A Specific Interaction between SecA2 and a Region of the Preprotein Adjacent to the Signal Peptide Occurs during Transport via the Accessory Sec System*

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Background: Transport of preproteins by the accessory Sec system requires a specific sequence (the AST domain) adjacent to the signal peptide.

Results: The AST domain interacts specifically with SecA2, and this interaction is modulated by the accessory Sec proteins Asp1–3.

Conclusion: The Asp-dependent AST-SecA2 interaction is essential for transport.

Significance: A better understanding of accessory Sec transport may aid the development of novel antimicrobials.

The accessory Sec systems of streptococci and staphylococci mediate the transport of a family of large, serine-rich glycoproteins to the bacterial cell surface. These systems are comprised of SecA2, SecY2, and three core accessory Sec proteins (Asp1–3). In Streptococcus gordonii, transport of the serine-rich glycoprotein GspB requires both a unique 90-residue N-terminal signal peptide and an adjacent 24-residue segment (the AST domain). We used in vivo site-specific photo-cross-linking to identify proteins that interact with the AST domain during transport. To facilitate this analysis, the entire accessory Sec system of S. gordonii was expressed in Escherichia coli. The determinants of GspB trafficking to the accessory Sec system in E. coli matched those in S. gordonii, establishing the validity of this approach. When the photo-cross-linker was placed within the AST domain, the preprotein was found to cross-link to SecA2. Importantly, no cross-linking to SecA was detected. Cross-linking of the N-terminal end of the AST domain to SecA2 occurred regardless of whether Asp1–3 were present. However, cross-linking to the C-terminal end was dependent on the Asps. The combined results indicate that full engagement of the AST domain by SecA2 is modulated by one or more of the Asps, and suggest that this process is important for initiating transport.

The serine-rich repeat (SRR)2 glycoproteins of Gram-positive bacteria are a unique family of bacterial adhesins that have a significant impact on biofilm formation and disease (1–10). The SRR glycoproteins all have a similar domain organization, with an unusually long N-terminal signal peptide followed by a short SRR domain, a ligand binding domain, a long SRR domain, and a C-terminal LPXTG cell wall anchoring motif. Despite the similar domain organization, the ligand binding domains are surprisingly diverse in terms of amino acid sequences, predicted structures, and binding properties (11–15).

The SRR glycoproteins are transported to the bacterial cell surface by a specialized transporter known as the accessory Sec (aSec) system (3, 16–18). These specialized transporters are invariably comprised of five proteins: SecA2, SecY2, Asp1, Asp2, and Asp3. In some streptococcal species such as Streptococcus gordonii, the aSec system may include one or two additional proteins (Asp4 and Asp5). Studies of the aSec system in S. gordonii have indicated that this transport system has some striking similarities to, and some important differences from, the general Sec system. SecA2 resembles SecA both in primary amino acid sequence and in the presence of conserved residues and domains, except it lacks the C-terminal SecB binding region (17). SecA2 can hydrolyze ATP and is likely to function as the transport motor (19). SecY2 resembles SecY and is likely to form the transmembrane channel (20). Asp4 or Asp5 might interact with SecY2 and function similarly to SecE or SecG, but this has not yet been verified (21).

The most notable difference between the Sec and aSec systems is that the latter has three proteins (Asp1–3) that are essential for transport (22, 23), but have no obvious counterpart in the general Sec system. Although recent studies in both S. gordonii and Streptococcus parasanguinis have begun to uncover interactions that occur among the Asps (termed Gap1–3 in S. parasanguinis) (23, 24), the precise role of these proteins in transport remains obscure. Asp2 and Asp3 can bind the serine-rich regions of GspB (the SRR glycoprotein of S. gordonii), suggesting that one or both Asps might function as chaperones to facilitate targeting (25). However, why the aSec system should require two specialized chaperones is unknown.

There are also distinct differences in the preproteins that are transported by the aSec and general Sec systems. We have previously used a truncated version of GspB (GspB736FLAG; Fig. 1A)
as a model substrate to analyze both the means by which the aSec system recognizes the preprotein substrate, and the features of the preprotein that facilitate trafficking to the aSec versus Sec system. Although the carbohydrate moieties on GspB strongly impede Sec transport, they are not necessary for recognition by the aSec system, since nonglycosylated GspB<sub>B.g</sub>-36FLAG is efficiently transported by the aSec system. Instead, an atypical 90 amino acid signal sequence at the N terminus of GspB is required for transport (26). Trafficking is largely determined by glycine residues in the hydrophobic core of the signal peptide (the H region; Fig. 2), which interfere with export via the Sec pathway and concomitantly facilitate aSec transport (27). Replacing one or more of the glycines with amino acids that increase the H region hydrophobicity or α-helicity (properties known to be important for the interaction of signal peptides with SecA and SecY), can re-route the protein, so that it is transported more readily by the general Sec system. Conversely, replacing one of the glycine residues with proline (G75P), which decreases the helical propensity, abrogates Sec transport but only partially interferes with aSec transport. Whether the H-region glycine residues affect trafficking by facilitating a direct interaction with an aSec component, or simply render the preprotein unrecognizable by the general Sec system (a “Sec avoidance” mechanism), is not yet known.

