Dual Role for Pilus in Adherence to Epithelial Cells and Biofilm Formation in *Streptococcus agalactiae*

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Abstract

*Streptococcus agalactiae* is a common human commensal and a major life-threatening pathogen in neonates. Adherence to host epithelial cells is the first critical step of the infectious process. Pili have been observed on the surface of several gram-positive bacteria including *S. agalactiae*. We previously characterized the pilus-encoding operon gbs1479-1474 in strain NEM316. This pilus is composed of three structural subunit proteins: Gbs1478 (PilA), Gbs1477 (PilB), and Gbs1474 (PilC), and its assembly involves two class C sortases (SrtC3 and SrtC4). PilB, the bona fide pilin, is the major component; PilA, the pilus associated adhesin, and PilC, are both accessory proteins incorporated into the pilus backbone. We first addressed the role of the housekeeping sortase A in pilus biogenesis and showed that it is essential for the covalent anchoring of the pilus fiber to the peptidoglycan. We next aimed at understanding the role of the pilus fiber in bacterial adherence and at resolving the paradox of an adhesive but dispensable pilus. Combining immunoblotting and electron microscopy analyses, we showed that the PilB fiber is essential for efficient PilA display on the surface of the capsulated strain NEM316. We then demonstrated that pilus integrity becomes critical for adherence to respiratory epithelial cells under flow-conditions mimicking an *in vivo* situation and revealing the limitations of the commonly used static adherence model. Interestingly, PilA exhibits a von Willebrand adhesion domain (VWA) found in many extracellular eucaryotic proteins. We show here that the VWA domain of PilA is essential for its adhesive function, demonstrating for the first time the functionality of a prokaryotic VWA homolog. Furthermore, the auto aggregative phenotype of NEM316 observed in standing liquid culture was strongly reduced in all three individual pilus mutants. *S. agalactiae* strain NEM316 was able to form biofilm in microtiter plate and, strikingly, the PilA and PilB mutants were strongly impaired in biofilm formation. Surprisingly, the VWA domain involved in adherence to epithelial cells was not required for biofilm formation.

Introduction

Group B Streptococcus (GBS, *Streptococcus agalactiae*) is a common colonizer of the gastrointestinal and urogenital tracts of up to 40% of healthy individuals [1]. However, in certain circumstances, GBS can become a life-threatening pathogen causing invasive infections in human neonates [2,3]. Epidemiological studies have documented how commonly GBS are transmitted from “carrier” mothers to newborn infants [4]. The clinical symptoms of acute GBS disease are pneumonia, septicemia, and meningitis. The lung is the portal of entry in clinical symptoms of acute GBS disease are pneumonia, septicemia, and meningitis. The lung is the portal of entry in circumstances, GBS can become a life-threatening pathogen in neonates [15]. The class A sortase is the ubiquitous housekeeping enzyme that anchors LPXTG proteins to the cell wall. The class B, C, and D involved in different functions [15]. The class A sortase is the ubiquitous housekeeping enzyme that anchors LPXTG proteins to the cell wall. The class B, C, and D involved in different functions [15]. The class A sortase is the ubiquitous housekeeping enzyme that anchors LPXTG proteins to the cell wall. The class B, C, and D involved in different functions [15].
D sortes are specifically involved in iron acquisition, pilus assembly and developmental processes including sporulation [15–17].

Sortase-mediated pilus assembly was first demonstrated in Corynebacterium diphtheriae [18,19] and these pioneer studies revealed the existence of 3 conserved genetic elements found within the major pilin subunit and necessary for pilus formation; i) the pilin motif (WxxxxVxVYPK); ii) the E-box domain (YxLxETxAPxG); and iii) the cell wall sorting signal (LPxTG followed by a hydrophobic domain and a positively charged tail). The current model for pilus assembly is as follows: the major subunit is assembled into a pilus by a cis-encoded sortase that catalyzes the covalent attachment between the conserved pilin motif lysine residue of one subunit with the conserved threonyl residue LPxTG motif of another subunit. In addition, one or more accessory subunits are incorporated into the pilus by an unknown mechanism, but requiring pilus-specific sortase as well as the E-box domain within the major pilin subunit. Then, during a final step, the pilus fiber is covalently linked to the peptidoglycan by either the pilus-specific sortase or the housekeeping sortase. This mechanism of pilus assembly catalyzed by class C sortases has now been demonstrated in several gram-positive pathogens using similar genetic and biochemical analyses [20–27].

