Review

Glycosylation and biogenesis of a family of serine-rich bacterial adhesins

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Glycosylation of bacterial proteins is an important process for bacterial physiology and pathophysiology. Both O- and N-linked glycan moieties have been identified in bacterial glycoproteins. The N-linked glycosylation pathways are well established in Gram-negative bacteria. However, the O-linked glycosylation pathways are not well defined due to the complex nature of known O-linked glycoproteins in bacteria. In this review, we examine a new family of serine-rich O-linked glycoproteins which are represented by fimbriae-associated adhesin Fap1 of Streptococcus parasanguinis and human platelet-binding protein GspB of Streptococcus gordonii. This family of glycoproteins is conserved in streptococcal and staphylococcal species. A gene cluster coding for glycosyltransferases and accessory Sec proteins has been implicated in the protein glycosylation. A two-step glycosylation model is proposed. Two glycosyltransferases interact with each other and catalyse the first step of the protein glycosylation in the cytoplasm; the cross-talk between glycosylation-associated proteins and accessory Sec components mediates the second step of the protein glycosylation, an emerging mechanism for bacterial O-linked protein glycosylation. Dissecting the molecular mechanism of this conserved biosynthetic pathway offers opportunities to develop new therapeutic strategies targeting this previously unrecognized pathway, as serine-rich glycoproteins have been shown to play a role in bacterial pathogenesis.

Introduction

Protein glycosylation is recognized as one of the major post-translational modifications and plays important roles in cell–cell recognition, adhesion and intracellular sorting (Drickamer & Taylor, 1998). Protein glycosylation is common in eukaryotes. More recently it has become clear that protein glycosylation occurs in all three domains of life (Schmidt et al., 2003). Prokaryotes, including bacteria and archaea, are able to glycosylate proteins (Moen & Vanderleyden, 1997; Upreti et al., 2003). More than 70 bacterial glycoproteins have been reported so far. Some are well characterized in Gram-negative bacteria, and a number of glycosylated proteins have also been documented and studied in Gram-positive bacteria (Upreti et al., 2003). Most characterized glycoproteins are virulence factors of medically important pathogens. The glycan variations of a few bacterial glycoproteins are closely associated with bacterial infection (Banerjee et al., 2002; Hamadeh et al., 1995), suggesting that bacterial protein glycosylation plays an important role in pathogenesis (Moen & Vanderleyden, 1997; Szymanski et al., 2003; Szymanski & Wren, 2005). Thus, the protein glycosylation and biosynthesis pathways may be new targets for therapeutic and prophylactic measures.

The biogenesis of glycoproteins is mediated by multiple genes clustered in so-called ‘glycosylation islands’ (Arora et al., 2001; Power et al., 2003; Szymanski et al., 2003). There are two basic types of protein glycosylation systems: N-linked glycosylation (oligosaccharides are usually linked to the amide nitrogen of asparagine residues of the peptide chain) and O-linked glycosylation (oligosaccharides are usually linked to the hydroxyl group of serine or threonine residues of peptide chains) (Szymanski & Wren, 2005). These two mechanisms can be differentiated by the mode in which the oligosaccharides are transferred to proteins. In the N-linked protein glycosylation system, an oligosaccharide is preassembled on a lipid carrier before being transferred en bloc to selected asparagine residues of acceptor proteins by an oligosaccharyltransferase (Szymanski & Wren, 2005). The N-linked glycosylation pathway is well defined in Campylobacter jejuni (Wacker et al., 2002). The classic O-linked protein glycosylation mechanism involves the sequential transfer of glycans directly from nucleotide sugar precursors to acceptor proteins. Bacteria produce a diverse array of O-linked glycoproteins. Recently one O-linked glycosylation pathway was characterized in the Gram-negative bacterium Neisseria gonorrhoeae (Hegge et al., 2004), which shares numerous common biosynthetic genes with N-linked glycosylation pathways and does not fit the bill as a classic O-linked glycosylation pathway. The biosynthetic mechanism in most bacteria is still unclear. Here we summarize...
recent findings on the structure, genetics and function of serine-rich O-linked glycoproteins of streptococci and staphylococci, and propose a biosynthetic pathway that represents an emerging mechanism for the bacterial protein glycosylation system.