Perhaps the most unique aspect of aSec transport is that it requires a specific 24-residue segment known as the accessory Sec transport (AST) domain adjacent to the signal peptide (20). A leucine residue at the C-terminal end of the AST domain (L114) is especially critical for aSec transport. Replacement of L114 with the helix-breaking residues glycine or proline nearly abolishes transport by the aSec system, whereas replacement with hydrophobic residues such as isoleucine, valine, or phenylalanine has no adverse effect. Extensive mutational analysis of the AST domain indicates that it might form an amphipathic helix. Experimental evidence also suggests that the AST domain affects both the targeting of GspB to the SecY2 translocon and gating of the transmembrane channel, but precisely how the AST domain might facilitate these processes is undetermined.

Although the AST domain is essential for GspB transport, no component of the aSec system has been found to bind specifically to this region of the preprotein (25). We therefore chose to use site-specific photo-cross-linking, to capture proteins that may interact only transiently with the AST domain during transport. This powerful technique has been increasingly employed to monitor protein-protein interactions in vivo (28–30). The key to this method is the incorporation of a photactivatable amino acid at a single, selected site in a protein of interest, through a process of “amber suppression” (31, 32). In brief, this method involves the placement of a TAG stop codon at a specific site within a gene of interest, and the co-expression of a nonsense suppressor tRNA along with a unique amino-acyl tRNA synthetase. The latter is engineered to transfer an “unnatural” amino acid such as benzoylphenylalanine (BzF), which is provided in the culture medium. Once the BzF-labeled protein is expressed, it can be photoactivated and cross-linked to any closely-associated proteins by exposing the culture to 350–365 nm light. The amber codon suppression technique has been utilized in eukaryotes, E. coli and mycobacteria but, to our knowledge, it has not yet been used in any Gram-positive bacterial species. Because of the documented efficiency, and lack of toxicity, of amber suppression in E. coli, we opted to express the S. gordonii aSec system along with the amber suppression components in E. coli. The combined results indicate that the aSec system is functional in E. coli, and that trafficking of GspB<sub>B.g</sub>-36FLAG and variant preproteins recapitulates that in S. gordonii. In addition, results in both E. coli and S. gordonii indicate that the AST domain interacts specifically with SecA2 to facilitate transport.

**EXPERIMENTAL PROCEDURES**

**Cloning and Expression of the S. gordonii aSec Operon in E. coli—**The entire aSec operon was cloned from the S. gordonii chromosome by using a marker rescue strategy. A unique Swa restriction site exists just downstream of asp5 (Fig. 1) and a second Swa site was incorporated just upstream of gspB (736flag by using forward primer: 5′-AAAAATGCATTTAAATTAG-TAGAGGGGATTAC-3′ (Swa site is underlined) and reverse primer 96XR (20) to amplify the 5′- end of gspB. The resulting fragment was digested with NsiI and XhoI and used to replace the NsiI-Xhol fragment of pB736fla/96X (20). The plasmid (pB736flag/96X/NsiSwa) was used to transform the gspB deletion strain PS1740 (23) resulting in strain PS2163 (Fig. 1B). Chromosomal DNA was extracted from PS2163, digested with SwaI, treated with T4 DNA ligase, and then used to transform E. coli TOP10 cells. The 23 kb plasmid carrying gspB736flag and the entire accessory Sec operon (pSwaR) was stably maintained as long as the host strain was grown at or below 30°C. Attempts to manipulate the plasmid, such as selective deletions of one or more genes, generally led to re-arrangements. However, a stable Δasp1–3 derivative plasmid (pSwaΔKpn) was obtained by deletion of a KpnI fragment. Plasmids pSwaR and pSwaΔKpn were used to transform E. coli strain C43/DE3. To assess the expression, glycosylation and transport of GspB<sub>B.g</sub>-736FLAG strains were grown at 30°C for 18 h in LB containing chloramphenicol (15 μg/ml), diluted 5-fold into fresh medium, and then incubated 90 min at 30°C. Periplasmic proteins were obtained by osmotic shock (33). The glycosylation of GspB<sub>B.g</sub>-36FLAG was assessed by lectin blot analysis, using biotinylated succinyllated wheat germ agglutinin as described (34).

