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Cellular aspects of the distinct M protein and SfbI
anchoring pathways in Streptococcus pyogenes
Cellular aspects of M protein and SfbI anchoring to *Streptococcus pyogenes* wall

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Summary

Wall-anchored surface proteins are critical for the *in vivo* survival of *Streptococcus pyogenes*. Cues in the signal sequence direct the membrane translocation of surface proteins: M protein to the septum, and SfbI to the poles. Both proteins are subsequently anchored to the wall by the membrane bound enzyme sortase A. However, the cellular features of these pathways are not fully understood. Here we show that M protein and SfbI are anchored simultaneously throughout the cell cycle. M protein is rapidly anchored at the septum, and in part of the cell cycle, is anchored simultaneously at the mother and daughter septa. Conversely, SfbI accumulates gradually on peripheral peptidoglycan, resulting in a polar distribution. Sortase is not required for translocation of M protein or SfbI at their respective locations. Methicillin-induced unbalanced peptidoglycan synthesis diminishes surface M protein but not SfbI. Furthermore, overexpression of the division regulator DivIVA also diminishes surface M protein but increases SfbI. These results demonstrate a close connection between the regulation of cell division and protein anchoring. Better understanding of the spatial regulation of surface anchoring may lead to the identification of novel targets for the development of anti-infective agents, given the importance of surface molecules for pathogenesis.
Introduction

*Streptococcus pyogenes* is an important human pathogen, responsible for 500,000 deaths per year worldwide (Carapetis *et al*., 2005). Infection ranges from a mild strep throat or skin infection, to severe invasive conditions, including toxic shock syndrome, septicemia, and necrotizing fasciitis or “flesh-eating disease”. Untreated infection may lead to sequela including rheumatic heart disease and glomerulonephritis (Cunningham, 2000). *S. pyogenes* employs an impressive array of wall-anchored virulence factors that are critical for its *in vivo* survival (Bisno *et al*., 2003, Marraffini *et al*., 2006, Nobbs *et al*., 2009). Wall-anchored surface proteins possess a conserved C-terminal anchor domain comprised of an LPXTG motif followed by a hydrophobic region and a few positively charged amino acids (Fischetti *et al*., 1990, Schneewind *et al*., 1992). The anchoring domain is recognized by sortase, a membranal transpeptidase, which cleaves the LPXTG motif between the threonine and glycine residues (Mazmanian *et al*., 1999), and connects the freed threonine to the peptidoglycan precursor lipid II (Perry *et al*., 2002, Marraffini *et al*., 2006).

While the biochemical aspects of the sorting reaction have been studied in detail, the spatial regulation of this process is not as well understood (Marraffini *et al*., 2006, Hendrickx *et al*., 2011, Spirig *et al*., 2011). Proteins are anchored to the wall of *S. pyogenes* at two distinct locations, the septum and the poles (Carlsson *et al*., 2006). Anchoring of M protein to the cell wall takes place exclusively at the septum (Cole & Hahn, 1962, Swanson *et al*., 1969), where newly anchored M protein localizes within areas of active lipid II export and wall synthesis (Raz & Fischetti, 2008). Since the
synthesis of new cell wall is restricted to the septum of *S. pyogenes*, septal anchoring leads to the coating of the entire cell surface with M protein. In contrast to M protein, SfbI (also known as protein F, or PrtF), which is a major fibronectin binding protein in certain streptococcal strains (Talay *et al.*, 1992, Hanski & Caparon, 1992), displays a polar distribution (Ozeri *et al.*, 2001).

The signal sequence directs surface proteins for translocation at their ultimate cellular location (Carlsson *et al.*, 2006, DeDent *et al.*, 2008). Signal sequences containing a YSIRK-G/S motif are targeted to the septum, while signal sequences lacking this motif are targeted to the poles. Although the YSIRK-G/S motif is required for efficient signal sequence processing (Bae & Schneewind, 2003), mutations in this motif do not affect targeting to the septum (Carlsson *et al.*, 2006, DeDent *et al.*, 2008). Surface proteins possessing a YSIRK-G/S motif are found in *S. pyogenes* (Carlsson *et al.*, 2006), *Staphylococcus aureus* (DeDent *et al.*, 2008) and *Streptococcus pneumoniae* (Tsui *et al.*, 2011), are rare in *Enterococcus faecalis* (Kline *et al.*, 2009), and are absent from *Listeria monocytogenes* (Bruck *et al.*, 2011).

*S. pyogenes* sortase A (SrtA), which is involved in the anchoring of both M protein and SfbI (Barnett & Scott, 2002), localizes to a number of membranal foci (Raz & Fischetti, 2008). These foci are preferentially associated with the division septum, near sites of active M protein anchoring, and are recruited to daughter septa at an early stage of the division cycle. The distribution pattern of *S. pyogenes* pilus-specific sortases, SrtB and SrtC (Barnett & Scott, 2002, Barnett *et al.*, 2004, Mora *et al.*, 2005), is unknown at present.
In this study we use deconvolution immunofluorescence microscopy to study the two anchoring pathways of *S. pyogenes*. We show that M protein and SfbI are anchored simultaneously throughout the cell cycle. The anchoring of M protein is restricted to the septum, and occurs simultaneously at the closing mother septum and the forming daughter septa at certain stages of the cell cycle. SfbI on the other hand, is anchored at large peripheral areas, and its gradual accumulation on peptidoglycan results in polar distribution. Sortase is not required for the correct localization of M protein and SfbI translocation. Methicillin-induced unbalanced peptidoglycan synthesis, disrupts the proper assembly of the septum, and results in a marked reduction in surface M protein, but not SfbI. Overexpression of DivIVA also disrupts the septum, and results in a decrease in surface M protein, but an increase in SfbI.

**Results**

**Localization of M protein and SfbI anchoring sites**

In an attempt to better understand the two anchoring pathways in *S. pyogenes*, we used 3D structured illumination microscopy (3D-SIM) to follow the anchoring of M protein and SfbI (Fig. 1A). As expected, M protein (red) covers the entire surface of log phase D471 cells, although at this high resolution the distribution appears somewhat irregular, while SfbI (green) is present at the poles. It is of note however, that while the majority of cells displayed a similar level of M protein fluorescence, the SfbI fluorescence varied greatly between different poles.

The cellular location of active M protein anchoring has previously been studied by digesting existing surface proteins with trypsin, and following the regeneration of
new M protein in medium without trypsin (Cole & Hahn, 1962, Swanson et al., 1969). We found that complete removal of SfbI required the use of pronase in addition to trypsin, both of which were included in the growth medium (Fig. 1B, no regeneration). These proteases cannot cross the S. pyogenes cell wall, and their effect is therefore limited to the outer surface of the bacterium. Following a wash and 2 minutes regeneration in medium without proteases, M protein was detected strictly at the septum, while SfbI was detected in patches over a relatively large peripheral area (Fig. 1B, 2 minutes regeneration). A Z-stack view detailing the distribution of M protein and SfbI through the different layers of a representative streptococcal chain is presented in Figure 1C.

