cloned from an oral isolate³ and similarly exhibits nanomolar affinity to human platelets⁴ and a broad range of human sialoglycans¹⁷. SrpA was also identified in an oral isolate, and binds to platelets with similar avidity as GspB and $Hsa^{12, 14}$, suggesting a high-affinity interaction. As compared with these adhesins, however, SrpA appears to have markedly lower affinity for the defined sialoglycans that have been tested to date¹⁶. These data indicate that the specific sialoglycan attached to the GPIb receptor and recognized by SrpA has yet to be identified.

In recent work, we combined X-ray crystallography with functional binding measurements to identify the structural basis for sialoglycan binding by both $SrpA_{BR}^{18}$ and $GspB_{BR}^{19, 20}$. These studies included a high-resolution costructure of $SrpA_{BR}$ bound to a non-human N glycolylneuraminic acid (Neu5Gc)-based sialyl galactoside disaccharide¹⁸. Here, we further explore the binding repertoire of $SrpA_{BR}$, using a range of Neu5Ac-based α 2-3 linked sialoglycans that are low-affinity ligands: sialyl-T antigen (Neu5Acα2–3Galβ1–3GalNAc; Fig. 2A), 3'-sialyllactose (Neu5Acα2–3Galβ1–4Glc; Fig. 2B), 3'-sialyl-N-acetyllactosamine (3'-sialyllactosamine; Neu5Acα2–3Galβ1–4GlcNAc; Fig. 2C), sialyl-LewisX (Neu5Acα2– 3Galβ1–4(Fucα1–3)GlcNAc; Fig. 2D) and a sialyl galactose disaccharide that is common to these trisaccharides (Neu5Acα2–3Gal; the β-galactose form is shown in the crystal structure, Fig. 2E). These studies indicate that the binding region of SrpA differs

significantly from GspB and Hsa in its interactions with sialoglycans. Moreover, SrpA is likely to accommodate either multiple ligands or larger, more complex ligands, as compared to these homologs.

EXPERIMENTAL PROCEDURES

Sialoglycan reagents

Sialyl-T antigen²¹ and sialyl galactose²² were synthesized as described previously. We purchased 3'-sialyllactose (Santa Cruz Biotechnology), 3'-sialyllactosamine (Carbosynth), and sialyl-Lewis^X (Calbiochem).

Expression and Purification of SrpABR

SrpA_{BR} was expressed and purified with significant modifications of our previously described procedure¹⁸. The SrpA_{BR} was expressed with tandem N-terminal His₆ and maltose binding protein (MBP) tags, using the isopropyl β-D-1-thiogalactopyranoside (IPTG) inducible pSV278 vector and E. coli BL21($DE3$). Cells were grown in Terrific Broth at 37 °C until an OD₆₀₀ of 2.0 was reached. The cells were then cooled on ice for 20 min, after which time expression was induced with 0.5 mM of IPTG. Srp A_{BR} was expressed at 18 °C for 24 hrs. Cells were harvested by centrifugation, washed with Tris-HCl buffer (pH 7.5, 0.1 M) and pellets were stored at −20 °C before use. Frozen pellets were resuspended in binding buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl) containing PMSF (1 mM), leupeptin (2 μg/mL), pepstatin (2 μg/mL) and cells were lysed by sonication. The lysate was clarified by centrifugation at 38,500 \times g for 40 min. His₆-MBP-tagged SrpA_{BR} was purified at 4 °C on Ni-NTA agarose resin (Qiagen) using a batch procedure by adding Ni-NTA agarose resin to clarified lysate, incubating for 2 hrs, washing with binding buffer, and eluting tagged protein with binding buffer containing imidazole (250 mM). The eluted protein was concentrated in a 10 kDa MW cut-off concentrator and exchanged into binding buffer. The $His₆-MBP$ affinity tag was cleaved with thrombin, and the cleaved affinity tag was separated from SpA_{BR} by a passage over Ni-NTA agarose column equilibrated with the binding buffer.

Crystallization and Structure Determination

SrpA_{BR} (16 mg/mL in 20 mM Tris-HCl, pH 7.2) was mixed with sialyl-T antigen, 3'sialyllactose, 3'-sialyllactosamine, sialyl-Lewis^X, or the disaccharide sialyl galactose and incubated on ice for 1 hr prior to crystallization. Cocrystals of $SrpA_{BR}$ with each sialoglycan were grown using the sitting drop vapor diffusion method in CombiClover 4 Chamber plates at room temperature (\sim 23 °C) by equilibrating droplets containing 1 µL of proteinsialoglycan complex (14.4 mg/mL Srp A_{BR} , 10 mM sialoglycan, 18 mM Tris-HCl, pH 7.2) and 1μL of reservoir solution over a reservoir solution (50 μL) that contained Ca(CH₃CO₂)₂ (0.2 M), sodium cacodylate, pH 6.5 (0.1 M) and PEG 8000 (18%). Crystals were cryoprotected in reservoir solution containing 5 mM sialoglycan and cryocooled by plunging into liquid nitrogen. Data were collected at −180 °C on a MARCCD detector at LS-CAT ID-G at the Advanced Photon Source, processed using the HKL Suite²³, and converted with the CCP4 suite²⁴ (Table 1).