**Subcloning of the aSec Operon—**To facilitate the separate expression of GspB<sub>B.g</sub>-36FLAG and the aSec system, a Sall fragment spanning from the middle of gspB to codon 220 of gspB (i.e. a Sall-Swal-Sall fragment; Fig. 3) was subcloned into pCOLA-Duet (Novagen), generating the plasmid pCOLAαSec. The subcloned fragment in pCOLAαSec was found to have a 1 kb insertion of an IS10 mobile genetic element in gtfA. The insertion apparently did not have a polar effect on the expression of the downstream genes, since the glycosylation of GspB<sub>B.g</sub>-36FLAG (which requires the expression of both GtfA and GtfB) could be restored by supplying gtfA on a separate plasmid (data not shown).

To generate an otherwise isogenic plasmid that lacked asp1–3, attempts were made to similarly subclone the Sall fragment from pSwaΔKpn into pCOLADuet, or to delete the asp1–3 KpnI fragment from pCOLAαSec/ghtfA::IS10. For
unknown reasons it was not possible to recover the aSecΔasp1−3 fragment in pCOLAΔduet using either of these methods. However, it was possible to subclone the Sall fragment from pCOLAαSec/gftA::IS10 into pVg891 (35), and then delete the asp1−3 KpnI fragment from this construct, resulting in plasmids pVg891αSec/gftA::IS10 and pVg891αSecΔKpnI/gftA::IS10, respectively.

**Construction of pCOLA-pBpF**—A BamHI-XhoI fragment containing the amber suppression components was removed from pEVOL-pBpF and ligated to pCOLADuet that had been digested with BamHI and Sall. This plasmid was used instead of pEVOL-pBpF in experiments where the aSec components were expressed from pVg891αSec/gftA::IS10 or pVg891αSecΔKpnI/gftA::IS10, since pVg891 and pEVOL-pBpF have the same origin of replication and selective marker.

**Construction of Amber Codon Variants of gspB736flag and BdD**—The gspB736flag or BdD coding regions were amplified from plasmid pBflagR or pBflagRΔ145−597 (26), respectively, using forward primer 5′-GGAATTCATATG-TTTTTTAAACGTCAAAAGGTCT-3′ and reverse primer 5′-TAATTTTAAATTTAATTCTGTAGTCATCGTCATCCTTGTAG-3′. The PCR products were digest with Ndel and PciI, and then cloned in pCDFDuet (Novagen) that had been digested similarly. Amber codons were incorporated into the gspB736flag or BdD coding sequences using a two-stage PCR strategy. The presence of only the intended change in any segments generated by PCR was confirmed by DNA sequence analysis (Sequetech).

**Amber Codon Suppression and Photocross-Linking**—E. coli C43/DE3 was transformed by electroporation, using simultaneous combinations of two or three plasmids as indicated. Transforms were selected on medium containing chloramphenicol (15 μg/ml), spectinomycin (50 μg/ml), and/or kanamycin (50 μg/ml) as appropriate. Strains were grown at 37 °C for 18 h, diluted 5-fold into fresh medium, and then incubated at 37 °C for 90 min. BzF (Bachem) and IPTG (Allstar) were each added to 100 μl of the culture by transferring two 50 μl samples to a large petri dish (144 mm²) and exposing to UV light with occasional mixing for 20 min. The other half of the culture was not exposed to UV but was otherwise processed similarly. After centrifugation for 20 min at 3200 × g, cells were lysed by suspension in Celllytic reagent (Sigma) followed by sonication. GspB736flag or BdD cross-linked products were purified using anti-FLAG antigen (Sigma). The eluted proteins were separated by SDS-PAGE, stained with Coomassie, and then submitted for analysis by tandem mass spectrometry (Stanford PAN Lab). In some cases, cell lysates or purified cross-linked complexes were treated with 2 units (100 ng) of thrombin for 3 h at room temperature, and the thrombin-treated complexes were analyzed by Western blotting using either anti-FLAG or anti-SecA2 polyclonal antibodies.

**Construction of the GspB736FLAG/spAST Chimera**—Chromosomal DNA was extracted from S. pneumoniae strain TIG4. The putative AST domain of psrP was amplified by PCR, using the forward primer 5′-AGTACTCGAGAAGGATGCTTGGC-3′ and the reverse primer 5′-CCGGATCCGCTGAATTCTTTAGATATGCACC-3′. The PCR product was digested with XhoI and BamHI and then used to replace the XhoI-BamHI fragment of pB736flag/96X118B (20), to generate plasmid pB736flag/spAST. The NsiI-SpeI fragment spanning codons 1 to 296 of gspB736flag/spAST was then used to replace the NsiI-SpeI fragment containing the wild-type sequence in pS326B602 (36), to generate pS326B602/spAST, which was used to transform S. gordonii strain PS1226 (19). A double crossover between sequences in pS326B602/spAST and the chromosome of PS1226 to generate PS2812 (M99 5′-spAST::pB736flag/spAST::pB736flagC ΔsecA2) was verified by sequence analysis of the chromosomal DNA. A secA2 resistance marker just upstream of gspB does not affect the expression or transport of GspB (36).