**M protein and SfbI are anchored simultaneously throughout the cell cycle**

To gain further insight into the anchoring of M protein and SfbI, we followed the localization patterns of these proteins as a function of the cell cycle stage. For this purpose D471 cells were treated with proteases as described above, washed, and resuspended in medium without proteases for two minute, allowing the regeneration of surface proteins. Numerous DeltaVision images were processed as average-intensity 3D-projections. A population of 781 cells was obtained by analyzing all the cells whose growth axis was parallel to the slide (and thus the septum was perpendicular to the slide), and that had no signal interference from other cells. Each cell was assigned to one of 6 groups based on cell length. While this division is arbitrary, it provides useful information about cell populations in different stages of the division cycle. Since S. pyogenes grows in chains, a clear-cut distinction of the
end of one cell cycle and the beginning of another is not immediately apparent. The first division stage in our analysis was chosen in a manner roughly analogous to that described by Higgins and Shockman (Higgins & Shockman, 1976), in which most cells reveal only preliminary peptidoglycan assembly at the forming septum, and where the mother septum is often not completely closed. A plot profile was generated for each cell, displaying the fluorescent signal intensity relative to the cellular position along the growth axis (see experimental procedures section). Whenever the chain orientation made it possible to determine which pole represented the previous division site (often for stages 1-4, but less often for stages 5-6), that pole was aligned to the left. A representative cell for each division stage, as well as its individual plot profile, are presented (Fig. 2A, left and middle columns). The mean intensity and standard deviation values for each cellular position among all the cells in the group were calculated, and these values are presented as population plots (Fig. 2A, right column). The experiment was repeated two more times, with smaller cell populations, yielding comparable results (not shown). The major cellular regions referred to in this study are presented in Figure 2B.

The population plots show that M protein and SfbI are anchored throughout the cell cycle. At the earliest stage (stage 1), M protein is observed primarily at the closing mother septum (here presented on the left). As the cell cycle progresses (stages 2-4), M protein anchoring activity is progressively redistributed to the forming daughter septum, and newly anchored M protein can often be observed simultaneously at both mother and daughter septa. Eventually, M protein is primarily anchored at the daughter septum (stage 5-6). SfbI on the other hand is anchored at a
broad peripheral region. In early division (stage 1), SfbI is preferentially anchored at
the pole distal to the previous division site. As division progresses (stages 2-4) SfbI is
anchored at the old pole as well as the inter-septal region. At the late stages of
division (stages 5-6) SfbI is anchored at both poles. Active division septa consistently
show the least amount of SfbI anchoring.

To test in a more direct manner whether M protein and SfbI are anchored
simultaneously to the surface of individual cells, the total M protein fluorescence of
each cell in the population was plotted against the total SfbI fluorescence of the same
cell (Fig. 2C). Only relatively few cells were found on the upper-left portion of the
plot (representing high M protein fluorescence but no SfbI fluorescence), and the
lower right portion (representing high SfbI fluorescence but no M protein fluoresce).
Although the cells presented a wide range of fluorescence intensities, most fell in the
middle region, indicating that both proteins were anchored. While these data
represent the outcome of 2 minutes of protein anchoring activity rather than real-time
anchoring, the prevalence of cells displaying both proteins, as well as the anchoring
of the two proteins in all stages of the division cycle, indicate that the two proteins are
anchored in parallel throughout the cell cycle.

M protein is anchored simultaneously at the closing mother septum, and forming
daughter septa

Ovococci are a group of bacteria with slightly elongated coccus morphology,
which divide in a single plain, and often form chains of organisms (Zapun et al.,
2008). This situation makes the overlap of two successive division cycles possible,
with the initiation of daughter septa assembly before the mother septum is completely
closed (Higgins & Shockman, 1976, Gibson et al., 1983). Early examinations of the
anchoring of M protein to the cell wall of *S. pyogenes* following trypsinization and 15
minutes of protein regeneration suggested that simultaneous anchoring at the mother
and daughter septa is possible (Cole & Hahn, 1962). To study the possibility of
simultaneous anchoring at these locations in more detail, we used population-level
analysis. While our initial population-level analysis is in agreement with
simultaneous anchoring (Fig. 2A), this method averages the fluorescence signal
across the cells in the group, and is therefore inadequate for this purpose.

We therefore developed a method for automatically analyzing the distribution of
M protein in individual cells. To minimize the possibility that sequential anchoring at
the mother and then daughter septa would be interpreted as simultaneous anchoring,
protease treated cells were suspended directly in medium without proteases and
allowed the regeneration of surface proteins for only 30 seconds before fixation
(omitting the one-minute wash step, in which some protein anchoring does occur).
Each cell in a population of 1,039 was assigned to one of six groups according to
length, and its M protein distribution plot was generated as described above. The
resulting plots were analyzed individually using MATLAB (see experimental
procedures section), and assigned to one of four categories: M protein anchored at the
mother septum alone, the daughter septum alone, both septa, or no septal anchoring
(Fig. 3A). To supplement the automated analysis, the entire population was also
classified into similar categories by direct observation of the cells (Fig. 3B). While
the manual method classified slightly more cells as presenting simultaneous
anchoring at both septa, the two methods were generally in good agreement. The majority of cells in the youngest cell population (stage 1) displayed M protein only at the mother septum. With the progression of the cell cycle (stages 2-4) anchoring activity was gradually redistributed to the daughter septum, with roughly 20-30% of stage 2 cells, 30-40% of stage 3 cells, and 20-30% of stage 4 cells displaying M protein at both mother and daughter septa. At the later division stages (stages 5-6) M protein was found almost exclusively at the daughter septum. The analysis of M protein anchoring was repeated with an additional population of 890 cells, resulting from four different cultures, yielding comparable results (Fig. S1A). Additionally, this same analysis was applied to the cell population described in Figure 2, which was allowed two minutes of protein regeneration (Fig. S1B). As expected, this population showed a higher occurrence of M protein detected at both mother and daughter septa. This increase may be attributed in part to the brighter fluorescent signal, yet with longer regeneration time, some sequential anchoring at the mother followed by the daughter septa, cannot be ruled out.

M protein anchoring following 30 seconds regeneration was also visualized using 3D Structured Illumination Microscopy (3D-SIM). For this purpose the cells were labeled with M protein specific antibodies (green), and a vancomycin Rhodamine red conjugate (red), which preferentially labels sites of active peptidoglycan synthesis (Daniel & Errington, 2003). Sites of M protein anchoring localized to regions strongly labeled with vancomycin Rhodamine red. In cells displaying both mother and daughter septa, M protein was often observed at both locations (Fig. 3C). These results suggest that in rapidly dividing S. pyogenes cells, simultaneous anchoring of
M protein to the mother and daughter septa during the initial stages of daughter septum formation is a common occurrence.

**Polar distribution results from gradual accumulation of SfbI on peripheral peptidoglycan**

As discussed above, although SfbI was anchored throughout the peripheral region of the cell, the final distribution was distinctly polar, and displayed great variations in fluorescence intensity between different poles. One hypothesis that may explain how diffuse SfbI anchoring could lead to the observed distinctly polar distribution is that SfbI is anchored at a relatively constant pace, and accumulation of more SfbI on older poles leads to the observed difference in fluorescence intensity. To test whether there is a correlation between the amount of SfbI found on the pole and its age we used a fusion protein between Green Fluorescent Protein (GFP) and the binding domain of the phage lysin PlyC (GFP-PlyC/BD) in a “pulse-chase” type experiment. The binding domain of the PlyC lysin binds tightly to the cell wall carbohydrate (Nelson et al., 2006), making it recognizable by its GFP fluorescence. D471 cells were grown in TH+Y media to OD\textsubscript{600} 0.15, at which stage purified GFP-PlyC/BD was added to the medium for 30 minutes (‘pulse’). Cells examined at this time point were brightly fluorescent in the green channel (Fig. 4A, ‘before chase’). The cells were then washed and incubated for one hour in medium lacking this fusion protein (‘chase’), at which time the green fluorescence was limited to poles that were already formed during the ‘pulse’ (Fig. 4A, ‘1 hour chase’). These poles, which are at least two generations old, showed the most SfbI labeling, while younger poles showed considerably less SfbI
labeling. The prevalence of SfbI on old poles labeled with GFP-PlyC/BD is apparent when the two fluorescent signals are examined side by side (Fig. 4B).