Cocrystals of SrpA_{BR} with each sialoglycan were isomorphous with the unliganded structure. As a result, all structures were determined by removing solvent molecules from the unliganded $SrpA_{BR}$ structure (PDB entry $5EQ2^{18}$) and transferring the coordinates directly into each data set. Each model was subjected to an initial round of rigid body refinement in Phenix²⁵. Sialoglycans were placed into the model after this initial structure determination and were retained at an occupancy of 1.0 during the course of the refinement process. Refinement proceeded using iterative rounds of $Coot^{26}$ and Phenix²⁵ until the R_{free} and the geometry was considered reasonable for each model (Table 2).

Binding of GST-tagged BRs to Platelet Monolayers

The production of glutathione S -transferase (GST)-tagged $SrpA_{BR}$ variants and the binding of GST-tagged BRs to platelet monolayers by ELISA were performed as previously described 27 . SrpA variant proteins were validated as stably folded using two complementary methods. First, we monitored degradation and chaperone-association using SDS-PAGE, where the chaperone-associated behavior identifies proteins that have exposed hydrophobic regions, and lowered stability. Second, we monitored monodispersity using size exclusion chromatography, where proteins with reduced stability will exhibit polydispersity, as reflected by a broadening of the peak width (not shown). Variants that exhibited either chaperone-associated behavior or polydispersity were not used for further analysis. Only variants that lacked chaperone-associated behavior and exhibited monodispersity were considered to be stably folded and were used in the assays.

Binding of platelet GPIbα to immobilized GST-SrpA_{BR} and GST-GspB_{BR}

GST or GST-BR fusion proteins were diluted to 500 nM in PBS and 50 μL applied to microtiter wells. Plates were incubated 18 hr at 4°C. Unbound proteins were removed by aspiration, and wells were rinsed with 100 μL TEN buffer (10 mM Tris, 1 mM EDTA, 100 mM NaCl, pH 8). An extract enriched for glycocalicin (the soluble extracellular domain of GPIbα) was prepared by sonication of washed human platelets, followed by incubation at 37°C for 1 h to allow the proteolytic cleavage of GPIbα and release of glycocalicin, as described.28 Insoluble debris was removed by centrifugation, followed by passage of the platelet extracts through a 0.45 μm filter. The platelet proteins were diluted to 60 μg/mL in TEN buffer containing Complete protease inhibitors (Roche) and 1X Blocking Reagent (Roche), with subsequent four-fold dilutions in the same buffer. Diluted platelet lysates (50 μL) were added to wells, and plates were incubated at 23 °C for 1 h with vigorous rocking. Wells were washed three times with PBS, and bound GPIba was detected by ELISA, using a rabbit monoclonal anti-CD42b (abcam), followed by a peroxidase-conjugated goat antirabbit IgG (Sigma), and colorimetric detection with α -phenylenediamine (Sigma). The negative control was background subtracted from the measurements.

RESULTS

Relative binding of SrpA_{BR} to platelets

A K_D value for the interaction between SrpA and platelet receptors has not been reported. However, SrpA has previously been implicated as mediating the attachment of S. sanguinis to platelet GPIba at levels that are roughly comparable to the binding of S. gordonii strain

M99, which binds via an interaction between GspB and GPIba, with a reported affinity of 23.8 nM⁴. We therefore used an ELISA assay to compare the relative binding of purified GST-tagged Srp A_{BR} or purified GST-tagged Gsp B_{BR} to intact platelets (Fig. 3A) or GPIba in platelet lysates (Fig. 3B) over a range of concentrations. These studies showed that $SrpA_{BR}$ binds to intact platelets more strongly than does $GspB_{BR}$ under the conditions tested (Fig. 3A). Moreover, the interaction is clearly mediated by the GPIbα glycoprotein, with SrpA_{BR} exhibiting ~10-fold higher avidity for GPIba than $GspB_{BR}$ (Fig. 3B).