We previously carried out a detailed structural and functional analysis of the pilus locus gbs1479-1474 (also referred to as PI-2A in GBS strain NEM316 [22]). This locus encodes a pilus composed of three structural subunit proteins Gbs1478 (PiA), Gbs1477 (PiB), and Gbs1474 (PiC) whose assembly involves two class C sortases (SrtC3 and SrtC4). PiB, the bona fide pilin, is the major component; PiC is a minor associated component mainly localized at the base of the pilus; and PiA is the pilus associated adhesin located at intervals along the pilus backbone. We previously showed that PiA mediates adherence of GBS NEM316 to the pulmonary epithelial cell line A549 independently of pilus formation [22]. The apparently paradoxical situation of a pilus that carries the adhesive property and yet is dispensable for binding was reported previously in the Escherichia coli Pap pilus model system [28] and more recently in the pneumococcal pilus [29]. We postulated that, in the absence of pilus, PiA behaves as a classical LPXTG-containing adhesin anchored to the cell wall by the housekeeping class A sortase SrtA to mediate adherence to cultured epithelial cells.

Bacteria often exist within natural systems in an entirely different form (sessile) from those grown in laboratory conditions (planktonic). Sessile bacteria appear to be protected in hostile environments by growing as colonies embedded in an extracellular matrix of carbohydrate or exopolysaccharide called biofilm. The pattern of biofilm development involves bacterial attachment to a solid surface, the formation of microcolonies, and then differentiation into exopolysaccharide-encased communities to form a mature biofilm. Many gram-negative pathogens use their pili to promote attachment and aggregation to host cells, that eventually develop into mature biofilm resulting in host tissues colonization [30]. Pilus contribution in biofilm formation was recently shown in gram-positive bacteria such as E. faecalis, S. pyogenes (GAS), and S. pneumoniae [11,26,31,32].

In this work, we investigated the roles of the housekeeping sortase A in pilus assembly in GBS and that of the pilus structure to resolve the paradox of a pilus dispensable in adherence assays although containing an adhesin subunit. We characterized the functional role of the von Willebrand adhesion domain found in the PilA adhesin. We adapted a biofilm formation assay for GBS and thus uncovered an essential role of GBS pilus in this process.

Roles of GBS PI-2A Pilus

The pilus is anchored to the cell-wall by the housekeeping class A sortase

Our previous functional characterization of the pilus locus in S. agalactiae [22] raised the question of the role of the housekeeping class A sortase (SrtA) in pilus biosynthesis but did not answer it since transcription of the PI-2A pilus locus was dramatically reduced in the srtA mutant of strain NEM316 [22]. This mutant was made by insertion of a promoterless aphA-3 kanamycin cassette within srtA by allelic replacement to generate a strain that synthesizes a truncated SrtA protein deleted of its carboxylic half (i.e., 127 out of 248 amino acids) including the catalytic TLXTC sequence [33]. Complementation of the srtA mutant with the wild-type gene inserted ectopically on NEM316 chromosome did not result in wild-type levels of pilus expression (data not shown), although restoring the correct localization of two model LPXTG proteins, Alp2 and ScpB [33].