**Identification of serine-rich glycoproteins**

The first serine-rich glycosylated protein was identified from an oral streptococcal bacterium, *Streptococcus parasanguinis* FW213 (Wu et al., 1998; Wu & Fives-Taylor, 1999). *Streptococcus parasanguinis* expresses peritrichous fimbriae on the cell surface (Fig. 1). Initial screening of nonadherent mutants revealed that they lacked fimbriae (Fachon-Kalweit et al., 1985), suggesting the importance of fimbriae in bacterial adhesion. The protein responsible for the formation of fimbriae was identified by immunological approaches. A series of monoclonal antibodies were produced by immunizing mice with the wild-type FW213. Specific monoclonal antibodies (mAbs) F51, D10 and E42 that react with wild-type bacteria but do not react with the nonadherent bacteria were identified (Fachon-Kalweit et al., 1985; Stephenson et al., 2002) and used to detect bacterial adhesins. Interestingly, all these monoclonal antibodies label the bacterial fimbriae and recognize a 200 kDa protein, suggesting that the 200 kDa protein is a fimbriae-associated protein. Therefore we named the fimbrial protein as Fap1 (fimbriae-associated protein 1).

The identity of Fap1 was elusive until the protein was purified in 1998. The purification and characterization of the Fap1 protein helped isolate and clone the full-length *fap1* gene. Fap1 has been shown to be involved in bacterial fimbrial formation and adhesion, as a knockout mutant failed to form fimbriae and had reduced ability to adhere (Wu et al., 1998). Recently we characterized nonadherent mutants of *Strep. parasanguinis* FW213 generated by random chemical mutagenesis and found that all the mutants had defects in *fap1* or in genes related to Fap1 biogenesis (Zhou et al., 2008). These results demonstrate the importance of Fap1 in bacterial adhesion. The initial cloning and sequencing of the *fap1* gene proved to be challenging because *fap1* possesses extensive repetitive sequences. The sequencing of the large repetitive region was completed by using a transposon mutagenesis strategy which allows us to sequence one 400–600 bp repetitive region at a time and assemble them manually based on the exact transposition insertion sites (Wu & Fives-Taylor, 1999). The unusual repetitive nature of Fap1 was initially thought to be artificial; however, the discovery of Fap1-like proteins in other streptococci, such as *Streptococcus gordonii* strains M99 and DL1 (Bensing & Sullam, 2002; Takahashi et al., 2002; Vickerman et al., 2007), *Streptococcus cristatus* (Handley et al., 2005) and the sequenced genome of *Streptococcus pneumoniae* TIGR4 (Tettelin et al., 2001), has confirmed our findings and stimulated researchers’ interest in the area.

Fap1 was found to be glycosylated, as purified Fap1 stained poorly with the protein stain Coomassie brilliant blue but strongly with the carbohydrate stain periodic acid–Schiff’s reagent (Wu et al., 1998). Other serine-rich adhesins shown to be glycosylated include GspB of *Strep. gordonii* (Bensing et al., 2004), SrpA of *Streptococcus sanguinis* (Plummer et al., 2005) and SraP of *Staphylococcus aureus* (Siboo et al., 2008). The glycosylation status of serine-rich proteins recognized from other Gram-positive bacterial genomes is unknown. Given the fact the predicted serine-rich proteins are localized adjacent to glycosylation and secretion loci similar to the one initially identified from *Strep. gordonii* M99 (Bensing & Sullam, 2002), it is likely that other serine-rich proteins are similarly modified.