To quantify the relations between the age of the pole and the amount of SfbI fluorescence, the signal distributions of SfbI and GFP-PlyC/BD were analyzed in a population of 714 cells, derived from two separate experiments (each experiment supported the final result, when analyzed individually). Each cell was divided into two regions: a “young pole” consisting of the “mother septum” and “inter-septal” regions (Fig. 2B), and an “old pole” equivalent to the “pole” region (Fig. 2B). The signal at the “daughter septum” region at the middle of the cell (Fig. 2B) was not analyzed, and this region served as buffer between the two poles. For each pole, the average SfbI and GFP-PlyC/BD fluorescence were determined, and the two values were plotted against each other (Fig. 4C). Young poles (blue) were not yet formed at the time of the GFP-PlyC/BD “pulse” and thus did not display substantial GFP-PlyC/BD fluorescence. The majority of these poles displayed only a modest amount of anchored SfbI. Old poles (red) represent a heterogeneous group with ages ranging from one to several generations. The majority of one-generation-old poles was created during the “chase” period, and thus displayed only modest GFP-PlyC/BD fluorescence. Poles that are two or more generations old were already formed during the “pulse” period, and thus displayed a high GFP-PlyC/BD fluorescence. Since the growth of *S. pyogenes* is not synchronized, a third group of poles are those that were only partly formed during the “pulse”, and thus displayed an intermediate level of GFP-PlyC/BD fluorescence. The average fluorescence values for the poles in each of these four groups are presented in Figure 4D. Both direct examination (Fig. 4C), and
comparison of the average SfbI fluorescence associated with each age group (Fig. 4D), reveal a direct correlation between the level of SfbI fluorescence and the age of the pole.

To supplement the “pulse-chase” results, D471 cells were grown in medium containing trypsin and pronase as described above, washed, and incubated in medium without proteases for 50 minutes to regenerate surface proteins. Since M protein is only anchored to newly synthesized peptidoglycan at the septum, areas devoid of M protein represent poles that were already fully formed at the time of protease treatment, and are therefore at least two generations old (Fig. 4E). When the labeling intensity of SfbI in these cells was determined, the “oldest” poles (with the least M protein labeling) displayed the most SfbI labeling, poles that were one generation old displayed weaker labeling, and newly formed poles displayed very weak SfbI labeling (see Fig. 7 for a model representation). The combined results of the two approaches suggest that gradual accumulation of SfbI on peripheral peptidoglycan, leads to the observed polar distribution, with older poles displaying a greater amount of SfbI. These observations also suggest that poles remain active sites of SfbI anchoring for at least two generations following their formation.

Translocation of M protein and SfbI at distinct locations is maintained in the sortase mutant AR01

Our next aim was to better understand the mechanisms underlying the differences between M protein and SfbI anchoring patterns. We first tested whether deletion of sortase A, which anchors both proteins to the cell wall (Barnett & Scott, 2002), could
affect their translocation pattern. The lack of M protein and SfbI anchoring to the cell wall of the sortase mutant strain AR01 was validated by fractionation and Western blot analysis (Fig. S2A). Although no anchoring took place, both proteins were translocated across the plasma membrane, and remained trapped to some extent in the cell wall. M protein and SfbI accumulated at distinct locations on the surface of untreated AR01 cells, and did not substantially colocalize (Fig. S2B). Regeneration of surface proteins following protease treatment revealed that M protein and SfbI first became visible on the surface of AR01 in cellular locations resembling that of wild type D471, namely M protein was translocated at the septum and SfbI at the cell periphery. A Z-stack view of a representative streptococcal chain is presented in Figure S2C. These results indicate that sortase is not required for spatially correct translocation of surface proteins.

Unbalanced peptidoglycan synthesis induced by methicillin results in a marked reduction in the cellular amount of M protein but not SfbI

At the initial stages of protein sorting, a nascent surface protein is cleaved near the C-terminus and covalently attached to lipid II through a sortase-mediated transpeptidation reaction. The ensuing complex serves as substrate for penicillin binding proteins (PBPs), resulting in covalent attachment of the surface protein to the cell wall. PBPs assemble peptidoglycan through two distinct activities, namely the polymerization of glycan chains through transglycosylation, and the cross-linking of these chains through transpeptidation. β-lactam antibiotics, which inhibit the transpeptidation activity but not the transglycosylation activity of PBPs, do not
directly affect the sorting reaction (Ton-That & Schneewind, 1999). An interesting feature of \( \beta \)-lactam antibiotics is their ability to bind various PBPs with different affinities, resulting in unbalanced peptidoglycan synthesis (Williamson \textit{et al.}, 1980, Gutmann \textit{et al.}, 1981, Pucci \textit{et al.}, 1986, Lleo \textit{et al.}, 1990). In particular, methicillin was shown to specifically inhibit septal peptidoglycan synthesis in several ovococci, resulting in the formation of rod shaped cells (Lleo \textit{et al.}, 1990, Perez-Nunez \textit{et al.}, 2011). We reasoned that if similar morphological effects could be induced in \textit{S. pyogenes}, the manner in which M protein and SfbI are anchored during unbalanced peptidoglycan synthesis could shed light on the differences between the two anchoring pathways.

\textit{S. pyogenes} D471 cells were grown in the presence of trypsin and pronase to digest existing surface proteins until OD\textsubscript{600} 0.5 was reached, at which time the culture was diluted 1:4 into a similar medium containing ascending concentrations of methicillin. Following one-hour growth, during which time the phenotypic effects of methicillin on peptidoglycan synthesis became apparent, the cells were washed and resuspended in medium containing a similar concentration of methicillin, but lacking proteases. Following 10 minutes of surface protein regeneration, the cells were fixed and the anchoring patterns of M protein and SfbI were determined by immunofluorescence (Fig. 5A). Morphological defects, resulting from unbalanced peptidoglycan synthesis, became apparent at 0.2 \( \mu \)g/ml methicillin, as many cells displayed an elongated or rod-shaped morphology with multiple septa. At higher methicillin concentrations the cells became bulbous with no obvious septa. In rod-shaped cells, M protein was typically anchored at several septa per cell, albeit at a
reduced quantity compared to untreated cells, while SfbI was anchored both at the poles and at the inter-septal regions. At higher methicillin concentrations the amount of M protein on the cells was diminished substantially, while significant anchoring of SfbI was still observed. Western blot analysis of fractionated cells revealed a similar pattern, where the amount of M protein was greatly reduced at methicillin concentrations above 0.2 µg/ml, while the amount of SfbI remained substantial (Fig. 5B). At 3 µg/ml methicillin, the level of SfbI anchoring dropped as well, however this reduction may be due to a general decline in cellular functions. To verify that methicillin did not directly interfere with the sorting reaction, cell cultures treated in a similar manner were harvested, and boiled in 2% SDS. The SDS supernatant was collected (Fig. 5C, “SDS-soluble”), and the cells were lysed using the phage lysin PlyC, thereby releasing covalently bound surface proteins (Fig. 5C, “SDS-insoluble”). Treatment with methicillin did not result in the release of substantial quantities of M protein or SfbI following boiling in SDS, indicating that the covalent attachment of surface proteins by sortase was not affected. The anchoring pattern of M protein and SfbI on methicillin-induced rod-shaped cells was further studied by 3D-SIM (Fig. 5D). These images show in greater detail the anchoring of M protein at multiple septa per cell, and the anchoring of SfbI, not only at polar regions, but also at inter-septal regions. 
Overexpression of the cell division protein DivIVA results in aberrant cellular morphology, a reduction in the level of M protein, and an increase in SfbI.