Structure of SrpABR with Neu5Ac-based trisaccharides

To explore a range of low-affinity ligands that likely exhibit specific binding to SrpA, we focused on α2-3 linked Neu5Ac-based sialoglycans. In all of these sialoglycans, the identity and the linkage of the first two sugars are identical (Neu5Acα2-3Galβ), with variability in the identity and the linkage to the third sugar in the larger glycans, and the presence of a fourth sugar in sialyl-Lewis^X. Our studies included two trisaccharides known to decorate platelets (sialyl-T antigen (Fig. 2A), 3'-sialyllactosamine (Fig. 2C)) and the sialyl galactose disaccharide (Fig. 2E). We additionally examined the interaction with two low-affinity ligands not reported to be associated with platelets (3'-sialyllactose (Fig. 2B) and sialyl-Lewis^X (Fig. 2D)). The rationale for evaluating the trisaccharides that have not been found on platelets is that these can still provide insights as to the chemical properties of the physiological receptor.

We determined the high-resolution X-ray crystal structures of $SrpA_{BR}$ in complex with sialyl-T antigen, 3'-sialyllactose, 3'-sialyllactosamine, sialyl-Lewis^X and the sialyl galactose disaccharide (Fig. 4, Tables 1&2). Following isomorphous replacement, unambiguous electron density for each sialoglycan was readily apparent (Fig. 5, Table 3). For all the sialoglycans, the subjective quality of the electron density was the most unambiguous for the sialic acid. Moreover, with the exception of 3'sialyllactosamine, the contour level of the electron density where the hexose ring of each carbohydrate was associated with contiguous density was highest for the sialic acid (Table 3). This is consistent with the many direct hydrogen-bonding interactions between Neu5Ac and SrpA_{BR}. The hydrogen bonds involve the side chains of Gln 344, Thr 346, and Arg 347 (Fig. 6–9) and the main chain carbonyls of Arg 342 and Gln 344 (Fig. 8). Validating this binding mode, we previously showed that Thr 346 and Arg 347 are important for the interaction between SrpA and platelets via sitedirected mutagenesis¹⁸. The electron densities for additional sugars were almost universally weaker (Fig. 5, Table 3), but were consistent with favorable conformations of each sialoglycan. For the trisaccharides and sialyl-Lewis^X (Fig. 5A–D), elaborations on both the galactoside and the third sugar could be observed in the initial electron density when contoured at or below 2σ. For the sialyl galactose disaccharide, the electron density of the galactose was quite weak and could only be observed when contoured below 2σ even after refinement, suggesting mobility of the galactose (Fig. 5E). The quality of the electron density for the galactose in the context of the disaccharide was subjectively worse than the quality of the electron density for galactoside in the context of the trisaccharides, suggesting that interactions between $SrpA_{BR}$ and the third sugar reduces the mobility of the galactoside.

As we previously observed for $SrpA_{BR}$ crystallized with a Neu5Gc-based sialyl galactoside disaccharide¹⁸, only one of the protomers in the dimer was associated with clear electron density for sialoglycan ligand, with diffuse electron density apparent in the binding site in the second protomer of the dimer. Crystal packing interactions likely prevent binding of the ligand to the other binding site in the dimer, and previously-reported mutagenesis disrupting dimerization was not associated with changes in binding affinity¹⁸, indicating no cooperativity between the sites. The single site sialoglycan binding to the dimer is depicted in Fig. 4.

Binding of di-, tri-, and tetrasaccharides to SpA_{BR} did not require backbone adjustments as compared to the unliganded structure, as evidenced by the rms deviation between sialoglycan-bound and unliganded SrpA_{BR} ranging from $0.21 - 0.30$ Å. A single side chain rotamer change, in Lys 296, accompanied glycan binding in the 3'-sialyllactosamine and sialyl-Lewis^X-bound SrpA. In each case, this made a weak hydrogen-bonding interaction with the third sugar; however the electron density suggested that this residue was somewhat mobile in all structures.

Differences in interaction between SrpA and Neu5Ac- or Neu5Gc-based sialoglycans

Naturally occurring sialic acid has different forms. Humans exclusively synthesize the Nacetylneuraminic acid (Neu5Ac) form; other animals also hydroxylate its nucleotide activated form, cytidine 5'-monophosphate-N-acetylneuraminic acid (CMP-Neu5Ac), on C11, resulting in the Neu5Gc form29 (compare Fig. 2 E, F). Previous binding studies of SrpA to purified, defined glycans identified that SrpA preferentially binds the Neu5Gc form of sialic acid¹⁶. Comparison of the costructure of $SrpA_{BR}$ in complex with the Neu5Gcbased sialyl galactoside disaccharide¹⁸ to each of the costructures of $SrpA_{BR}$ with each of the Neu5Ac-based sialyl galactose compounds identifies a favorable hydrogen-bonding interaction between the OH of Tyr 368 of SrpA_{BR} and the unique hydroxyl of Neu5Gc (Fig. 6) that is absent in the Neu5Ac-based sialoglycans (Fig. 2A–E). This may explain the preference of SrpA for binding Neu5Gc. Attempts to verify this experimentally were unsuccessful due to the misfolding of site-directed variants, as indicated by evidence of protein degradation and co-purification with chaperones (not shown).