To characterize the role of SrtA in pilus synthesis, we therefore constructed a catalytic mutant of SrtA by in frame-modification of the TLXTC signature sequence encompassing the critical cysteyl residue (TLTCTDPE to TAAAPGRAE replacement in the catalytic site). This new mutant named SrtA* exhibited phenotypes similar to those of the previously characterized SrtA mutant (Figure 1A–1C). It is unable to anchor the classical LPXTG protein Alp2 on the bacterial surface as shown by immunofluorescence (Figure 1A) or by Western blotting (Figure 1B, left panel). The ScpB protein was found in larger amounts in the supernatants of the SrtA* and SrtA+ mutants compared to the wild-type strain NEM316 (Figure 1B, right panel). As expected, the binding to human fibronectin- and fibrinogen-coated plates was similarly affected in both mutant strains (Figure 1C). Of note, the surface properties of SrtA* and SrtA+ were macroscopically different from that of the parental strain: they bound less to polypropylene-, MaxiSorp-, or glass-matrices and their pellets obtained after centrifugation were smooth (data not shown).

We then tested expression levels of the major pilin subunit PiB in the SrtA+ mutant by immunoblotting on whole bacteria. As shown in Figure 1D, the level of PiB in the SrtA+ mutant was

**Author Summary**

Streptococcus agalactiae (Group B Streptococcus) is a leading cause of sepsis (blood infection) and meningitis (brain infection) in newborns. Most bacterial pathogens have long filamentous structures known as pili or fimbriae, which are often involved in the initial adhesion of bacteria to host tissues but also in bacteria–bacteria interactions, resulting in biofilm formation. Our previous functional characterization of the pilus locus in S. agalactiae showed that it encodes a major pilin and two minor pilin subunits that are covalently polymerized by the action of two enzymes belonging to the sortase C family. One of the accessory pili is responsible for the adhesive property of the pilus. However, this initial study raised two major questions that were addressed in the present work: i) what anchors the pilus to the cell wall and ii) what is the function of the pilus fiber itself. We showed that the pilus is essential for optimal display of the pilus-associated adhesin and overcomes the masking effect of the capsule. Pilus integrity was shown to be critical in adherence assays under flow conditions. We also report that GBS can form biofilms and that pili play an important role in this process.

**Results**

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We then tested expression levels of the major pilin subunit PiB in the SrtA+ mutant by immunoblotting on whole bacteria. As shown in Figure 1D, the level of PiB in the SrtA+ mutant was
similar to that found in the wild-type strain. To unravel the role of the various sortases in the pilus assembly process, we monitored pilus polymerization in the various GBS sortases mutants by immunoblotting using specific anti-PilB polyclonal antibody (Figure 2). *S. agalactiae* isogenic strains were grown to the same optical density (OD$_{600}$=2) and the cultures were separated into three fractions (medium, cell-wall, and membrane) that were electrophoresed on 4–12% gradient SDS-PAGE and probed with the PilB antiserum (upper panel). The same fractions were also probed with an antiserum raised against the secreted protein Bsp [34] used as an internal loading control (lower panel). Pilus polymers are readily detected in the various fractions of the wild-type strain but, as previously reported [22], their polymerization requires either SrtC3 or SrtC4 (Figure 2). PilB monomer could be detected in the culture medium fractions as a band of about 80 kDa, the lower band at 60 kDa being a degradation product. As previously shown, pili are not expressed in the SrtA$^-$ mutant (Figure 2). In the SrtA* mutant, the pilus polymers are only found in the membrane and medium fractions, but not in the cell wall fraction. This result demonstrates that the housekeeping class A sortase is not necessary for pilus polymerization but is absolutely required for anchoring the pilus to the cell wall.
Pilus integrity is essential for PilA display at the cell surface