DivIVA plays a major role in the regulation of septum placement in Gram-positive organisms. *Bacillus subtilis* DivIVA is targeted to areas of negatively curved membrane at the poles and the septum, recruits MinC and MinD to the poles, and thus prevents the assembly of polar division rings. In the absence of DivIVA, division is severely inhibited and septa are misplaced (Cha & Stewart, 1997, Edwards & Errington, 1997, Marston & Errington, 1999, Lenarcic *et al.*, 2009, Ramamurthi & Losick, 2009). *S. pneumoniae* and *E. faecalis* do not possess a MinCD system, nevertheless, DivIVA is important for the correct placement of the septum through a mechanism that is not completely understood (Fadda *et al.*, 2003, Fadda *et al.*, 2007, Ramirez-Arcos *et al.*, 2005). We postulated that overexpression of *S. pyogenes* DivIVA could similarly interfere with septum placement, and thus provide an additional way to test the dependence of protein anchoring on a functioning septum.

To test the effects of DivIVA overexpression, three plasmids carrying a spectinomycin (spec) selectable marker were constructed: pAR291_DivIVA, which expresses DivIVA under the control of the highly active M protein promoter, pAR287_GFP-HT, which expresses a GFP-HaloTag fusion protein under the control of the same promoter (unrelated protein control), and pAR161, an empty vector. For microscopy studies, overnight cultures of D471 harboring pAR291_DivIVA and pAR161 were diluted 1:50 into TH+Y+spec medium containing trypsin, and pronase. At OD$_{600}$ 0.5, the cells were washed, incubated in medium without proteases for 5 minutes, and the anchoring of M protein and SfbI was examined. Cells harboring...
pAR291_DivIVA had a reduced growth rate, and displayed aberrant morphology, including enlarged volume and irregular shape (Fig. 6A). The amount of M protein associated with these cells was greatly reduced compared to control samples. When observed on cells, M protein was still associated with the septa despite the aberrant morphology. SfbI was found in wall-associated patches, and displayed no clear cellular localization preference, although the exact distribution pattern was difficult to ascertain due to the irregular shape of these cells. For Western blot analysis, overnight D471 cultures harboring pAR161, pAR287_GFP-HT, or pAR291_DivIVA, were diluted 1:50 into TH+Y+spec medium, and were fractionated upon reaching OD$_{600}$ 0.5. The amount of M protein in cells harboring pAR291_DivIVA was greatly reduced compared to cells containing pAR161, or pAR287_GFP-HT (Fig. 6B). Conversely, the amount of SfbI was increased in cells containing pAR291_DivIVA. The reduction in the level of M protein was not due to the introduction of the M protein promoter on the DivIVA expression plasmid, since cells harboring pAR287_GFP-HT, which possess a similar promoter, did not display altered M protein expression level. When examined by fluorescent microscopy, cells harboring pAR287_GFP-HT were brightly fluorescent in the green channel, confirming the functionality of the M protein promoter used in this study (not shown). It is of note that disruption of proper septum assembly by methicillin or the overexpression of DivIVA, both resulted in a marked reduction in the level of surface M protein.
Discussion

Anchoring of surface proteins to the cell wall of streptococci and staphylococci is divided into septal and peripheral anchoring pathways (Carlsson et al., 2006, DeDent et al., 2008). We used S. pyogenes M protein and SfbI as a model system to better understand the relations between the division cycle and protein anchoring through the two pathways, in an attempt to reach a more unified view of these processes. We took two general approaches: the first was largely descriptive and followed the two anchoring pathways throughout the cell cycle using population-level analysis, while the second tested how interference with the proper assembly of the division septum affects protein anchoring. We found that M protein and SfbI are anchored simultaneously throughout the cell cycle. M protein anchoring is strictly limited to the septum and occurs simultaneously at the mother and daughter septa at certain stages of the cell cycle. Conversely, SfbI is anchored in patches throughout the peripheral peptidoglycan, with some preference for the poles, and its accumulation over time results in polar distribution. Figure 7 summarizes these observations in a model form.

We also found that sortase A is not required for localized translocation of the two proteins. Methicillin-induced unbalanced peptidoglycan synthesis, disrupts the proper assembly of the septum, and results in a marked reduction in surface M protein, but not SfbI. Overexpression of DivIVA also disrupts the septum, and results in a decrease in surface M protein, but an increase in SfbI.
The relations between the anchoring of surface proteins and the division cycle

The observed overlap in the anchoring of M protein at the mother and daughter septa of *S. pyogenes* is in agreement with a model proposing the simultaneous synthesis of peptidoglycan at these locations. Electron microscopy studies of cell division in enterococci revealed that initiation of peptidoglycan synthesis at future division sites begins before the mother septum is completely closed, and is independent of the completion of chromosome replication (Higgins & Shockman, 1976, Gibson *et al.*, 1983). The localization pattern of the *S. pneumoniae* division factors FtsZ (Morlot *et al.*, 2003), FtsA (Lara *et al.*, 2005), and FtsW (Morlot *et al.*, 2004), often shows simultaneous labeling at the mother and daughter septa. The labeling pattern of fluorescent vancomycin, which preferentially labels regions of lipid II export and cell wall synthesis, also shows simultaneous localization at the mother and daughter septa of both *S. pneumoniae*, and *S. pyogenes* (Daniel & Errington, 2003, Ng *et al.*, 2004, Raz & Fischetti, 2008). The penicillin binding proteins PBP1b, PBP2a, PBP2b, and PBP2x, all show simultaneous localization at the mother and daughter septa of *S. pneumoniae* at certain stages of the cell cycle (Morlot *et al.*, 2003, Morlot *et al.*, 2004, Zapun *et al.*, 2008). Examination of the locations where M protein is regenerated following protease digestion, has previously been used to study septal peptidoglycan synthesis (Cole & Hahn, 1962). While this method does not provide real-time results, in this work we have reduced the regeneration time to the bare minimum of 30 seconds, and analyzed the anchoring pattern of this protein in large populations of cells. Our finding that M protein is anchored simultaneously at
the mother and daughter septa, strongly supports the model of simultaneous peptidoglycan synthesis at these locations in *S. pyogenes*.

In contrast to M protein, SfbI is anchored in patches to large peripheral regions of the peptidoglycan, with preference for the poles. The polar distribution of SfbI on *S. pyogenes* cells is the result of two processes: preference for anchoring at non-septal regions, and gradual accumulation on preassembled peptidoglycan, which results in a correlation between the amount of SfbI anchored and the pole’s age. Our results also demonstrate that SfbI is actively anchored at the poles for at least two generations following their formation. Expanding on this basic model, it should be noted that the expression level of non-YSIRK-G/S proteins might also affect their observed distribution. A low expression level may result in substantial labeling only on poles that are far apart, while a high expression level may result in the anchoring of a substantial quantity of proteins even on relatively young poles.