Sialoglycan interactions with SrpA

Interestingly, there are no direct hydrogen-bonding interactions between $SrpA_{BR}$ and the second, third, or fourth carbohydrate in any of the sialoglycans tested. Instead, these sugars exhibited two types of relatively non-selective interactions with SrpA. The first type of interaction between SrpA_{BR} and these sialoglycans is achieved via positioning of an apical CH from the third sugar above the π -ring of Phe 294 (Fig. 7), which forms a stabilizing CH/ π-stacking interaction. CH/π-stacking interactions are common in protein-carbohydrate interactions³⁰ and are proposed to promote weak facial selectivity for their ligands³¹. Notably, changing the linkage from 1➔3 (sialyl-T antigen) to 1➔4 (3'-sialyllactosamine, 3'-sialyllactosamine) and the identity of the sugar from N-acetylgalactosamine (GalNAc; sialyl-T antigen) to glucose (Glc; 3'-sialyllactose) or N-acetylglucosamine (GlcNAc; 3' sialyllactosamine) is accompanied by a 180° rotation of the third sugar around the glycosidic linkage, which maintains the CH/ π -stacking interaction. In comparison, the sialyl-Lewis^X

tetrasaccharide appears to be slightly rotated, which places the glycosidic linkage between the GlcNAc and the fucose (Fuc) above the Phe 294 tetrapole.

The second type of interaction between $SrpA_{BR}$ and the second and the third sugars of the trisaccharides is indirect, water molecule-mediated interactions (Fig. 8). The water-mediated hydrogen-bonding network extending from these sugars is extensive and structurally conserved in the costructures with all three trisaccharides (Fig. 8A–C) and only slightly altered with the sialyl-Lewis^X tetrasaccharide (Fig. 8D). Close examination of this region identifies that the water molecule network extends through a shallow but extended surface cleft of $SrpA_{BR}$. While the functional relevance of this cleft has never been tested in SrpA, surface clefts of proteins are hallmarks of ligand-binding sites.³² Interestingly, comparison of sialoglycan-bound $SrpA_{BR}$ or $GspB_{BR}$ to their unliganded counterparts identifies that positions occupied by hydroxyl groups of bound sialoglycans superimpose with water molecules in the unliganded structures. In considering this more extended water molecule network and the cleft adjacent to the bound trisaccharides, it appears that the available ligand-binding surface on the protein could be larger than required to accommodate a trisaccharide.

Molecular properties of the SrpA sialoglycan binding site

While the physiological sialoglycan ligand for SrpA remains unknown, structural comparison of SrpA with mammalian sialic acid binding immunoglobulin-like lectins (Siglec) offers one intriguing hypothesis. Mammalian Siglecs share a fold with SrpA and its bacterial homologs, but despite the similar fold, these two families of sialic acid binding proteins bind sialoglycans at distinct sites.^{33–39} The bacterial homologs use the YTRY motif to anchor sialic acid above a strictly conserved arginine residue, which is Arg 347 in SrpA, Arg 484 in GspB, and Arg 340 in Hsa (Fig. 1). The mammalian Siglecs also bind at a conserved arginine (Arg 97 of Siglec-1 (sialoadhesin), Arg 124 of Siglec-5, or Arg 124 of Siglec-7, Fig. 9A), however this conserved mammalian arginine is not analogous to the bacterial arginine. Indeed, in a structural overlay of bacterial and mammalian Siglecs, the critical sialoglycan-binding arginines of each family are spatially separated by \sim 15 Å and appear to be parts of distinct sialoglycan binding pockets (Fig. 1, Fig. 9A–C). Although spatially separated, both the mammalian arginine and the bacterial arginine are located on the F-strand of the Ig fold.