**Construction of SecA2 Chimeras**—The secA2 chimera CH1 was generated and cloned in pMSP3535 using the same strategy previously described for construction of secA-secA2 chimeras (19). In brief, primer pairs spa2F (5′-AAGGATCCCCAGGAGTGCAGATTGAGCTAGG-3′) and IRA2R were used to amplify the pneumococcal secA2 NBD1, PPXD, and NBD2 regions, and primers IRA2F and 3553MCS were used to amplify the C-domain of S. gordonii secA2. The two first-stage PCR reactions were combined and amplified using primers spa2A2 and 3′A2, and the second-stage PCR product was then cloned in pMSP3535 (37). For CH2, primers spa2F and spPPXDR (5′-CATACACGTGCTCTAGGAGATATTTGACATG-3′) were used to amplify the pneumococcal secA2 NBD1 and PPXD regions. The PCR product was digested with BamHI and PmlI and then used to replace the corresponding fragment in pMSPsecA2 (a PmlI site is present at the 3′-flank of the PPXD region). To facilitate the construction of CH3, a silent mutation was first introduced at codon 222 of secA2 and spPPXDR (5′-ATCTGGCCGCC-CCKCGTGTGYAGTGC-3′) and reverse primer 5′-GACTGRA-CACGGMGGCCGCAGCGM-3′). The silent mutation generates a unique Ascl restriction site (underlined) in the 5′-flank of the PPXD (pM99secA2/asc). The pneumococcal secA2 PPXD region was then amplified using primers gpAsCF and the reverse primer spPPXDR. The PCR product was digested with
Ascl and PmlI, and used to replace the corresponding region of pM99secA2/asc. The resulting plasmids were used to transform the ΔsecA2 strains PS1226 and PS2812 (described above).

Analysis of GspB736FLAG Secretion from S. gordonii—Strains were grown at 37 °C for 18 h in Todd-Hewitt broth containing 60 μg/ml erythromycin. Proteins from S. gordonii culture supernatants or protoplasted cells were prepared and analyzed by Western blotting as described previously (20).

RESULTS

Functional Expression of the aSec System in E. coli—To assess the validity of studying the S. gordonii aSec system in E. coli, we first examined the transport of GspB736FLAG when co-expressed with the entire aSec system. It should be noted that attempts to assemble the aSec system by combining the individual genes into single or multiple plasmids were largely unsuccessful. However, it was possible to retrieve the entire operon with the native gene organization by using a marker rescue strategy (see “Experimental Procedures”). With the entire aSec operon present, GspB736FLAG became glycosylated and was transported to the E. coli periplasm (Fig. 1B, lane 1). Upon deletion of aspI−3, GspB736FLAG migrated with a higher apparent mass, which is indicative of more extensive glycosylation, and remained mainly in the cytosol (lanes 3 and 4). These results indicate that the S. gordonii aSec system can indeed function in E. coli.

The expression of BzF-labeled GspB736FLAG along with the entire aSec system was subsequently examined. Although UV-induced mass shifts were evident (data not shown), characterization of the cross-linked products tended to be complicated by the mass heterogeneity that results from carbohydrate modification (glycosylated GspB736FLAG ranges from 120 to 150 kDa). It was therefore more feasible to assess cross-linking to the AST domain in the absence of glycosylation, but it was then necessary to determine whether the preprotein features that affect trafficking of the non-glycosylated preprotein in S. gordonii were recapitulated in E. coli. As mentioned above, transport of GspB736FLAG via the general Sec system in S. gordonii is inhibited not only by the carbohydrate moieties associated with the mature region, but also by glycine residues in the H region of the signal peptide. In the absence of the S. gordonii aSec system and glycosyl transferases, some transport of non-glycosylated GspB736FLAG was evident, as determined by both cleavage of the signal peptide and the appearance of the mature protein in the periplasm (Fig. 2, lanes 1 and 2). The signal peptide appeared to be cleaved at the normal cleavage site (between residues 90 and 91), since the periplasmic form was ~10 kDa smaller than the unprocessed preprotein. Moreover, a variant of GspB736FLAG that has an alteration predicted to interfere with signal peptidase cleavage (A90R/E91P; reference 26), did not show the same mature protein, but instead appeared to be cleaved somewhat heterogeneously (lane 3). Substitution of one or more of the H-region glycine residues with residues predicted to increase the H region hydrophobicity or helical propensity resulted in more efficient transport (lanes 4−6), whereas substitution with proline nearly abolished transport (lane 7). These results indicate that, as in S. gordonii, the native hydrophobic core of the signal peptide is not optimal for transport by the general Sec system of E. coli, and that Sec transport can be blocked by placement of a proline residue in the H region.