The PI-2A pilus of *S. agalactiae* is composed of three structural subunits PilA (Gbs1478), PilB (Gbs1477), and PilC (Gbs1474). PilB is the major constituent of the pilus fiber; PilC is a minor associated component mainly localized at the base of the pilus; and PilA is the pilus-associated adhesin located at intervals along the pilus backbone [22,27]. Immunogold electron microscopy revealed abundant surface staining and pilus structures extending largely beyond the capsule in strain NEM316 (Figure 3A). Using the previously characterized mouse monoclonal antibody S9 directed against the type III capsule polysaccharide [35], we carried out a triple-labeling experiment to detect simultaneously the PilB pilin, the PilA-associated adhesin, and the capsule. Wild-type (WT) and isogenic mutant bacteria were stained with: i) mouse mAb S9 followed by 5 nm gold-labeled IgG; ii) with rabbit pAb anti-PilB followed by 10 nm gold-labeled IgG, and iii) with rabbit pAb anti-PilA followed by 20 nm gold-labeled IgG. The mAb S9 decorates the external layer of the capsule [36] and its thickness in strain NEM316 was estimated to be ≈50 nm on ultrathin sections by transmission electron microscopy (Figure S1). In the absence of the PilB backbone pilin, the PilA adhesin is found at the cell surface without detectable pili (Figure 3A, bottom panel). As expected, the absence of the PilA accessory protein did not prevent pilus formation and in this mutant the pili are even longer than in wild-type strain (Figure 3A, bottom panel, and Figure S2). Strikingly, in the absence of the PilC ancillary protein, pili are longer but also more extended (Figure 3A, bottom panel). Of note, a significant amount of pili produced by *ApilA* and *ApilC* mutants were released in the culture medium compared to the parental strain (Figure S2). These immuno electron micrographs were subjected to quantitative analysis and the results are shown in Table 1. Pili were shown to be longer in both *ApilA* and *ApilC* compared to the wild-type strain. In addition, immunofluorescence analyses clearly shows that pili are not only longer but also thicker in the *ApilA* and *ApilC* (Figure S2). Immunoblotting analysis on whole bacteria confirmed the specificity of all four antisera (PilA, PilB, PilC, and S9) and showed that in the absence of the pilus backbone (*ApilB*), PilA cannot be detected at the bacterial surface (Figure 3B). PilA accessibility at the bacterial surface is also reduced in the *ApilC* mutant. Previous transcriptional and western blot analyses showed that deletion of *pilB* or *pilC* does not affect expression of *pilA* [22]. Altogether, these results reinforce the idea that pilus integrity is essential for efficient PilA display at the bacterial surface.

PilB-dependent display of PilA is essential for adherence to epithelial cells under flow-conditions

We previously showed that PilA mediates adherence of *S. agalactiae* strain NEM316 to the human alveolar epithelial cell line A549 independently of pilus formation [22]. Indeed, the apiliated *pilB* mutant is as adherent as the wild-type strain to A549 cells (Figure 4A) and the role of the pilus fiber in bacterial adhesion therefore remains to be characterized. A major defence in the lung is constituted by the mucociliary clearance apparatus. Goblet and glandular cells beneath the epithelium produce mucus that lines the epithelial layer of the air conducting pathways. Mucus is moved through the conducting pathways as fast as 1 cm/min by bronchial epithelial cell cilia to the trachea and later towards the mouth. We reasoned that surface display of PilA adhesin could be important in more stringent adherent conditions, e.g., in the presence of liquid fluid mimicking the mucociliary movement in the lung. A major limitation of the standard adhesion model is that it neglects the local fluid mechanic environment encountered in the organism. We therefore examined the role of the various pilus components under defined shear stress condition by analyzing *pilA*,
pilB, and pilC mutants. Adherent human alveolar epithelial cells (A549) were grown on glass slides and placed in a laminar flow chamber observed under an inverted microscope (for experimental details see [37]). *S. agalactiae* labeled with fluorescent 5-chloromethylfluorescein diacetate (CMFDA) was introduced in the chamber under a controlled flow. Before introduction of bacteria little or no fluorescence was detected. A low shear stress value (0.04 dynes/cm²) mimicking the mucus flow in the lung was selected. We showed that all three pilus mutants were significantly decreased for adherence as much as the *srtA* mutant under low shear stress (Figure 4B and Figure S3). The structural component of the pilus is therefore necessary for efficient adhesion in the presence of a shear stress reproducing the conditions encountered by the bacteria in the lung.