Interestingly, SrpA appears to contain arginine residues corresponding to both a bacterial and a mammalian sialoglycan binding pocket (compare Fig. 9A–C); the bacterial sialoglycan-binding arginine of SrpA is Arg 347 and the mammalian-like sialoglycanbinding arginine is Arg 350 (Fig. 9B). SrpA is the first known example of either a mammalian or a bacterial Siglec that contains an arginine corresponding to the critical sialic acid binding residue of the other family. Indeed, the GspB and Hsa bacterial Siglecs that are homologs of SrpA do not have an arginine analogous to the mammalian sialoglycan binding arginine (Fig. 1).¹⁹ In addition, the structure of $Gs p B_{BR}$ has been reported, and shows that the sialoglycan binding pocket is capped by a helix (Fig. 9C) that physically covers the mammalian binding site.¹⁹

If Arg 350 of SrpA were important for platelet binding, it could mean that a second, mammalian-like sialoglycan binding site is present in this bacterial adhesin, which suggests additional possibilities for the natural SrpA ligand. One possibility is that the mammalianand bacterial sialoglycan binding sites of are connected so that SrpA can bind a larger and more complex sialoglycan ligand than has been previously tested, in particular a ligand that contains two sialic acids. A second possibility is that both the mammalian-like and bacteriallike binding cooperate to bind two glycans through a multivalent interaction.

Functional validation of SrpA residues required for platelet binding

To test which residues of the sialoglycan binding site in $SrpA_{BR}$ are important for binding to the physiological receptor, we assessed the impact of selected site-specific variations on the binding of purified $SrpA_{BR}$ to immobilized platelets (Fig. 10). First, we assessed the impact of mutating Phe 294, which would disrupt the CH/π -stacking interaction. Second, we assessed the impact of mutating Asn 361 and Thr 363, which contribute to the stabilization of the network of water-mediated hydrogen-bonding interactions between SrpA and each of the tri- and tetrasaccharides, but could also interact directly with a larger sialoglycan. Finally, we assessed the impact of mutating Arg 350, which does not interact either directly or indirectly with any of the trisaccharides or the sialyl-Lewis^X tetrasaccharide and is not conserved in bacterial Siglecs (Fig. 1), but which is analogous to the critical sialic acidbinding arginine of mammalian Siglecs. All variants folded correctly, as described in the experimental procedures (not shown). Moreover, the variants altered surface residues that are not conserved in SrpA homologs with different ligand specificity (Fig. 1), indicating that this domain can accommodate a range of amino acids at these positions. We compared the platelet binding of each variant to platelet binding exhibited by wild-type SrpA_{BR} and the SrpA_{BR}^{R347E} variant, a variant with the critical arginine of the bacterial YTRY sialic acidbinding motif altered. The Srp A_{BR}^{R347E} variant has previously been demonstrated to significantly reduce platelet binding while retaining a folded structure²⁷, thus allowing it to serve as a control.

 $\text{SrpA}_{\text{BR}}^{\text{F294A}}$, designed to disrupt CH/ π -stacking, exhibited a modest but statistically significant reduction in platelet binding as compared to wild-type $SrpA_{BR}$. Srp A_{BR} ^{T363A} and SrpA_{BR}^{N361A}, designed to disrupt either the water molecule network or interactions to a larger physiological ligand, exhibited different effects on platelet binding. $\rm SrpA_{BR}^{T363A}$ was associated with platelet binding that was only slightly reduced, as compared to wild-type SrpA_{BR}. In contrast, SrpA_{BR}^{N361A} showed a substantial reduction in binding to platelets. The SrpA_{BR}^{R350E} variant, designed to eliminate the mammalian-like sialoglycan-binding arginine, also displayed substantial loss of binding as compared to wild-type $SrpA_{BR}$. These mutagenesis studies identify Asn 361 and Arg 350 as important for high-affinity binding to platelet receptors. As previous studies identified both Thr 346 and Arg 347 to be key for platelet binding, and as Thr 346, Arg 347, Asn 361 and Arg 350 lie along a contiguous surface, the most reasonable interpretation of these results is that the functional SrpA receptor-binding pocket is much larger than is necessary to accommodate the ligands tested here (Fig. 9B). The features of this binding region could either be interpreted as two separate binding pockets for trisaccharides, or as a single binding pocket that binds a larger, more complex glycan.

DISCUSSION

The nature of the physiological sialoglycan receptor of SrpA during endocarditis

SrpA has been demonstrated to bind to human platelet $GPIb¹⁸$, a highly sialylated glycoprotein. Moreover, SrpA contains strong sequence identity to GspB and $Hsa⁴⁰$, which both exhibit sialidase-sensitive platelet binding properties⁴¹. The sequence identity between SrpA, GspB, and Hsa extends to a loosely conserved YTRY sequence motif that binds to sialic acid (Fig. 1). The sequence is FTRT in $SrpA⁴⁰$. The central Thr-Arg residues of this motif (Thr 346 and Arg 347) have been shown to be critical for platelet binding by SrpA, and these side chains directly interact with sialic acid in the crystal structures¹⁸. However, a defined high-affinity ligand for SrpA has not, to date, been identified¹⁶.