The effects of signal peptide and AST mutations on transport of non-glycosylated GspB736FLAG in the absence or presence of the aSec system was also assessed (note that the expression of non-glycosylated GspB736FLAG in the presence of the aSec system was facilitated by a fortuitous non-polar insertion in gtfA, as described under “Experimental Procedures”). With the aSec system present, most of the wild-type GspB736FLAG underwent transport and processing of the signal peptide (Fig. 3, lane 5). Transport of the L114G variant appeared reduced by comparison (lane 6), but was not abolished. However, since GspB736FLAG/L114G was transported (albeit inefficiently) by the Sec system (lane 2), it was not possible to determine the extent to which the L114G substitution impacted aSec transport. We therefore assessed the effect of the AST domain mutation in combination with the signal peptide mutation G75P,

FIGURE 1. The S. gordonii accessory Sec system and preprotein substrates. A, GspB is the native, cell wall-anchored (CWA) substrate. SP, signal peptide; AST, accessory Sec transport domain; SRR1 and SRR2, serine-rich repeat regions 1 and 2, respectively; BR, the ligand-binding region. GspB736FLAG is a model aSec substrate that is freely secreted into the S. gordonii culture medium. Bdd is a derivative of GspB736FLAG that lacks residues 145−597 (26). B, map of the accessory Sec operon in S. gordonii PS2163, and expression of the aSec system in E. coli. The Swal restriction site upstream of gspB736flag was engineered in to facilitate rescue of the entire operon. The transport components are indicated by orange arrows, the subunit is indicated in green, and glycosyl transferases are indicated in yellow. The integrated plasmid pVA891 has a chloramphenicol resistance gene and pACYC origin for selection and replication, respectively, in E. coli (35). A 23 kb plasmid that corresponds to the circularly-ligated Swal chromosomal fragment (pSwR), or a Δasp1−3 derivative plasmid (pSwDΔKpn), was used to transform E. coli strain C43/DE3. Periplasmic proteins (peri) or whole-cell lysates (wcl) were analyzed by Western blotting with anti-FLAG, anti-SecB or anti-MBP (maltose-binding protein) as indicated. SecB (a cytoplasmic protein) and MBP are negative and positive controls for periplasmic proteins, respectively. Glycosylated forms of GspB736FLAG were detected by lectin blot analysis, using succinylated wheat germ agglutinin (not shown).
which selectively blocks Sec transport. As expected, the G75P variant was transported only when the aSec system was present (lane 7 versus lane 3). However, the G75P/L114G double mutant was not transported by either pathway (lanes 4 and 8). This indicated that the L114G mutation interferes specifically with aSec transport. The combined results demonstrate that trafficking of non-glycosylated GspB736FLAG in E. coli is similar to trafficking in S. gordonii and, importantly, that aSec transport is dependent on the preprotein AST domain.

**Cross-linking to the AST Domain of GspB**

We next examined cross-linking to the AST domain of non-glycosylated GspB736FLAG in the absence or presence of the aSec system. The photoactivatable amino acid BzF was initially incorporated at position 114, since studies in S. gordonii had indicated that bulky hydrophobic residues at this position did not negatively impact transport (20). In the absence of the aSec system, two major cross-linked products were evident, migrating at 105 and 120 kDa (Fig. 4, lane 2), corresponding to GspB736FLAG cross-linked to E. coli proteins with apparent masses of ~110 and 20 kDa, respectively. A minor cross-linked product migrating at 150 kDa, corresponding to GspB736FLAG and a 50 kDa E. coli protein, was also evident. When the aSec system was present, one additional cross-linked product migrating at 200 kDa was seen (lane 4), corresponding to a mass shift of ~100 kDa.