**Protein anchoring and the localization of sortase**

M protein and SfbI are both anchored to the cell wall by sortase A (Barnett & Scott, 2002). We previously studied the distribution pattern of sortase in *S. pyogenes* and found that it localizes to a number of membrane-bound foci in each cell. Consistent with the anchoring pattern of M protein described here, sortase foci are preferentially associated with division septa, are recruited to the forming septa at an early stage, and are often present at both mother and daughter septa simultaneously (Raz & Fischetti, 2008). Despite their relative abundance at the division septa, sortase foci are not strictly confined there, and some foci are regularly seen at other cellular
locations. One way of explaining the apparent bias in sortase distribution towards septal anchoring is that the streptococcal septum is constantly being split to form peripheral peptidoglycan, which can no longer facilitate the anchoring of YSIRK-G/S proteins, and therefore the time-window available for septal anchoring is very limited. The need for efficient anchoring at the septum is met through the coupling of protein anchoring to septal peptidoglycan synthesis, which ensures a high level of lipid II, PBPs, and possibly other septal factors. Localized translocation of M protein at the septum is also likely to play an important role in ensuring efficient anchoring (Carlsson et al., 2006). The higher prevalence of sortase foci at the septum is therefore likely to play a role in mediating efficient anchoring at this location. SfbI on the other hand, is anchored gradually over a relatively large cellular area, and its anchoring may therefore require a lower concentration of sortase.

The model presented here for the anchoring of surface proteins to the cell wall of \textit{S. pyogenes} differs substantially from the model proposed for \textit{Streptococcus mutans} (Hu et al., 2008) and \textit{E. faecalis} (Kline et al., 2009). These studies showed that sortase A colocalizes with SecA in a single membranal microdomain termed ExPortal. The ExPortal, first described in \textit{S. pyogenes}, is a membranal microdomain enriched in anionic lipids, in which the secretion related ATPase SecA and the membranal protease/chaperone HtrA localize, and facilitates the secretion and maturation of the streptococcal secreted protease SpeB (Rosch & Caparon, 2004, Rosch & Caparon, 2005, Rosch et al., 2007). A different study however, found that SecA is distributed throughout the streptococcal membrane (Carlsson et al., 2006), and the reason for this difference is not clear. Recently, \textit{S. pneumoniae} SecA, and
SecY were found to be dynamically localized to both the septum and periphery suggesting the absence of an ExPortal in this organism (Tsui et al., 2011). *S. pneumoniae* sortase A displays a punctate pattern, but does not show preferential distribution to the septum (Tsui et al., 2011). Further study is needed to fully understand the relation between protein translocation and anchoring in *S. pyogenes*, as well as the molecular mechanisms underlying the distinct anchoring patterns observed in different ovococi.

### The effects of septum disruption on protein anchoring

To better understand the mechanisms governing the anchoring of M protein and SfbI to the wall of *S. pyogenes*, we interfered with two processes: protein sorting, and septum assembly. The deletion of sortase did not alter the cellular locations where M protein and SfbI were translocated, indicating that sortase does not play a role in determining the site of translocation. Conversely, interference with the placement of the septum through the use of methicillin or the overexpression of DivIVA resulted in an increase in cell size, a reduction in the number of septa, and a reduction in the amount of M protein found at the cell surface. The level of SfbI was only little changed following methicillin treatment, and was increased when DivIVA was overexpressed. These data suggest that a functioning septum is critical for the anchoring of YSIRK-G/S-type proteins. When septum formation is prevented, the anchoring of YSIRK-G/S-type proteins does not become delocalized but rather, these proteins are absent from the bacterial surface. The mechanism underlying this phenomenon is currently under investigation. While inhibition of M protein...
anchoring by the overexpression of DivIVA was linked to severe deformities in cellular morphology, the possibility that DivIVA is involved in the regulation of surface protein anchoring in a more direct manner should not be completely ruled out. Of particular note is that S. pneumoniae DivIVA is localized to both the division septum, and to a certain extent, the poles (Fadda et al., 2007).

One interesting phenomenon was the formation of rod-shaped cells displaying multiple septa, following exposure to an intermediate concentration of methicillin, similar to the effect observed in Lactococcus lactis (Perez-Nunez et al., 2011). In contrast to L. lactis and most other ovococci however, S. pyogenes lacks PBP2b, RodA, MreC and MreD homologues, which are important for peripheral peptidoglycan synthesis in ovococci (Zapun et al., 2008). This distinct peptidoglycan synthesis mechanism adds cell wall material at the splitting septum, and is responsible for the slightly elongated shape of many ovococci. Formation of S. pyogenes rod-shaped cells suggests therefore that the septal mechanism for peptidoglycan synthesis may be sufficient to facilitate the coccus-to-rod transformation. The specific molecular mechanisms involved in this alteration of S. pyogenes shape however, are not fully understood at present.

The formation of rod-shaped S. pyogenes cells provided us with an interesting model, in which to test the relations between surface protein anchoring and cell shape. We found that M protein was regularly anchored simultaneously at multiple septa along the rod, which were analogous to mother and daughter septa in untreated cells. Inhibition of septum closure therefore intensified the propensity to simultaneously anchor M protein at these locations. We also found that SfbI was not
anchored solely at the poles of the rod, but was also anchored at the cylindrical inter-
septal regions. This suggests that SfbI is not necessarily targeted to the poles as such, 
but rather, its anchoring pattern may best be described as exclusion from the septum. 
Interestingly, a distribution pattern of exclusion from the septum was also observed 
for the *S. pneumoniae* D,D-carboxypeptidase PBP3 (Morlot *et al.*, 2004). Studying 
the manner in which such a distribution pattern is achieved may provide clues for the 
regulation of SfbI translocation. Further insight into the manner by which non-
YSIRK-G/S-type proteins are anchored may come from studies dealing with rod-
shaped bacteria such as *L. monocytogenes* (Bierne *et al.*, 2004, Rafelski & Theriot, 
2006, Bruck *et al.*, 2011). A direct comparison however, may be complicated by the 
different manner peptidoglycan synthesis is regulated in these organisms. 
Recently, a screen of *S. aureus* transposon integration library revealed that 
disruption of proteins containing an abortive infectivity (ABI) domain results in 
reduced expression level of YSIRK-G/S type surface proteins (analogous to *S. 
pyogenes* M protein), but does not affect the expression of proteins that do not contain 
this motif. Interestingly, these mutants also display thicker septal peptidoglycan, and 
it is not clear whether the reduction in septal protein expression is a direct effect or 
the result of defects in septal peptidoglycan synthesis (Frankel *et al.*, 2010). 

**Conclusions**

When considering the two distinct anchoring pathways, each appears to have its 
own unique characteristics and advantages. Septal anchoring allows efficient coating 
of the entire surface of the cell from the moment the peptidoglycan is formed.
Immediate and extensive coating is likely to be important for the proper function of M protein and possibly other septum-anchored proteins. Consider that anchoring of M protein through the peripheral anchoring pathway may result in large areas of the streptococcal cell wall being devoid of this molecule at any given time. In the absence of M protein and other such molecules, these areas may be subject to opsonization, leading to the elimination of these bacteria through phagocytosis (Perez-Casal et al., 1992). Peripheral anchoring of SfbI on the other hand has the unique characteristic of creating cellular polarity. The extent to which the polar localization of virulence factors is important for the pathogenic process and survival of S. pyogenes is not clear at present. It was shown however, that polar distribution of SfbI resulted in reciprocal clustering of integrins on the surface of host cells (Ozeri et al., 2001), and future studies are likely to uncover additional examples. The ends of a streptococcal chain may have a better opportunity for contact with the host, and thus a higher concentration of certain virulence factors at these locations may be advantageous. Polar localization of proteins is a recurring theme in Gram-negative and Gram-positive bacteria, and has been shown to play a role in the general functions of the bacteria, as well as in pathogenesis (Shapiro et al., 2002, Shapiro et al., 2009, Rudner & Losick, 2010).