Both the sialyl-T antigen and 3'-sialyllactosamine tested in these studies are known to decorate platelets; however, the full repertoire of platelet glycans is not known at this time, which hinders efforts to identify the receptor. Here, we identified two additional surface residues, Asn 361 and Arg 350, that are distal to the FTRT sequence and that impact SrpA affinity to platelets (Fig. 10). Asn 361 and Arg 350 are on a surface of SrpA that also contains the FTRT sequence (Fig. 9B), with the two arginines (Arg 347 and Arg 350) separated by \sim 15 Å. This receptor-binding surface in SrpA appears significantly larger than sialyl-T antigen-binding pocket on the GspB homolog (Fig. 9C).

One interpretation of these data is that SrpA improves affinity to platelets via a multivalent interaction, where SrpA simultaneously binds one sialoglycan above Arg 347 and a second sialoglycan (or another ligand) above Arg 350 (Fig. 11A). In this model, the two sialoglycans would be adjacent on the GPIb receptor. If this were the case, it might be anticipated that two glycans would bind to SrpA during crystallization. However, crystal contacts closely approach Arg 350, which would prevent a ligand from binding at this site. As a result, the lack of observed sialoglycan binding near Arg 350 in these crystal structures does not rule out the possibility of a multivalent interaction.

An alternative would be that SrpA binds a larger glycan (Fig. 11B) that interacts with both Arg 347 and Arg 350 simultaneously. Consistent with this second mechanism, both the mammalian-like Arg 350 and the bacterial-like Arg347 in SrpA are located on the F-strand of the Ig fold¹⁹ and are arranged in a way that they could be a part of the same binding site. If SrpA does bind to a larger glycan during endocardial infection, the identity of this ligand is not immediately apparent because the full repertoire of sialoglycans present on platelets or GPIb is not well characterized.

One possibility could include sulfate elaborations of sialoglycans. Although not reported on GPIb to date, experimental support for a role of sulfated sialoglycans in binding comes from studies using sialoglycan arrays¹⁶. These arrays showed that sulfated substitutions of di- and trisaccharide sialoglycans bind to SrpA with similar or slightly increased affinity as compared to the non-sulfated versions¹⁶.

Other possibilities for the true receptor are longer core 1 structures, and branched core 2 structures, potentially disialylated. Previous studies of the platelet glycoprotein GPIbα

identified a core 2 hexasaccharide comprised of sialyl-T antigen with β1–6 glycosidic linkage between the GlcNAc in 3'-sialyllactosamine Siaα2–3Galβ1–4GlcNAc and the core GalNAc in the sialyl-T antigen^{42, 43} (this structure is shown in the conceptual model in Fig. 11B). This branched core 2 structure was reported to be the most abundant O-linked glycan on platelet GPIbα. If this disialylated hexasaccharide adopted an elongated conformation, its two sialic acids would be separated by \sim 15 Å, a distance ideal for interaction with Arg 347 and Arg 350. The central part of the glycan could then interact with Asn 361. This hexasaccharide is not readily available, making it difficult to experimentally validate this possibility at the present time.

Within these candidates for the SrpA receptor in humans, a Neu5Gc-containing sialoglycan cannot be excluded a priori. Although Neu5Gc is not synthesized in humans²⁹, there are some reports that low levels of Neu5Gc can be incorporated into the human glycan repertoire after absorption from dietary sources^{$44–46$}. While there is no known correlation between a Neu5Gc-containing diet and endocardial infection caused by S. sanguinis, the observed additional hydrogen-bonding interaction between SrpA and Neu5Gc would suggest that a Neu5Gc-incorporated sialoglycan would be the preferred ligand over a Neu5Ac-incorporated sialoglycan.

Conclusions

Here, we determine that the SrpA adhesin contains a larger ligand binding surface than is necessary to bind a trisaccharide. This surface contains residues critical for platelet binding, including Thr 346, Arg 347, Arg 350, and Asn 361. The ligand binding site appears to be a composite of the binding sites identified in the bacterial homolog GspB and the mammalian Siglecs, both of which bind to sialoglycans. These data allow the proposal of a model where SrpA binds with much higher affinity to human platelets than to synthetic trisaccharides either by recognizing multiple, adjacent sialoglycans (increasing affinity with multivalency), or by recognizing a much larger sialoglycan (increasing affinity with additional contacts).

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ABBREVIATIONS

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Figure 1.

Sequence alignment of the Siglec domain of SrpA, GspB, and Hsa. Residues conserved in all three homologs are marked with a colon (:). The YTRY sialic acid-binding motif is highlighted with a green box. As discussed in the text, this motif contains a conserved arginine residue (Arg 347 of SrpA, Arg 484 of GspB and Arg 340 of Hsa) that makes direct hydrogen-bonding interactions to the sialic acid. Additional key residues discussed in the text are highlighted with arrows. These include the putative Neu5Gc selectivity residue Tyr 368 (highlighted with a red arrow), and the putative glycan binding residues Phe 294, Arg 350, Asn 361, and Thr 363 (highlighted with a blue arrow). The cation binding residues are highlighted with yellow boxes.