The von Willebrand Adhesion type A (VWA) domain of PilA is required for adherence to epithelial cells

*In silico* analysis of GBS PilA adhesin revealed the presence of a von Willebrand factor type A domain (*VWA* found at amino acids 228 to 585) located upstream from the putative pilin motif (YPK). This VWA domain is flanked by two Cna-B type domain found in a *S. aureus* collagen-binding surface protein (Figure 5A). However, the Cna-B regions do not mediate collagen binding but forms a stalk that presents the ligand binding domain away from the bacterial surface [38]. VWA domains in extracellular eukaryotic proteins mediate adhesion via metal ion-dependent adhesion sites (MIDAS). Binding of Mn²⁺ and Mg²⁺ to the MIDAS region in eukaryotic proteins have been demonstrated by crystallographic structures. Divalent cations were shown to stabilize the α₁β₁ integrin I domain [39]. Of note, the critical serine and aspartate residues known to interact with divalent cations are conserved in the VWA domain of PilA (Figure 5A). Many homologues have been identified in bacterial genomes but their role have not been characterized [40]. Multiple sequence alignments of prokaryotic and eukaryotic VWA-domains is shown in Figure S4.

We sought to determine whether the VWA domain of PilA was involved in PilA-mediated adherence. To test this hypothesis, we constructed a PilA mutant named ΔVWA in which the first 180 amino acids of the 358 amino acids VWA domain was replaced by a 9-aa residue-long hemagglutinin epitope tag (HA tag) allowing the detection of the mutant protein with specific anti-HA monoclonal antibody (Figure S4). Of note, the putative pilin motif YPK of PilA allowing its incorporation into the pilus fiber is

| Strain          | WT | Δ pilA | Δ pilB | Δ pilC |
|-----------------|----|--------|--------|--------|
| **Experiment 1**|    |        |        |        |
| Sample number   | 48 | 39     | 50     |        |
| Mean length in μM | 0.29 | 0.42* | 0.65*  |        |
| Standard Deviation | 0.134 | 0.155 | 0.299  |        |
| **Experiment 2** |    |        |        |        |
| Sample number   | 25 | 38     | 28     |        |
| Mean length in μM | 0.296 | 0.503* | 0.582* |        |
| Standard Deviation | 0.107 | 0.185 | 0.243  |        |

*Using the Mann-Whitney Test, the two-tailed P value is < 0.0001 between WT and mutant strains considered extremely significant.

Table 1. Quantification of pilus length in *S. agalactiae* NEM316 and mutant derivatives.

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The number of fluorescent bacteria per mm² was same strains labeled with fluorescent CMFDA under liquid flow experiment of at least 3 independent experiments. (B) Adherence of the results are presented as mean value (±SD) is representative of 3 independent experiments. (B) Adherence of the various mutants are relative values. The level of adherence of the WT strain is arbitrarily reported as 100, and the level of adherence of the various mutants are relative values. The results presented as mean value (±SD) from one representative experiment of at least 3 independent experiments. (B) Adherence of the same strains labeled with fluorescent CMFDA under liquid flow (0.04 dynes/cm²). The number of fluorescent bacteria per mm² was determined (see Experimental Procedures). The level of adherence of the WT strain is arbitrarily reported as 100 and the level of adherence of the various mutants are relative values. The results presented as mean value (±SD) is representative of 3 independent experiments. doi:10.1371/journal.ppat.1000422.g004