**Serine-rich glycoproteins are conserved in oral streptococci and many pathogenic streptococci and staphylococci**

The predicted Fap1 protein contains 2570 amino acid residues (Fig. 2). It has a long signal sequence (SS) at the N terminus, and a classic cell wall anchor domain (CW) at the C terminus. Two serine-rich repeat regions (RI and RII) flank a non-repeat region II (NRII). Another non-repeat region (NRI) is located between SS and RI. The two repeat regions contain SESVSESVSI repeats and constitute 80% of the entire amino acid sequence (Fig. 2). Fap1-like serine-rich proteins are also recognized from many other streptococci, staphylococci and lactobacilli. All serine-rich proteins contain two repeat regions; the amino acid sequences within the two regions are slightly different. The alternate amino acids are quite diverse. For instance,
Fap1 has a large number of glutamic acid (E) residues whereas GspB has alanine (A) residues instead (Table 1). Analysis of the Fap1-like proteins has revealed that all the Fap1-like serine-rich proteins are very acidic, with an isoelectric point (pI) below 6 (Table 2). Interestingly, the NRII regions of different Fap1-like proteins are diverse; they may be either acidic or basic. Most serine-rich proteins, like Fap1, have an acidic NRII region. Only a few Fap1-like proteins contain basic NRII regions. They include Strep. pneumoniae, Strep. gordonii, Lactobacillus johnsonii NCC533 and Staphylococcus haemolyticus. Paul Sullam and co-workers have proposed that the pI value of NRII may direct the binding of the serine-rich proteins to host receptors, based on their observations that basic NRII regions of Hsa and GspB of Strep. gordonii mediate bacterial adhesion to human platelets whereas the acidic NRII region of Streptococcus agalactiae fails to support the adhesion (Takamatsu et al., 2005a, 2006). However, as they have pointed out, SrpA, a serine-rich protein of Strep. sanguinis with an acidic NRII, also mediates sialic-acid-dependent bacterial adhesion (Plummer et al., 2005). Further, Fap1, another protein containing an acidic NRII region, does not mediate bacterial binding to human platelets at all. Together these results suggest that more complex mechanisms may be involved in NRII-mediated bacterial adhesion.

We have identified and characterized an eleven-gene cluster located adjacent to the fap1 locus of Strep. parasanguinis FW213. This gene cluster contains genes encoding glycosyltransferases, a nucleotide-sugar synthetase and genes involved in protein secretion (Wu et al., 2007a, b) (Fig. 3). A similar locus was initially identified from Strep. gordonii M99 (Bensing & Sullam, 2002) and has subsequently been shown to be important for glycosylation and biogenesis of GspB (Bensing et al., 2004a; Takamatsu et al., 2004a, b, 2005b). A search of completed and ongoing bacterial genome projects led to interesting findings: fap1-like gene clusters and related glycosylation and secretion loci were present in the genomes of oral streptococci, Strep. gordonii DL1 Challis (Vickerman et al., 2007), Strep. sanguinis (Xu et al., 2007) and Strep. cristatus (Handley et al., 2005), and in important human pathogens, including Strep. agalactiae (Glaser et al., 2002), Strep. pneumoniae (Tettelin et al., 2001), Staphylococcus aureus (Baba et al., 2002), Staphylococcus epidermidis (Zhang et al., 2003) and Staphylococcus haemolyticus (Takeuchi et al., 2005). Interestingly, this type of gene cluster is also found in some commensal bacteria, such as L. johnsonii NCC533 (NCBI genome site) and Streptococcus salivarius (Pombert et al., 2008). Collectively the gene cluster contains two regions. One core region is composed of seven genes: secY2, gap1–3 (designed as asp1–3 in other organisms), secA2, gtf

Table 1. Serine repeat sequences in the family of serine-rich proteins

| Species and strain (protein name) | Serine repeat sequence |
|-----------------------------------|------------------------|
| Strep. parasanguinis (Fap1)       | SESVSESVSI SESVSESVSI  |
| L. johnsonii NCC533 (LJ1711)     | SQSESI SESL/1)SEVSSESV |
| Strep. sanguinis Sk36 (SrpA)      | SASISA SASTSASTSASV   |
| Strep. cristatus CC5A (SrpA)      | SV(A/SV(/A))SQ SASAVSASQ |
| Strep. agalactiae 2603V/R (Srr-1) | SVSM SAST/(M)         |
| Strep. agalactiae J48 (Srr-2)     | SSS SESISSESVSN        |
| Staph. aureus (SraP)              | STST STSLSD            |
| Staph. aureus NCTC 8325 (SAOUHSC_02990) | STS SVSTS          |
| Staph. epidermidis ATCC 12228 (SE2249) | SSEEK SDNSASTSL   |
| L. johnsonii NCC533 (LJ_0391)    | SLS SL/(M)SNVSMSMSE   |
| Strep. gordonii M99 (GspB)        | SAS SASTSASTSASTSASV  |
| Strep. gordonii DL1 (Hsa)         | SA/(L/S) SAST/(E)SASTSASV |
| Strep. pneumoniae TIGR4 (PsrP)    | SASE/(T) SASTSASA/(E)  |
| Staph. haemolyticus JCSC1435 (SH0326) | STST STSVSD    |