 \sim OH

 Ω

B) sL (Neu5Ac α 2-3Gal β 1-4Glc β)

 C) sLn (Neu5Ac α 2-3Gal β 1-4GlcNAc β)

D) sLe^X [Neu5Ac α 2-3Gal β 1-4(Fuc α 1-3)GlcNAc β]

F) Neu5Gc-Gal (Neu5Gc α 2-3Gal β) HO HO OH O_2C HO OH HO NH HO \overline{a} -OH \overline{h}

Figure 2.

Sialoglycans used in this study. (A) sialyl-T antigen (sTa), (B) 3'-sialyllactose (sL), (C) 3' sialyllactosamine (sLn), (D) sialyl-Lewis^X (sLe^X), (E) Neu5Ac-Gal. (F) Neu5Gc-Gal. Shaded boxes highlighting the third sugar have the same color as the stick structures in Figs. 3, and $5 - 7$.

Figure 3.

Comparative binding of $SrpA_{BR}$ and $GspB_{BR}$ to human platelets or to platelet GPIba. (A) Binding of GST-BRs (5–500 nM) to platelet monolayers. Measurements were background subtracted. (B) Binding of platelet GPIbα to immobilized GST-BRs. Data are reported as the mean \pm standard deviation, with n=4.

Figure 4.

Overview of a trisaccharide-bound $SrpA_{BR}$ dimer. The two polypeptide chains forming the dimer are depicted with the N-terminal Siglec domain colored cyan and the C-terminal Unique domain colored pink. Dimerization is mediated by the Unique domain. The sialyl-T antigen trisaccharide bound to the Siglec domain is shown in yellow sticks. Constraints of the crystal packing interactions likely result in the binding of only one sialoglycan molecule to the dimer. Divalent cations proposed to be physiologically relevant are shown as yellow spheres; additional ions from the crystallization conditions are not shown.

Figure 5.

Electron density for Neu5Ac-based sialylated trisaccharides binding to $SrpA_{BR}$ (A) sialyl-T antigen (B) 3'-sialyllactose (C) 3'-sialyllactosamine (D) sialyl-LewisX (E) sialyl galactose. In each panel, $|F_0| - |F_c|$ electron density calculated after the removal of the sialoglycan and three cycles of refinement in Phenix²⁵ is contoured at 3σ (*black*) and 1.8σ (*white*) and is overlaid with the final model of the Siglec domain of each $SrpA_{BR}$ costructure. In all panels, the Neu5Ac is colored grey, the galactoside is colored navy, oxygens are colored red, and nitrogens are colored blue. For the trisaccharides, the third sugar differs in both identity and linkage and is colored yellow (1➔3GalNAc, sialyl-T antigen), green (1➔4 Glc, 3' sialyllactose) or *magenta* (1→4 GlcNAc, 3'-sialyllactosamine and sialyl-Lewis^X). The fucose of sialyl-Lewis^X (Fuca1 \rightarrow 3GlcNAc) is colored white.

Figure 6.

A comparison of Neu5Ac and Neu5Gc-based disaccharides bound to SrpABR. The Neu5Gcbased sialyl galactoside disaccharide is shown in black, the Neu5Ac-based sialyl galactose disaccharide is shown in *lavender*, oxygens are colored *red*, and nitrogens are colored *blue*. Hydrogen bonding interactions have the same color as each disaccharide. The synthetic Neu5Gc-based sialyl galactose makes an additional hydrogen-bond to Tyr 368 as compared to the Neu5Ac-based glycans (black dotted line).

Figure 7.

Overlay of trisaccharides bound to SrpA_{BR}. The views in (A) and (B) are separated by 90 $^{\circ}$. The trisaccharides are colored as in Figure 5, with the conserved glycans (Neu5Ac, grey; Gal, navy) represented by the positions in sialyl-T antigen, and the direct hydrogen-bonding contacts between Neu5Ac and SrpA_{BR} are the same in all three trisaccharides. Carbon atoms and bonds of the third sugar are colored yellow (sialyl-T antigen) green (3'-sialyllactose) or magenta (3'-sialyllactosamine), respectively. Oxygens are colored *red*, and nitrogens are colored blue. Direct interactions include both hydrogen-bonding interactions between the protein and the Neu5Ac, and a CH/ π stacking interaction (*blue* dashed line) between the hexose ring of the third sugar and the side chain of Phe294.

Figure 8.