located behind the VWA domain and left intact in the mutant (Figure 3A). This new isogenic pilAAVWA mutant displayed similar growth characteristics in Todd-Hewitt broth compared to the parental strain at 37°C (data not shown). Dot-blot analysis on whole bacteria using a commercial anti-HA antibody showed that the HA epitope is located at the bacterial surface of the ΔVWA HA-expressing bacteria. No signal could be detected in the parental strain confirming the specificity of the HA antibody (Figure 3B). Interestingly, introduction of the srtC3C4 mutation in the ΔVWA strain to abrogate pilus polymerization caused the disappearance of the HA signal (Figure 5B). This result strongly suggests that HA detection on the bacterial surface of ΔVWA mutant depends on its incorporation into the pilus fiber. Western-blot analysis of cell wall extracts from isogenic mutants showed the presence of the HA-tagged PilAAVWA protein in the pilus polymers of the ΔVWA strain but this incorporation is abolished in the ΔVWA/SrtC3C4− mutant where a single 60-kDa protein, i.e. the predicted size of monomeric PilAAVWA protein, is present (Figure 5C). We also analyzed the interaction between HA-tagged PilA mutant and the major pilin subunit PilB using pull-down experiment with HA agarose beads and immunoblotting with anti-PilB antibody. As shown in Figure 5D, the HA-tagged PilAAVWA physically interact with PilB polymer in cell wall extracts. No signal was detected with wild-type extracts as control for HA specificity (data not shown). Immunolocalization of the PilAAVWA-HA protein in the pilus fiber was demonstrated by scanning electron microscopy (Figure 5E). Double-labeling experiments were performed on the parental ΔVWA and its isogenic SrtC3C4− mutant using rabbit anti-PilB polyclonal antibody followed by 10 nm gold-labeled IgG (thin arrows) and then with rat anti-HA monoclonal antibody followed by 20 nm gold-labeled IgG (arrow heads). HA staining was detected at various locations including the base and the tip (Figure 5E) and was similar to that of PilA staining (Figure 3A and [22,27]). No staining was detected in the absence of pilus polymerization in the ΔVWA/SrtC3C4− double mutant (Figure 5E). Altogether these results indicate that the PilAAVWA protein is produced, folded, and incorporated into the pilus fiber like the intact PilA protein.

Finally, we examined the ability of the PilAAVWA mutant to bind to human epithelial cells from alveolar (A549) and intestinal (TC7) origins. Standard adhesion assays showed that the ΔVWA mutant is strongly reduced for adherence to both A549 and TC7 cell lines compared with the parental strain NEM316, to a level similar to that obtained with the pilA and the SrtA* mutants (Figure 6). Collectively, these results show that the VWA domain of PilA is essential for PilA adhesive property.

The PI-2A pilus is involved in biofilm formation

We initially observed that all pilus mutant strains remained in suspension after an overnight culture in Todd-Hewitt broth whereas strain NEM316 sedimented at the bottom of the tube (data not shown). This result suggested a role of the PI-2A pilus locus in bacterial aggregation and possibly in biofilm formation. We thus began to assay the ability of S. agalactiae to form biofilms on microtiter polystyrene plate as previously described [41]. In this assay, staining with 0.1% crystal violet (CV) for 15 min enables the visualization of attached, sessile cells after bacterial biofilms have formed in microtiter plate wells. Biofilm assays were carried out under various conditions to determine the optimum experimental conditions. Various media (TH, THY, BHI, LB, RPMI 1640), temperature (30°C and 37°C), and time points (24 to 48 h) were used in preliminary experiments but only LB and RPMI 1640 media supplemented with 1% glucose at 37°C for 24 h produced uniform biofilms (data not shown). In enriched media such as TH, THY, BHI bacteria grew better than in LB or RPMI media but failed to evenly adhere over the surface, instead forming pellets at the bottom of the well. It appears that a nutritionally rich environment does not favor S. agalactiae biofilm formation on polystyrene but that nutritionally limited environment increases sessile growth. We also compared the ability of S. agalactiae to form biofilms on different surfaces. Polystyrene surface was more suited than polystyrenechloride or glass surfaces on which S. agalactiae adhered poorly. Thus, the optimal conditions to see biofilm formation with strain NEM316 were as follows: overnight culture
in Todd-Hewitt medium, dilution in LB medium supplemented with 1% glucose to obtain an initial OD 600 nm of 0.05, inoculation of sterile polystyrene 96-well plate and growth at 37 °C for 24 h. Biofilm formation of \textit{S. agalactiae} strain NEM316 and its isogenic pilus mutants were assayed accordingly (Figure 7). We hypothesized that the sortase A mutants, unable to attach to the polystyrene surface, would be defective for biofilm formation as recently reported for \textit{S. gordonii} srtA mutant [42]. Indeed, both \textit{S. agalactiae} \textit{srtA} and \textit{srtA*} mutants were unable to form biofilm (Figure 7). We showed that \textit{pilA} and \textit{pilB} mutants were as strongly impaired as the \textit{srtA} mutants for biofilm formation. The \textit{pilC} mutant that still forms pili was only slightly reduced for biofilm formation. Surprisingly, the \textit{pilAVWA} mutant readily forms thicker biofilm, as compared to the parental strain NEM316, although it is unable to adhere to epithelial cells (Figure 6 and Figure 7).