The data presented here suggest that factors related to the division ring are likely to play a role in promoting the anchoring of M protein. Over the recent years, the function of many of these division factors has been elucidated (Zapun et al., 2008, Adams & Errington, 2009, Shapiro et al., 2009, Rudner & Losick, 2010), and this knowledge may facilitate direct examination of their importance for surface protein
Defining the factors critical for the proper regulation of protein sorting could yield novel targets for the development of anti-infective agents, since pathogens lacking surface proteins are greatly impaired in their ability to cause disease (Marraffini et al., 2006, Maresso & Schneewind, 2008).

**Experimental procedures**

**Bacterial strains and culture conditions.** *Escherichia coli* strain DH5α was used for molecular cloning and recombinant protein expression. *S. pyogenes* strain D471 (an M6 serotype) was from the Rockefeller University collection. AR01 is a *srtA* knockout strain derived from D471 (Raz & Fischetti, 2008). *E. coli* was grown in Luria-Bertani (LB) medium, supplemented with 100 µg/ml ampicillin, when needed. *S. pyogenes* strains were grown in Todd-Hewitt medium (Difco) supplemented with 1% yeast extract (Fisher Scientific) at 37°C. Erythromycin was used at 15 µg/ml for *S. pyogenes*, and spectinomycin was used at 20 µg/ml for *E. coli* and 120 µg/ml for *S. pyogenes* when appropriate.

**Reagents and antibodies.** The M protein specific 10B6 monoclonal antibody (Jones et al., 1985) was used at a 1:10,000 dilution for Western blot analysis, and 1:1000 for immunofluorescence. SfbI-specific rabbit serum and pre-immune serum (Molinari et al., 1997), were used at a 1:1000 dilution. Goat anti-mouse IgG conjugated to either Rhodamine red (Jackson ImmunoResearch), or FITC (Sigma), were used at 1:1000. Goat anti-rabbit IgG, conjugated to either FITC (Sigma) or Alexa Fluor 647 (Invitrogen) were used at 1:1000. Wheat germ agglutinin (WGA)
Marina Blue conjugate (Invitrogen) was used at 5 µg/ml. DAPI (Sigma) was used at 1 µg/ml. Vancomycin was conjugated to NHS Rhodamine red according to manufacturer’s instructions (Thermo Scientific), and separated from unbound dye by thin layer chromatography. GFP-PlyC/BD (see below) was used at a final concentration of 50 µg/ml. All other reagents were purchased from Sigma unless otherwise noted.

**DNA manipulation.** Standard procedures were used for DNA manipulation and for *E. coli* transformations (Sambrook *et al.*, 1989). Transformation of *S. pyogenes* was performed according to Perez-Casal (Perez-Casal *et al.*, 1991). Plasmid DNA was isolated using QIAPrep spin miniprep kit (Qiagen). PCR amplification procedures were performed using either Vent or Phusion DNA polymerase (New England Biolabs). Oligonucleotides were from Eurofins. Restriction enzymes were from New England Biolabs. T4 DNA ligase was from Invitrogen.

**Construction of the plasmids.** For the construction of the GFP-PlyC/BD expression vector, pAR159, the binding domain of the PlyC phage lysin was amplified using the following primers: 5_PlyC-BD_XbaI (5’-GGCTCTAGATGAGCAAGATTAATGTAAACGTAGAAAATG-3’), 3_PlyC-BD_PstI (5’-CGCCCTGCAGTTACTTTTCATAGCCTTTCTGATAGCC-3’), and inserted into the XbaI and *PstI* sites of pBAD24 (Guzman *et al.*, 1995). Primers 5_H6_GFP_EcoRI (5’-CGCGAATTCTGAGTAAAGGAGAATCTCATCATTCCATTCCCTCCGACCAGTAAG-3”) and 3_GFP_KpnI (5’-GAGGATCTTTTGATAGTTCCATGCC-3”)
3’) were used to amplify the GFP_mut2 gene (Cormack et al., 1996). An N-terminal hexahistidine tag is encoded on the upstream primer. This PCR product was inserted into the EcoRI and KpnI sites of the above plasmid, yielding pAR159.

A modified version of the S. pyogenes shuttle vector pLZ12-spec (Husmann et al., 1995) was constructed by replacing the original multiple cloning site (MCS) between the EcoRI and SphI sites, with a DNA fragment formed by aligning primers 5_new_plzMCS (5’-AATTCCCAAGCTTCCCCAGATCTAAACCGGAAACAGCTGAAAC-3’) and 3_new_plzMCS (5’-CTTTCCATGGTTTCAGCTGTTTCCGC-GGTTTAGATCTGGGAAGCTG-3’), which contain the new MCS (EcoRI-HindIII-BglII-SacII-PvuII-NcoI-SphI). The resulting plasmid, termed pAR161, was used for the construction of pAR287_GFP-HT and pAR291_DivIVA.

For the construction of pAR287_GFP-HT, The HaloTag gene was amplified from pFN18A HaloTag T7 Flexi Vector (Promega) using primers 5_HaloL1_SacII (5’-CCCCCCGGGTGCATCTGCCCAGGCAGAAATCGGTACTG-3’) and 3_Halo_NcoI (5’-CCCCCCCATGGCTATCAGCCGAAATCCTGGACGTC-3’), and inserted into the SacII and NcoI sites of pAR161. The GFP_mut2 gene (Cormack et al., 1996) was amplified using primers 5_GFP_BglII (5’-GAGAGATCTATGATAAGGAGAAGAACTTTTC-3’) and 3_GFP_SacII (5’-GAGCCGCGTTTGTATAGTTCATCCCATGCCC-3’), and inserted into the BglII and SacII sites of the resulting plasmid. Finally, the M protein upstream region was amplified from genomic D471 DNA using primers 5_Mp_UTR_EcoRI (5’-CGCGAATTCCACAGCTAGCCGCAAGAACTC-3’) and 3_Mp_UTR_BglII (5’-
CCCAGATCTGCTCCTATGTATCTATTTTTTAGG-3’), and inserted into the EcoRI and
BglII sites, yielding pAR287_GFP-HT.

The construction of pAR291_DivIVA begun with a derivate of pAR161
containing a myc-tag (that is not expressed in the final construct), formed by aligning
primers 5_myc_tag (5’-GGGAACAAAAACTTATTCTGAAGAAGACCTGTAGC-3’) and
3_myc_tag (5’-CATGGCTACAGGCTTTCTCAAGAAATAAGTTTTTGTTCCGCG-3’), and
inserting the resulting double stranded DNA fragment into the SacII and NcoI sites
pAR161. The M protein upstream region was inserted into the EcoRI and BglII sites
of this plasmid as described for pAR287_GFP-HT. The DivIVA gene, including a
stop codon, was amplified from the genome of D471 using primers
5_DivIVA_BamHI (5’-CCCGGATCCATGGCACTTACACGCTAGAAATTAAAG-3’) and
3_DivIVA_SacII (5’-CCCGCGGTAGATATTTAATTTAAACGTTTGTGTTTCATGAG-
3’), and inserted into the BamHI and SacII sites of the resulting plasmid, yielding
pAR291_DivIVA.