Comparison of the direct and water mediated hydrogen-bonding interactions between SrpA_{BR} and (A) sialyl-T antigen (B) 3'-sialyllactose (C) 3'-sialyllactosamine and (D) sialyl-Lewis^X. Direct hydrogen-bonding interactions between $SrpA_{BR}$ and the sialic acid are observed for all three sialoglycans (light grey dotted lines). Some hydrogen-bonding interactions involve backbone atoms; for example Arg 342 (discussed in the text) has an interaction to the backbone carbonyl. The Cα of Arg 342 is marked with a *. Water molecules (red spheres) form hydrogen-bonding interactions that bridge the galactoside to Gln 344/Thr 363/Thr 346 (navy dashed lines) and bridge the third sugar to Asn 361 (dotted lines colored similarly to the third sugar). No direct or indirect interactions to the fucose of sialyl-Lewis^X are observed. Minimal alteration in water molecule structure and hydrogenbonding pattern is required to accommodate the different elaborations and linkages on the three trisaccharides (panels A–C). However, the rotation of sialyl-Lewis^X with respect to the remaining glycans alters the water molecule structure somewhat. The view is rotated 180° in the z-axis and 75° in the x-axis with respect to Figure 5; this view is rotated approximately 180° around the y-axis with respect to Figures 6B and 7B.

Figure 9.

The SrpA sialoglycan binding site has additional functional regions. $(A) - (C)$ Side-by-side comparison of the isolated Siglec domain of (A) mammalian Siglec-5 bound to 3' sialyllactose (PDB entry 2ZG3³³) (B) SrpA bound to 3'-sialyllactose and (C) GspB bound to sialyl-T antigen (PDB entry 5IUC). Residues equivalent to Arg 347 and Arg 350 of SrpA are highlighted. The view is rotated by 30° in both the x- and y-axes as compared to the view in Fig. 3 in order to better observe both sialoglycan binding sites. (A) In mammalian Siglec-5, sialic acid binds forms two hydrogen-bonding interactions to Arg 124 (*pink*) on the F-strand of the V-set Ig fold. Tyr 121 (cyan) is observed in the same position as the sialic acidbinding arginine of SrpA and its homolog GspB. (B) In SrpA, the sialic acid binds to the FTRT sequence motif and forms two hydrogen-bonding interactions to Arg 347 on the Fstrand $(cyan)$, but also contains Arg 350, analogous to the sialic acid binding sequence of siglec-5 (*pink*). This second, putative sialic acid binding site is \sim 15 Å distal and is unoccupied in these structures, with crystal packing interactions occluding access to this region. Additional residues that were investigated by site-directed mutagenesis are highlighted in *magenta*. (C) In the SrpA homolog GspB, only the SrpA-like sialoglycan binding site is observed, and is organized around Arg 484 (cyan). Instead of a second arginine residue, GspB has Val 487 (pink) at this position. Moreover, a helix unique to GspB (magenta) is in the same location as the binding resides of siglec-5. This helix provides a defined boundary for the GspB binding pocket that limits the size of the glycan and blocks access to Val 487.

Figure 10.

Binding of N-terminally GST-tagged SrpA_{BR} variants to platelet monolayers, as monitored by ELISA. Measurements were background subtracted for non-specific binding between GST and platelets. Measurements were performed in triplicate and are expressed as the mean ±S.D. All variants had binding that was significantly different from wild-type, as assessed as assessed by analysis of variance (ANOVA) combined with the Bonferroni multiple comparisons test, * p 0.02 , *** p 0.001 .

Figure 11.

Conceptual model for the binding of SrpA to platelet glycans. View is rotated 45° around the x-axis as compared to Fig. 9. Side chains that have a dramatic impact on platelet binding are shown as sticks. (A) Model for a multivalent interaction mode with sialyl-T antigen bound to the GspB-like binding site and 3'-sialyllactosamine bound to the mammalian-like binding site. (B) Model for an interaction with a larger, disialylated carbohydrate. Shown is the extended core 2 hexasaccharide Neu5Acα2–3Galβ1–3(Neu5Acα2–3Galβ1–4GlcNAcβ1– 6)GalNAcα.

Table 1 Crystallographic data collection statistics

Raw diffraction data can be accessed using doi:10.15785/SBGRID/###, where ### indicates the SBGrid entry number listed above.

 a The completeness of the outer shell data (from 1.83–1.80 Å resolution) for the 3'-sialyllactose costructure was influenced by the geometry used during data collection. The data are 71.6% complete in the second to last resolution shell (1.86 – 1.83 Å resolution).

Table 2

Crystallographic refinement statistics.

Table 3

Electron density of bound sialoglycans

Relative electron density contours where each sugar is associated with contiguous density. NA= not applicable.