Fig. 2. Schematic diagram of Fap1. SS, signal sequence; NRI and NRII, non-repeat regions I and II; RII and RII, repeat regions I and II; CW, cell wall anchor domain.
Table 2. pI values of serine-rich proteins

| Group   | Species and strain (protein name) | pI | NRII       |
|---------|----------------------------------|----|------------|
|         | Entire protein                   |    |            |
| Acid    | *Strep. parasanguinis* (Fap1)    | 3.26| 4.85 (179-505) |
|         | *L. johnsonii* NCC533 (LJ1711)  | 4.23| 4.61 (181-1412) |
|         | *Strep. sanguinis* Sk36 (SrPA)   | 4.96| 5.41 (242-446) |
|         | *Strep. cristattus* CC5A (SrPA)  | 5.6 | 6.21 (256-739) |
|         | *Strep. agalactiae* 2603V/R (Srr-1) | 4.94| 4.63 (200-641) |
|         | *Strep. agalactiae* J48 (Srr-2)  | 4.21| 4.53 (212-531) |
|         | *Staph. aureus* (SraP)           | 4.2 | 5.69 (231-751) |
|         | *Staph. aureus* NCTC 8325 (SAOUHSC_02990) | 4.04| 5.61 (229-751) |
|         | *Staph. epidermidis* ATCC 12228 (SE2249) | 4.03| 6.00 (263-790) |
| Basic   | *L. johnsonii* NCC533 (LJ_0391)  | 4.7 | 8.13 (232-361) |
|         | *Strep. gordonii* M99 (GspB)     | 3.96| 9.51 (237-603) |
|         | *Strep. gordonii* DL1 (Hsa)      | 4.15| 8.74 (221-449) |
|         | *Strep. pneumoniae* TIGR4 (PsrP) | 3.9 | 9.91 (122-379) |
|         | *Staph. haemolyticus* JCSC1435(SH0326) | 3.58| 9.25 (301-496) |

and gtf2 (gtfA and gtfB homologues); this core region is conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative glycosyltransferases. The number and identity of genes coding for putative glycosyltransferases are quite diverse, as depicted in Fig. 3. For instance, four glycosyltransferase-coding for putative glycosyltransferases are quite diverse, as glycosyltransferases. The number and identity of genes conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative conserved in every genome that contains Fap1-like serine-rich proteins (Fig. 3). The other region contains putative Table 2. pI values of serine-rich proteins

| Group   | Species and strain (protein name) | pI | NRII       |
|---------|----------------------------------|----|------------|
|         | Entire protein                   |    |            |
| Acid    | *Strep. parasanguinis* (Fap1)    | 3.26| 4.85 (179-505) |
|         | *L. johnsonii* NCC533 (LJ1711)  | 4.23| 4.61 (181-1412) |
|         | *Strep. sanguinis* Sk36 (SrPA)   | 4.96| 5.41 (242-446) |
|         | *Strep. cristattus* CC5A (SrPA)  | 5.6 | 6.21 (256-739) |
|         | *Strep. agalactiae* 2603V/R (Srr-1) | 4.94| 4.63 (200-641) |
|         | *Strep. agalactiae* J48 (Srr-2)  | 4.21| 4.53 (212-531) |
|         | *Staph. aureus* (SraP)           | 4.2 | 5.69 (231-751) |
|         | *Staph. aureus* NCTC 8325 (SAOUHSC_02990) | 4.04| 5.61 (229-751) |
|         | *Staph. epidermidis* ATCC 12228 (SE2249) | 4.03| 6.00 (263-790) |
| Basic   | *L. johnsonii* NCC533 (LJ_0391)  | 4.7 | 8.13 (232-361) |
|         | *Strep. gordonii* M99 (GspB)     | 3.96| 9.51 (237-603) |
|         | *Strep. gordonii* DL1 (Hsa)      | 4.15| 8.74 (221-449) |
|         | *Strep. pneumoniae* TIGR4 (PsrP) | 3.9 | 9.91 (122-379) |
|         | *Staph. haemolyticus* JCSC1435(SH0326) | 3.58| 9.25 (301-496) |

Serine-rich glycoproteins play important roles in bacterial adhesion, colonization and pathogenesis

Fap1 was identified as an adhesin in *Strep. parasanguinis* as it is responsible for bacterial binding to saliva-coated hydroxylapatite (SHA) (Wu et al., 1998), an in vitro tooth model. A 53 kDa salivary receptor has been identified, although the nature of the receptor is unknown (Wu et al., 2007b). Fap1 also mediates bacterial biofilm formation in vitro (Froeliger & Fives-Taylor, 2001). Glycosylation of Fap1 is involved in fimbrial assembly, bacterial adhesion and biofilm formation (Peng et al., 2008). It appears that the Fap1 polypeptide backbone is involved in bacterial adhesion whereas the protein glycosylation is responsible for the development of bacterial biofilm (Wu et al., 2007b). Recently we have analysed a variety of non-adherent chemical mutants; strikingly, all the mutants we characterized have defects in Fap1 or Fap1-related genes (Zhou et al., 2008), demonstrating that Fap1 is a key adhesion molecule. 