To optimize the yield of cross-linked products for purification and identification by mass spectrometry, we then examined cross-linking under a variety of conditions. Although the same three cross-linked products were consistently obtained, extensive fragmentation of GspB736FLAG within the ligand-binding domain occurred prior to and during purification (Fig. 4 and data not shown) making it difficult to track and recover the cross-linked protein by anti-FLAG affinity chromatography. The ligand-binding domain of GspB is highly susceptible to proteolytic degradation and is not essential for aSec trans-

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**FIGURE 2. Accessory Sec-independent transport of GspB736FLAG and selected variants in E. coli.** The amino acid sequence of the H and C regions of the signal peptide, along with the AST domain of GspB is indicated. The GspB736FLAG variants were expressed constitutively from pVA891 (26, 27), and were detected by Western blotting with anti-FLAG monoclonal antibodies. Lane 1, periplasmic proteins; lanes 2–8, whole-cell lysates.

**FIGURE 3. Transport of GspB736FLAG and selected variants in the absence or presence of the aSec system in E. coli (aSec–, or aSec+, respectively).** GspB736FLAG (wt) and variants with the substitutions L114G (LG), G75P (GP), and L114G/G75P (LG GP) were expressed constitutively from pVA891 (20, 27). The aSec components were expressed from pCOLAaSec/gtfA::IS10. Note that gtfA* is disrupted by an insertion sequence, so GspB736FLAG does not undergo glycosylation. Anti-FLAG monoclonal antibodies were used for the detection of GspB736FLAG by Western blotting of whole cell lysates.

**FIGURE 4. Cross-linking to position 114 in the AST domain of GspB736FLAG.** GspB736FLAG/L114TAG was expressed from pCDF, the BzF-specific amber suppression components (amino-acyl tRNA synthase and tRNA_CUA) were expressed from pEVOL-pBpF, and the aSec operon was expressed from pCOLAaSec/gtfA::IS10. GspB736FLAG was detected by Western blotting with anti-FLAG antibodies, and the full-length preprotein is indicated by a black arrow. Red asterisks indicate cross-linked complexes that occur in both the presence and absence of the aSec system. Red arrow indicates an aSec-specific cross-linked complex. The panel on the right shows a shorter exposure of the Western blot, in which the 105 kDa cross-linked complex is more readily visible.
port (Ref. 26). Additional studies were therefore carried out using a derivative of GspB736FLAG that lacks most of this region (GspB derivative D, or BdD; Fig. 1). The photo-activatable BzF was again incorporated at position 114 in the AST domain. UV-dependent cross-linked products of apparent masses of 55 and 70 kDa were obtained in the absence of the aSec system, and an additional cross-linked product migrating at 150 kDa was obtained when the aSec system was present (Fig. 5A). The observed masses of the three cross-linked proteins represent UV-induced mass shifts of 5, 20, and 100 kDa (since BdD has an apparent mass of 50 kDa), which is the same as those in the major cross-linked complexes seen with BzF-labeled GspB736FLAG (Fig. 4).

Cross-linking to BdD at three additional positions (94, 95, and 115) was also examined. BdD labeled at positions 94 and 95 showed UV-dependent cross-linking similar to that of BdD labeled at position 114, with products of apparent masses of 55 and 70 kDa generated in the absence of the aSec system (data not shown) and an additional 150 kDa product generated in the presence of the aSec system (Fig. 5B). BdD labeled at position 115 yielded cross-linked products of 55 and 70 kDa in both the absence and presence of the aSec system (data not shown and Fig. 5B, respectively), but no 150 kDa cross-linked complex was detected. Thus, labeling at positions 94, 95, or 115 in the mature region results in identical cross-linking patterns, whereas labeling at position 115 showed no aSec-specific cross-linked product.

Characterization of the Cross-linked Proteins—Since cross-linking to the 100 kDa protein was clearly dependent on the aSec system, the protein cross-linked to BdD/Q94BzF was purified and was analyzed by mass spectrometry. The proteins identified in the complex were BdD and SecA2. The 150 kDa complex generated by using BdD/L114BzF was also purified, and Western blot analysis indicated it also contained SecA2. Red asterisks indicate transport-dependent, pathway-independent cross-linked products.
Interactions with the AST Domain of GspB

prendiplasm, and occurs upon transport by either pathway. If so, the cross-linked complexes may actually contain mature BdD, and the masses of the two E. coli proteins are likely to be 15 and 30 kDa.

Dependence of the AST Interactions on Asp1–3—Our recent studies have indicated that two of the accessory Sec system components, Asp2 and Asp3, can bind GspB directly (25). We have proposed that these Asps might function as chaperones and, along with Asp1, facilitate the targeting of GspB to the SecA2/Y2 translocon. We therefore examined whether cross-linking of the AST domain to SecA2 and the two E. coli proteins occurred in the absence of Asp1–3. With the BdD/G75P variant labeled at position 94, the 55 and 70 kDa complexes were not detected in the absence of Asp1–3 (Fig. 6, lane 6). Moreover, there was little evidence of signal peptide cleavage, which confirms that the Asps are essential for aSec transport in E. coli. Thus, the results further support the possibility that formation of the 55 and 70 kDa complexes is dependent on transport to the periplasm. Surprisingly cross-linking to SecA2 was readily apparent. This indicated that the Asps were not necessary for preprotein targeting to SecA2.