Purification of GFP-PlyC/BD. An overnight culture of E. coli DH5α harboring
pAR159 was diluted 1:100 into one liter of LB supplemented with ampicillin, grown
to OD₆₀₀ 0.5, and induced with 0.2% L-arabinose at room temperature for 5 hours.
The construct was purified on a NiNTA column as previously described (Raz &
Fischetti, 2008). The eluted fraction was concentrated using an Amicon Ultra
centrifugal filter device with a cutoff limit of 5 kDa, and the buffer was changed to
PBS by repeated cycles of dilution in PBS and volume reduction. The final protein
concentration was 0.85 mg/ml.
**Fractionation of *S. pyogenes* cells and Western blot analysis.** Fractionation of the cells, and Western blot analysis were carried out as previously described (Raz & Fischetti, 2008). Samples were normalized to account for slight variations in OD. Fractionation by boiling in SDS was carried out as follows: One milliliter of culture at OD$_{600}$ 0.5 was harvested and washed with 30 mM Tris, pH 6.3. The bacterial pellet was suspended in 50 µl of 2% SDS, boiled for 10 minutes, and centrifuged for 2 minutes at 16000 rcf. The supernatant, containing non-covalently bound proteins, was supplemented with 12 µl of 5× SDS loading buffer. The cell pellet, containing covalently anchored proteins, was washed with 200 µl deionized water, suspended in 50 µl of 30 mM Tris, pH 6.3, containing 300 U/ml PlyC (Nelson *et al.*, 2006) for 15 minutes at room temperature, and then supplemented with 12 µl 5× SDS loading buffer.

**Regeneration of surface proteins following protease treatment.** For protein regeneration studies, overnight cultures were diluted 1:100 in TH+Y containing 0.35 mg/ml trypsin (Sigma) and 0.04 mg/ml pronase (Sigma). Unless otherwise noted, the cells were harvested upon reaching OD$_{600}$ 0.5 by one-minute centrifugation at 16,000 rcf, resuspended in TH+Y without proteases, and immediately spun again. The cells were immediately resuspended in medium without proteases, and incubated at 37°C for the stated amount of time. Each experiment typically consisted of three to four repeats, and was performed several times.
Fluorescent microscopy. Fluorescent microscopy procedures were carried out as previously described (Raz & Fischetti, 2008); however, the membrane and cell wall permeabilization steps were omitted since the antigens studied are exposed on the bacterial surface. Immunofluorescence experiments involving the use of methicillin or the overexpression of DivIVA were performed using polyclonal SfbI sera pre-adsorbed with the M1 serotype *S. pyogenes* strain SF370, which lacks a *sfbI* gene (Ferretti *et al.*, 2001). In those cases the cells were incubated sequentially with the M-protein-specific mouse monoclonal 10B6, anti-mouse Rhodamine red, rabbit anti-SfbI, and anti-rabbit FITC antibodies, for improved signal over background. Structured Illumination microscopy was performed on a DeltaVision OMX Blaze 3D-Structured Illumination Microscopy (3D-SIM) system (Applied Precision) fitted with an Olympus 100x/1.40 NA UPLSAPO objective, Photometrics Evolve EMCCD cameras, and 405, 488 and 568 lasers. 3D-SIM reconstruction and channel alignment were performed using the SoftWoRx algorithms and reconstructed images were exported as maximum projections.

Analysis of protein anchoring using MATLAB. For the purpose of signal analysis, average intensity projections were produced containing the signal data from all the Z-sections, using SoftWoRx (Applied Precision). These projections were converted into tiff format using ImageJ (http://rsb.info.nih.gov/ij/), and the signal distribution data were subsequently analyzed using MetaMorph offline (64-bit) version 7.7.5.0 (Molecular Devices). In each image, all the cells whose growth axis paralleled with the slide (and therefore the septum was perpendicular to the slide),
and that presented no signal interference from adjacent cells, were analyzed. Each cell
was confined in a rectangle whose long dimension parallels with the cell’s growth
axis. To analyze the fluorescence intensity of the antigens in different cellular
regions, each rectangle was then sub-divided into pixel-wide cross-sections, and the
total fluorescence of each section was calculated. These raw fluorescence distribution
data were further analyzed using MATLAB version 7.6.0 R2008a (MathWorks). To
create cell-specific plots the fluorescence intensity of each section was plotted as a
function of its cellular position, represented as the distance from the younger of the
two poles, or mother septum. Population plots were obtained by aligning all the cells
in each group, and calculating the mean fluorescence signal and standard deviation
values for each cellular position.

For the analysis of M protein anchoring at the mother and daughter septa,
presented in Figure 3, each cell was divided into 4 regions: (1) mother septum,
defined as the 0.13 µm region (3 pixels wide) adjacent to the previous division site
(on the left); (2) daughter septum, defined as the 0.21 µm region (5 pixels wide) at the
middle of the cell; (3) inter-septal region, located between the mother and daughter
septa; and (4) old pole, defined as the remainder of the cell (see Figure 2B). A mother
septum region was defined as positive for the anchoring of M protein if it passed a
threshold of 200 fluorescence units, and displayed double the minimal signal of the
inter-septal region. A daughter septum region was defined as positive for M protein
anchoring if, in addition to the above conditions, it also displayed double the minimal
signal of the polar region. In the youngest cell population, resolution constraints
sometime resulted in protrusion of the mother septum fluorescent signal into the inter-
septal region, preventing the achievement of the two-fold fluorescence difference required for the recognition of mother septum anchoring. To address this situation, cells that were negative for M protein anchoring using the above definitions, but passed the fluorescence threshold and displayed 5-fold mother septum fluorescence compared to the old pole, were defined as positive for mother septum anchoring.

For the analysis of SfbI signal distribution following the GFP-PlyC/BD “pulse-chase” experiment, the signal distribution of the different fluorescent channels was acquired as described above. A 5-pixel-wide region in the middle of the cell was left unanalyzed, while the regions on both sides were defined as “young pole” and “old pole” according to the GFP-PlyC/BD distribution along the chain. The average fluorescence intensity of each channel was calculated for each pole. The code used for all the MATLAB operations is available upon request.

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Abbreviations

WGA – Wheat Germ Agglutinin.
PBP – Penicillin Binding Protein.

RFU – Relative Fluorescence Units.

3D-SIM – 3D Structured Illumination Microscopy.

References:


Figure 1. The relative anchoring patterns of M protein and SfbI. An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y (A), or TH+Y containing trypsin and pronase (B), and fixed upon reaching OD$_{600}$ 0.5 (A, and B “no regeneration”). Protease treated cells were also harvested at OD$_{600}$ 0.5, washed, and resuspended in TH+Y for 2 minutes at 37°C prior to fixation (B “2 minutes regeneration”, and C). Specific antibodies were used to label M protein (red) and SfbI (green). The cell wall was stained with WGA marina blue (blue). 3D-SIM microscopy images are presented as maximum intensity projections composed of all the Z-sections (A and B), or as sequential Z-sections (C).
Figure 2. M protein and SfbI are anchored simultaneously throughout the cell cycle. (A) An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y containing trypsin and pronase and incubated at 37°C to OD$_{600}$ 0.5, at which point the cells were washed, and resuspended in TH+Y without proteases for 2 minutes at 37°C prior to fixation. The cells were labeled for M protein (red), SfbI (green) and wall (WGA, blue). Average intensity 3D-projections were made for numerous DeltaVision images, resulting in a population of 781 cells. These cells were divided according to cell length to 6 groups, representing different stages of the cell cycle. Each cell was divided into pixel-wide strips, parallel to the division plane, and the total fluorescence intensity was calculated for each strip. These data, when plotted against the relative position in the cell, produced the cellular distribution plots. Representative cells for each division stage are presented on the left, and their respective plots are presented on the middle column. For each division stage, the group’s average fluorescent signal distribution and standard deviation values are presented on the right. (B) A representation of the cellular regions of a streptococcal cell, as used in this study. (C) A plot displaying the total M protein fluorescence value for each cell in the population, plotted against its total SfbI fluorescence.
A: Automatic classification