Both GspB and Hsa have been separately identified from *Strep. gordonii* strains M99 and DL1. Hsa was isolated as a polysaccharide receptor (Takahashi et al., 2002) whereas GspB was discovered by its ability to bind to human platelets (Bensing & Sullam, 2002). Hsa was later shown to be involved in binding to human platelets as well (Takahashi et al., 2006; Xiong et al., 2008). Importantly, inactivation of either hsa or gspB results in a reduction in *Strep. gordonii*-mediated infective endocarditis in a rat model (Takahashi et al., 2006; Xiong et al., 2008). The interaction of GspB with host components has been studied extensively; GspB not only binds to human platelets but also interacts with a variety of salivary components such as the low-molecular...
mass salivary mucin MG2 and salivary agglutinin (gp340) (Takamatsu et al., 2006). The binding specificity relies on sialic acid residues of the receptor molecules (Takamatsu et al., 2005a, 2006). A very recent study has demonstrated that Hsa mediates sialic-acid-dependent binding of \textit{Strep. gordonii} to HL-60 cells differentiated into monocytes, granulocytes and macrophages via known phagocyte receptors such as CD11b, CD43 and CD50 (Yajima et al., 2008). All the binding is mediated by basic NRII regions of GspB and Hsa. The ability of \textit{Strep. gordonii} to survive in polymorphonuclear leukocytes following adhesin-mediated phagocytosis has been postulated as an important virulence determinant for infective endocarditis (Young Lee et al., 2006). Identification of new host receptors for the bacterial adhesin supports the function of Hsa in bacterial infective endocarditis. Hsa provides a selective advantage for \textit{Strep. gordonii} to outcompete \textit{Strep. sanguinis} in binding to SHA in vitro, which provides a means for \textit{Strep. gordonii} to persist within the oral cavity, despite the greater natural prevalence of \textit{Strep. sanguinis} in plaque and saliva (Nobbs et al., 2007). Among many adhesins of \textit{Strep. gordonii}, Hsa is a major one that directs the bacterial adhesion to immobilized gp340 and epithelial cells (Jakubovics et al., 2005). The oral streptococcus \textit{Strep. cristatus} possesses