Since G75P partially interferes with aSec transport (Fig. 3, lanes 5 and 7 and Ref. 27), we also examined cross-linking to the AST domain of BdD that has the wild-type H region. With BdD labeled at position 94, cross-linking to SecA2 was again readily detected in the absence of Asp1–3 (Fig. 6, lane 10). Cross-linking to SecA2 was likely the result of targeting to, but not transport by, the aSec system (as in lane 6), whereas cross-linking to the 15 and 30 kDa proteins was due to transport by the general Sec system (as in Fig. 5). In contrast, with BdD labeled at position 114, cross-linking to the two smaller proteins, but not SecA2, was detected in the absence of the Asps (lane 14). To re-examine the ability of BdD labeled at position 94 or 114 to cross-link to SecA2, we co-expressed the labeled preproteins with SecA2 alone. Again, cross-linking of SecA2 to BdD labeled at position 94 was readily detected, but cross-linking of SecA2 to position 114 was not (Fig. 7). Therefore, although SecA2 can bind the preprotein at position 94 independently of the Asps, cross-linking of SecA2 to position 114 of the AST domain (i.e. full engagement of the AST domain) requires the entire aSec system.

Interaction of the AST Domain with the PPXD Domain of SecA2—To identify the region of SecA2 that cross-linked to the AST domain at position 94 or 114, the 150 kDa complexes were treated with thrombin, which cleaves SecA2 at residue 490 (Fig. 8A) but does not cleave BdD. Western blot analysis of the resulting proteolytic fragments indicated that BdD was linked in both cases to the N-terminal two-thirds of SecA2 (data not shown). This region includes a domain similar to the preprotein binding domain (PPXD) of SecA, which has been implicated in the binding of both the signal peptide and mature region of preproteins (38–40). We reasoned that, if a highly specific interaction between the AST region and SecA2 was necessary for transport, it might be possible to detect such an export requirement in S. gordonii by simultaneously substituting the native AST domain of GspB and the AST-binding domain of SecA2 with a heterologous pair of regions.

When selecting candidate homologues for domain exchange, we noticed that the sequences of SecA2, as well as the putative AST domains of the SRR proteins, diverge fairly rapidly across species. For example, the SecA2 proteins from Streptococcus sanguinis SK36, Streptococcus pneumoniae TIGR4, and Streptococcus agalactiae COH1 show 91, 79, and 70% similarity (81, 61, and 49% identity), respectively, to that of S. gordonii M99. Although the S. sanguinis SecA2 can partially complement a secA2 mutant of S. gordonii, the S. pneumoniae SecA2 is unable to transport GspB (data not shown).

The putative AST domains vary comparably. A chimeric form of GspB736FLAG in which the AST domain was replaced with the corresponding region of the S. sanguinis homologue SrpA (GspB736FLAG/ssAST) was readily transported by S. gordonii (data not shown), whereas a preprotein chimera containing the putative AST domain of the pneumococcal homologue PsrP (GspB736FLAG/spAST) showed reduced transport (Fig. 8B, lane 2 versus lane 1). For the compensatory exchange experiments, we therefore chose to incorporate regions from the pneumococcal SecA2 into the S. gordonii SecA2, and then examined the ability of these SecA2 chimeras to transport either GspB736FLAG or GspB736FLAG/spAST. Two of the SecA2 chimeras (CH1 and CH2, Fig. 8A) were unable to transport either wild-type GspB736FLAG or the chimeric preprotein (data not shown). However, a third SecA2 chimera (CH3) that contains only the pneumococcal PPXD showed markedly reduced transport of the wild-type preprotein (lane 3 versus lane 1), but was better able to transport GspB736FLAG/spAST (lane 4 versus lane 3). These data support the possibility that the AST domain interacts with the PPXD of SecA2 to facilitate transport. However, since transport of the preprotein chimera by the SecA2 chimera was reduced, as compared with transport of the wild-
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Protein transport by the accessory Sec system of *S. gordonii* is distinctly different from transport via the general Sec system in that it requires a specific segment in the mature region of the preprotein, adjacent to the signal peptide, known as the AST domain. Our previous genetic analyses indicated that the AST domain of the streptococcal glycoprotein GspB might affect both targeting of the preprotein to the SecA2/Y2 translocon and gating of the SecY2 transmembrane channel. The studies presented here were undertaken to determine which aSec components might interact with the AST domain to facilitate these processes. With the preprotein labeled at position 94, 95, or 114 of the AST domain, cross-linking to SecA2 was detected. No cross-linking to SecA2 occurred when the preprotein was labeled at position 115, which is consistent with genetic data suggesting that position 114 is the C-terminal end of the AST domain.