- Mother only
- Both
- Daughter only
- N/A

B: Manual classification

- Mother only
- Both
- Daughter only
- N/A

C: vancomycin Rhodamine red, M protein

1 μm
Figure 3. M protein is anchored simultaneously at the mother and daughter septa. An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y containing trypsin and pronase and incubated at 37°C to OD$_{600}$ 0.5, at which point the cells were harvested, and resuspended in TH+Y without proteases for 30 seconds at 37°C prior to fixation. M protein was labeled using specific antibodies and average intensity 3D-projections were made for numerous DeltaVision images, resulting in a population of 1,039 cells. The M protein distribution plots of these cells were analyzed using MATLAB (see experimental procedures section) to determine the presence of newly anchored M protein at the mother and daughter septa (A). The location of M protein anchoring on the same cells was determined by direct observation (B). (C) Cells treated in a similar manner were labeled with vancomycin Rhodamine red conjugate (red), and M protein specific antibodies (green), and then visualized by 3D-SIM. Arrowheads represent mother septa and arrows represent forming daughter septa.
Figure 4. Gradual accumulation of SfbI results in polar distribution. An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y and grown to OD\textsubscript{600} 0.15, at which time GFP-PlyC/BD (green) was added to the medium for 30 minutes (“pulse”). The cells were then washed and suspended in TH+Y for one hour at 37°C (“chase”) prior to fixation. SfbI was stained using a specific serum (red), and the cell wall was stained using WGA marina blue (blue). (A) Representative cells before and after the “chase” period. (B) Streptococcal chains following the “chase” period alongside the separate fluorescent channels. (C) Average intensity 3D-projections were made for numerous DeltaVision images, resulting in a population of 714 cells. The signal distribution plots of these cells were analyzed using MATLAB. Each cell was divided into “young pole” (blue) and “old pole” (red) regions. For each pole, the mean GFP-PlyC/BD and SfbI fluorescence values were plotted against each other. (D) The “old pole” group was sub-divided into 3 groups according to GFP-PlyC/BD fluorescence, representing the estimated age of the pole. The mean SfbI fluorescence the standard deviation values are presented for each group, as well as for the “young pole” group in its entirety. (E) An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y containing trypsin and pronase, and incubated at 37°C to OD\textsubscript{600} 0.5, at which point the cells were washed, and resuspended in TH+Y without proteases for 50 minutes at 37°C prior to fixation. Specific antibodies were used to label M protein (red) and SfbI (green). The cell wall was stained with WGA marina blue (blue). Arrows denote wall regions that were already assembled at the time of protease treatment. DeltaVision images for (A), (B), and (E), are presented as maximum intensity 3D-projections.
Figure 5. Methicillin-induced unbalanced peptidoglycan synthesis results in a substantial reduction in the cellular amount of M protein compared to SfbI. An overnight *S. pyogenes* D471 culture was diluted 1:100 into TH+Y containing trypsin and pronase, and grown at 37°C to OD$_{600}$ 0.5. The cells were then diluted 1:4 into tubes containing TH+Y, trypsin, pronase, and ascending concentrations of methicillin. Following one hour, the cells were washed, and resuspended for 10 minutes in TH+Y containing a similar concentration of methicillin but lacking proteases, and then fixed. (A) M protein (red) and SfbI (green) were labeled using specific antibodies, and the cell wall was stained with WGA marina blue (blue). Deconvolution immunofluorescence images are presented as maximum intensity projections. (B) Similar cultures were fractionated into supernatant, wall, and spheroplast pellet, and processed by Western blot. (C) Additional cultures were harvested, boiled in 2% SDS, and separated into supernatant (“SDS Soluble” fraction), and cell pellet (“SDS insoluble” fraction), which was subsequently treated with the phage lysin PlyC to release wall-anchored proteins, prior to processing by Western blot. (D) Cells treated with 0.2 µg/ml methicillin, were visualized by 3D-SIM and are presented as maximum intensity projections.
Figure 6. Overexpression of DivIVA results in a marked decrease in the cellular amount of M protein and an increase in SfbI. (A) Overnight cultures of D471 cells harboring pAR291_DivIVA, or the empty vector pAR161, were diluted 1:50 into TH+Y+spec containing trypsin and pronase and grown to OD$_{600}$ 0.5. The cultures were then washed, resuspended in medium without proteases for 5 minutes, and then fixed. Specific antibodies were used to label M protein (red) and SfbI (green), and the cell wall was stained with WGA marina blue (blue). DeltaVision images are presented as maximum intensity projections. (B) D471 cells harboring pAR291_DivIVA, pAR287_GFP-HT (non-specific control), or the empty vector pAR161, were grown in TH+Y+spec to OD$_{600}$ 0.5, and fractionated into supernatant, wall, and spheroplast pellet fractions. Samples were examined by Western blot using antibodies specific for M protein and SfbI.
Figure 7. A model representation of the anchoring of M protein and SfbI. The regeneration of surface proteins during two division cycles is presented. M protein (red) is anchored exclusively to newly synthesized peptidoglycan at the septum (sites of active anchoring labeled yellow). Anchoring of M protein at daughter septa begins before the mother septum is completely closed, resulting in simultaneous anchoring at both locations. Following two generations, M protein is anchored to all newly synthesized, but not pre-existing, peptidoglycan. SfbI (green) is anchored over time in patches to peripheral peptidoglycan, with some preference to the poles. Following two generations, the oldest poles (I) display the most intense SfbI labeling, while one-generation-old poles show less labeling (II), and newly formed poles (III) show little SfbI labeling.
Figure S1. M protein is anchored simultaneously at the mother and daughter septa – analysis of additional populations. (A) Four overnight *S. pyogenes* D471 cultures were diluted 1:100 into TH+Y containing trypsin and pronase, harvested, and resuspended in TH+Y for 30 seconds at 37°C prior to fixation (each sample was processed separately). The fixed cells from the different cultures were mixed and processed together for microscopy. M protein was labeled using specific antibodies, and average intensity 3D-projections were made for numerous DeltaVision images, resulting in a population of 890 cells. The M protein distribution plots were analyzed using MATLAB to determine the presence of newly anchored M protein at the mother and daughter septa (see experimental procedures section). (B) The presence of newly anchored M protein at the mother and daughter septa, was analyzed in the cell population described in Figure 2 (following 2 minutes of protein regeneration), using MATLAB.
Figure S2. Distribution of M protein and SfbI in the sortase mutant, AR01. (A) Overnight cultures of the sortase deletion mutant AR01, and its parental strain D471, were diluted 1:100 into TH+Y, grown to OD$_{600}$ 0.5, and fractionated into supernatant (S), wall (W), and spheroplast pellet (P). Samples were separated by 10% SDS-PAGE, and examined by Western blot using antibodies specific for M protein or SfbI. (B) Overnight cultures of AR01 were diluted 1:100 into TH+Y or TH+Y containing trypsin and pronase, and grown to OD$_{600}$ 0.5. Protease treated cells were either fixed immediately upon reaching OD$_{600}$ 0.5 (no regeneration), or washed and resuspended in TH+Y without proteases for 5 minutes at 37°C prior to fixation (5 minutes regeneration). Specific antibodies were used to label M protein (red) and SfbI (green), and the cell wall was stained with WGA marina blue (blue). Images are presented as maximum intensity projections. (C) Sequential Z-sections of a representative protease-treated chain following 5 minutes regeneration.