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**Fig. 3.** Conservation of serine-rich glycoproteins and proteins contributing to their biosynthesis and secretion in streptococci, staphylococci and lactobacilli. DNA sequences of \textit{fap1} and \textit{fap1}-like loci were collected from the related NCBI websites. Genes coding for serine-rich repeat proteins linked to secY2–gtf loci were designated as follows: \textit{fap1} of \textit{Strep. parasanguinis}; \textit{srr-1}, SAK\_1493 and \textit{srr-2} of \textit{Strep. agalactiae}; \textit{psrP}, SPCG\_1750 and SPH\_1885 of \textit{Strep. pneumoniae} TIGR4, CGSP14 and Hungary 19A-6 strains; \textit{fimS} of \textit{Strep. salivarius} ATCC 25975; gspB or hsa of \textit{Strep. gordonii}; SA2447 of \textit{Staph. aureus} N315; SE2249 of \textit{Staph. epidermidis} ATCC 12228; SH0326 of \textit{Staph. haemolyticus} JCSC1435; and LJ-0391 of \textit{L. johnsonii} NCC533. Arrows indicate the direction of transcription.
Fap1-like protein SrpA, which was believed to be involved in the assembly of bacterial surface fimbrial structures (Handley et al., 2005); however, its precise function has not been explored. Strep. sanguinis has the ability to bind to human platelets; serine-rich glycoprotein SrpA mediates the binding (Plummer et al., 2005) although it has not been possible to demonstrate its involvement in bacterial infective endocarditis (unpublished data). Strep. pneumoniae PsrP has been implicated in invasive diseases by epidemiological studies; PsrP deficiency slows the development of bacteremia and increases the mouse survival rate (Obert et al., 2006). PsrP exhibits a tissue tropism. It is required for bacterial persistence in the lungs but not for colonization in the nasopharynx or replication in the bloodstream during sepsis (Rose et al., 2008). The basic NRRI region of PsrP mediates bacterial adhesion to human alveolar type II pneumocytes, A549 cells (Obert et al., 2006; Rose et al., 2008). Two serine-rich proteins, Srr-1 and Srr-2, have been described for Strep. agalactiae strains 2603V/R and J48 (Samen et al., 2007; Seifert et al., 2006). Serotype III delta/epsilon-positive strains of Strep. agalactiae possess Srr-2. They are highly virulent in an in vivo mouse model of neonatal sepsis in comparison with Srr-2-negative serotype III strains (Seifert et al., 2006), suggesting that Srr-2 likely contributes to bacterial pathogenesis. The function of the other serine-rich protein, Srr-1, has also been investigated. Srr-1 can interact with human keratin 4 and promotes adherence to epithelial cells, suggesting that Strep. agalactiae has developed a new means to interact with the host (Samen et al., 2007). Staph. aureus SraP also binds to human platelets via its N-terminal NRRI domain, and mediates the pathogenesis of infective endocarditis in a rabbit model (Siboo et al., 2005). In a search of all 22 sequenced genomes of Staph. aureus (http://www.ncbi.nlm.nih.gov/sites/entrez?Db=genomeprj&Cmd=Search&Term ToSearch=txid1280[orgn]), we found that all sequenced strains contain SraP-like serine-rich proteins. Serine-rich glycoproteins Srr-1 and Srr-2 of Strep. agalactiae and their putative glycosylation and biosynthetic loci are also found in all eight sequenced genomes. These observations are in stark contrast to what is documented for Strep. pneumoniae. In all 34 sequenced genomes of Strep. pneumoniae (http://www.ncbi.nlm.nih.gov/sites/entrez?Db=genomeprj&Cmd=Search&Term ToSearch=txid1313[orgn]), only six genomes have SrpP-like open reading frames. The prevalence of SraP in Staph. aureus and Srr-1 and Srr-2 in Strep. agalactiae strains highlights their importance.

In summary, the serine-rich proteins are important in bacterial adhesion via their NRRI domains; they constitute a new family of bacterial lectins, as many of them bind to sialic-acid-containing surface molecules and play an important role in bacterial pathogenesis. Interestingly, predicted serine-rich proteins also exist in probiotic bacteria such as Lactobacillus johnsonii NCC533, Lactobacillus reuteri and Streptococcus salivarius ATCC 25975, members of the normal microbial community. Therefore, this important family of proteins may be also required for bacterial fitness and normal physiology.

Glycosylation and biogenesis of serine-rich glycoproteins

Both O- and N-linked glycan moieties have been identified in glycoproteins of Gram-positive bacteria. The platelet-aggregation-associated protein (PAAP) of Strep. sanguinis (Erickson & Herzberg, 1993) is a well-defined streptococcal O-linked glycoprotein. Fap1 and its homologues constitute an important family of O-linked streptococcal glycoproteins and present a new biosynthetic mechanism for protein glycosylation (Peng et al., 2008; Stephenson et al., 2002; Wu et al., 2007a, b). It has been reported that accessory Sec proteins SecA2 and SecY2 play an important role in export of serine-rich glycoproteins GspB and Fap1 (Bensing et al., 2005; Chen et al., 2004). As the contribution of glycan moieties and the signal sequence to export of serine-rich glycoproteins has been reviewed recently (Rigel & Braunstein, 2008), we will focus on mechanisms related to glycosylation and maturation of serine-rich proteins.