In addition to SecA2, cross-linking to two *E. coli* proteins was detected. Cross-linking to these proteins only occurs if the labeled preprotein is transported (Figs. 4B and 5C), but the interactions are not aSec-specific. Cell fractionation experiments have indicated that these complexes are membrane-associated (data not shown), but sufficient quantities for identification of the constituent proteins have not yet been obtained. However, the combined results are consistent with the possibility that cross-linking to the two proteins occurs after the AST domain reaches the periplasm, and that the apparent masses of these proteins are therefore 15 and 30 kDa.

A notable outcome of these experiments is that no interaction with SecA was detected, even in cases where the labeled preprotein was transported by the general Sec system (i.e. in the absence of the aSec components, and when the preprotein carried the wild-type signal peptide H region). This is consistent with the fact that the AST domain is not required for, and does not interfere with, general Sec transport. This also implies that a more intimate or prolonged interaction between the AST domain and the translocon motor protein occurs during, or prior to, transport by the aSec *versus* Sec pathway, and that cross-linking between the AST domain and SecA2 does not merely reflect incidental contact.

One surprising result is that the interaction of SecA2 with the preprotein does not require Asp1–3, but is instead modulated by one or more of the Asps. That is, the interaction of SecA2 with the N terminus of the AST domain (position 94) did not require Asp1–3, whereas interaction with the C terminus (position 114) did. These results indicate that full engagement of the AST domain by SecA2 is dependent on one or more of the Asps, and help to explain why the Asps are essential components of the aSec system. The finding that SecA2 contacts position 94 (and presumably the adjacent signal peptide), but not position 114, in the absence of Asp1–3 suggests two possible roles for the Asps. Rather than targeting the preprotein to SecA2, the Asps might maintain the preprotein in a conformation in which the entire AST domain is accessible to SecA2, or the Asps may directly alter SecA2 such that the AST binding site is more exposed (Fig. 9). This latter possibility would be analogous to a proposed role for SecB in post-translational transport. In *E. coli*, the C-terminal SecB binding domain of SecA is thought to occupy or perhaps occlude a signal peptide binding site that resides within the preprotein binding domain of SecA (39). This “auto-inhibition” can be relieved by SecB interaction with the C-terminal domain as it delivers a preprotein to SecA. The Asps might similarly alter SecA2 conformation to unmask the AST binding site.

Although it was suspected that the AST domain might interact with SecY2 to facilitate gating (*i.e.* opening) of the transmembrane channel, cross-linking was detected only to SecA2 but no other aSec protein. This suggests that the previously-described “gating” and “targeting” functions of the AST domain may actually reflect two different interactions between the preprotein and SecA2. The two purported functions were based on the finding that some AST domain mutations could be compensated for by using a pre-gated, or partially opened, form of...
SecY2 (SecY2<sup>1382EN</sup>), whereas other mutations could not (20). The presumed gating mutations were generally alterations in the N-terminal end of the AST domain, whereas the presumed targeting mutations were generally located near the C-terminal end of the AST domain. Since cross-linking of SecA2 to position 94 of the AST domain occurs independently of aSec transport, whereas cross-linking of SecA2 to position 114 is correlated with aSec transport (Figs. 6 and 7), it is possible that the full engagement of the AST domain by SecA2 is analogous to the “trapping” step of general Sec transport initiation described by Gouridis et al. (41). The trapping event is a prolonged, high affinity interaction between mature regions of preproteins and SecYEG-bound SecA, and corresponds to an early translocation intermediate. In the general Sec system, this role of the mature domain is not sequence-specific, whereas aSec transport is strictly dependent on the AST domain.

Results of domain-swapping experiments in <i>S. gordonii</i> support the possibility that the AST domain interacts specifically with the PPXD domain of SecA2 to mediate transport. However, replacement of the <i>S. gordonii</i> SecA2 PPXD with the pneumococcal SecA2 PPXD did not fully restore transport of a chimeric preprotein that contained the putative pneumococcal AST domain. The most likely explanation for this is that, in addition to being precisely matched to the AST domain, the PPXD must be matched to the signal peptide. Alternatively, the SecA2 chimera CH3 might not optimally interact with another component of the aSec system such as SecY2. Indeed, the PPXD of SecA makes several contacts with SecY during transport (28, 30, 42). An additional possibility is that the heterologous AST domain does not interact properly with the GspB signal peptide. We are currently undertaking experiments to examine these possibilities directly.

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