Fap1 is the first protein in the family determined to be glycosylated (Wu et al., 1998). The purified Fap1 protein possesses several monosaccharide residues including glucose, N-acetylgalcosamine (GlcNAc), N-acetylgalactosamine and rhamnose (Stephenson et al., 2002). The glycan linkage to the Fap1 peptide backbone is not defined; however, it appears to be O-linked for the following reasons. N-Glycanases fail to release the glycan chains from Fap1 although the N-linked consensus sequence is present in Fap1 (Stephenson et al., 2002). Furthermore, the putative O-linked amino acid residues exist in Fap1, and the glycan chains can be released from a recombinant Fap1 by a condition used to cleave O-linked glycan moieties selectively (Bu et al., 2008). The GspB protein (Bensing et al., 2004b) and other members of this family, SrpA of Strep. sanguinis (Plummer et al., 2005) and SraP of Staph. aureus (Siboo et al., 2008), all react with the GlcNAc-specific lectin sWGA, suggesting that adding GlcNAc residues to the peptide backbone is common. We have shown that the glycosylation of Fap1 is initiated by addition of a GlcNAc residue by two glycosyltransferases, Gtf1 and Gtf2, of Strep. parasanguinis. Inactivation of either gtf1 or gtf2 leads to the expression of unglycosylated 360 kDa Fap1 precursor A. Gtf1- or Gtf2-deficient mutants fail to produce mature Fap1. Furthermore, the interaction between Gtf1 and Gtf2 is required for Fap1 glycosylation (Bu et al., 2008). Thus, we conclude that the formation of the Gtf1 and Gtf2 protein complex is important for this first step of Fap1 glycosylation.

What is intriguing though, is that deletion of gtf1 and gtf2 homologues gtfA and gtfB in Strep. gordonii abolishes the expression of native GspB although GtfA and GtfB have been shown to be involved in the transfer of the sugar residue to a recombinant GspB of Strep. gordonii M99 in the cytoplasm (Takamatsu et al., 2004b). Our work has also demonstrated the intracellular accumulation of glycosylated Fap1 (Chen et al., 2004), which supports the
hypothesis that the protein glycosylation occurs in the cytoplasm. Phylogenetic analysis indicates that Gtf1 and Gtf2 homologues are highly conserved in a variety of streptococcal and staphylococcal species (Fig. 4). Furthermore we have shown that the Gtf1 and Gtf2 homologues from 

Strep. sanguinis and Strep. pneumoniae can transfer an oligosaccharide containing GlcNAc residues to the recombinant Fap1 molecule (Bu et al., 2008), further highlighting the functional conservation of the Gtf complex. Thus we conclude that the formation of the Gtf1 and Gtf2 complex is required for glycosylation of serine-rich proteins.

A cluster of 11 genes flanking the fap1 locus has been identified in Strep. parasanguinis (Bu et al., 2008; Wu et al., 2007a, b). Mutagenesis of the downstream genes such as secY2, gap1 and gap3 generated another distinct high-molecular-mass (HMM) 470 kDa Fap1 precursor which migrates slower than the Fap1 precursor A; we designated it Fap1 precursor B (Li et al., 2008; Wu et al., 2007a). The mature 200 kDa Fap1 reacts with all Fap1-specific peptide and glycan antibodies. Interestingly, the 470 kDa Fap1 precursor reacts with peptide-specific antibody (mAb E42) and one glycan-specific antibody, mAb D10, but not with the mature Fap1-specific mAb, F51, suggesting that this Fap1 precursor is incompletely glycosylated or improperly folded. Similar HMM protein precursors are also recognized in mutants of Strep. gordonii deficient in secY2, gap1.

![Gtf1 phylogenetic analysis](image)

**Fig. 4.** Evolutionary relationships of glycosyltransferases Gtf1 and Gtf2. The phylogenetic analysis was performed using MEGA4. The length of the branches reflects the number of amino acid changes between different glycosyltransferases, as indicated by the bar. Bootstrap values are shown at the nodes. Glycosyltransferase sequences were obtained by searching all bacterial genomes and protein databases available from NCBI and TIGR.
and gap3 homologues (Takamatsu et al., 2005b), indicating that production of such a precursor is common in the biogenesis of serine-rich glycoproteins.

The presence of two distinct Fap1 precursors in the different mutants prompted us to hypothesize a two-step glycosylation pathway in which Fap1 progresses from apparent masses of 360 kDa to 470 kDa to 200 kDa based on SDS–PAGE analysis (Fig. 5). The first glycosylation step is controlled by a two-component glycosyltransferase complex, Gtf1 and Gtf2 (Bu et al., 2008), and occurs intracellularly, while the second step requires several accessory secretion components including SecA2 (Chen et al., 2004, 2006, 2007), SecY2 (Wu et al., 2007a) and glycosylation-associated proteins Gap1 and Gap3 (Li et al., 2008). In support of such a processing pathway, we find that only the 200 kDa protein is readily detectable in mature fimbriae (Wu & Fives-Taylor, 2001).

The expression of three Fap1 proteins, one with undetectable glycosylation (360 kDa), one incompletely glycosylated or improperly folded (470 kDa) and one fully glycosylated (200 kDa), is reminiscent of experimental findings in another oral streptococcus, Strep. salivarius (Weerkamp et al., 1986a, b). Two antigens associated with fimbriae have been identified in Strep. salivarius, AgB and AgC. Both antigens are predominantly associated with the cell wall in wild-type cells. However, AgC (AgC\textsubscript{in}) accumulates in high amounts in the cytoplasmic fraction of mutants that lack AgC\textsubscript{w}, the cell-wall associated form. This HMM AgC\textsubscript{in} molecule resembles 470 kDa Fap1 precursor accumulated intracellularly in secY2, gap1 and gap3 mutants of Strep. parasanguinis. AgB, AgC\textsubscript{in} and AgC\textsubscript{w} have apparent molecular masses of 380, 488 and ~250 kDa, respectively. Both AgC\textsubscript{in} (488 kDa) and AgC\textsubscript{w} (~250 kDa) are associated with carbohydrate as evidenced by staining with periodic acid–Schiff’s reagent, whereas AgB (380 kDa) contains little or no carbohydrate, and may represent the unglycosylated polypeptide backbone. However, the relationships among these three antigens and the corresponding genes coding for these HMM proteins of Strep. salivarius have not been identified. These antigens may have a common origin and represent a system analogous to Fap1. Biochemical analysis has revealed the presence of Fap1-like glycoprotein in Strep. salivarius (Levesque et al., 2001). Identification of fap1-related protein glycosylation and secretion loci in Strep. salivarius supports this assumption (Pombert et al., 2008). Together these observations indicate that the proposed two-step glycosylation and secretion model is relevant to the biogenesis of the serine-rich glycoprotein of Strep. salivarius.

This two-step model supports the concept that the glycosylation is coupled to the secretion (Chen et al., 2004; Wu et al., 2007a) of the protein. Interestingly, a Gap1 homologue of Strep. gordonii has been hypothesized as an accessory secretion protein as it affects mature GspB secretion (Takamatsu et al., 2004a, b). A HMM pre-GspB, but not mature GspB, has also been found in the cell wall fractions isolated from strains carrying mutations in each of the accessory secretory genes (secY2, asp1, asp2, asp3 and secA2). Although the nature of the pre-GspB is unknown, it is possible that this protein precursor is incompletely glycosylated. These results indicate that the accessory Sec system is required not only for the efficient export but also for the proper processing of GspB. The protein glycosylation is likely to contribute to this process. In this regard, GspB export and maturation (glycosylation) also appear to be coupled. Study of the precise structural differences between pre-GspB and mature GspB would help resolve the issue whether the protein glycosylation is directly or indirectly responsible for the processing of GspB. We have evidence supporting the notion that Gap1 is a putative glycosyltransferase and regulates Fap1 glycosylation and maturation (Li et al., 2008). Therefore, we conclude that glycosylation and secretion are coupled. In mycobacteria, Sec-dependent protein secretion mediates O-mannosylation, an analogous process prevalent in eukaryotic O-linked protein glycosylation (VanderVen et al., 2005). Thus, glycosylation of bacterial proteins appears to resemble the analogous process in eukaryotic cells. In eukaryotic cells, the exogenously supplied or endogenously generated nucleotide sugar building blocks are imported into the secretory compartments of cells, including the lumen of the endoplasmic reticulum and the Golgi apparatus. Transfer of a given oligosaccharide to a protein scaffold is controlled by specific localization of glycosyltransferases and mediated by protein secretion (Hounsell et al., 1996). Although the mechanism of prokaryotic glycosylation is not well defined and bacteria do not have specified organelles to carry out this function, it is possible that bacteria have developed an analogous system (such as subcellular localization) to control the function of the protein glycosylation complex. The proposed Fap1

![Fig. 5. A model for the Fap1 glycosylation and biogenesis pathway.](image-url)
glycosylation pathway is a dynamic process. The conserved sequence features indicate that this coupled glycosylation and secretion action is likely to be a common theme in biogenesis of serine-rich proteins in streptococci and staphylococci. Study of this newly identified biosynthetic pathway for bacterial glycosylation and secretion mechanism may lead to design of new therapeutic antimicrobial agents targeting the pathway.

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