**Discussion**

Our previous functional characterization of the pilus locus in \textit{S. agalactiae} [22] raised two major questions that were addressed in the present work: i) what is the role of the housekeeping class A sortase (SrtA) in pilus biosynthesis and ii) what is the function of the pilus fiber itself. Indeed, understanding the apparent paradox of a pilus carrying the adhesive property but yet dispensable for adherence remains a major challenge of the field.
We previously observed a down-regulation in transcription of the pilus genes in the $\text{srtA}^{-}$ mutant and therefore could not test the role of $\text{SrtA}$ in pilus biogenesis [22]. In this report, a new $\text{srtA}^{-}$ mutant ($\text{srtA}^{+}$) displaying all characteristics of the $\text{srtA}^{-}$ mutant but expressing wild-type levels of PI-2A pilus was constructed. Since high molecular weight polymers of pili were seen in the $\text{srtA}^{+}$ mutant, it is clear that the housekeeping sortase A is not involved in the polymerization process. This is in direct contrast to the effects of deleting both pilus-associated sortases $\text{SrtC}$ that abrogates the formation of pilus polymers (Figure 2). Shedding of pilus polymers in the culture medium of the $\text{srtA}^{+}$ mutant demonstrate a role of the housekeeping sortase A in the anchoring phase. A similar result was very recently reported in $\text{S. agalactiae}$ strain 515 [43]. As previously shown in $\text{C. diphtheriae}$ [44] and $\text{B. cereus}$ [21], these results support a two-stage model of pilus assembly where pili are first polymerized by a pilus-specific sortase and the resulting fiber is then attached to the cell wall by the housekeeping sortase. In contrast, $\text{SrtA}$ is dispensable for pilus assembly and localization to the cell wall in $\text{S. pneumoniae}$ [45]. Interestingly, the three pneumococcal $\text{RrgA}$, $\text{RrgB}$, and $\text{RrgC}$ proteins that assemble into the pilus each have a motif (YPRTG, IPQTG, and VPDTG respectively) that is divergent in the first amino acid position of the canonical LPxTG cell wall signature sequence (CWSS) recognized by the house-keeping sortase A which could account for differences in sortase specificity.

**Figure 6. Adherence of S. agalactiae pilus mutants to human pulmonary epithelial cells A549 and to human intestinal epithelial cells TC7.** Cells were infected at a MOI of 20 bacteria per cell for 1 h at 37°C and adherence frequencies were calculated from the numbers of bacteria remaining attached to the cells after the incubation period with respect to the number of inoculated bacteria. The level of adherence of the WT strain is arbitrarily reported as 100 and the level of adherence of the various mutants are relative values. The results are presented as mean value (±SD) from one representative experiment of at least 3 independent experiments (left panel). Immunofluorescence analysis of GBS adherence to A549 and TC7. Bacteria were revealed with specific rabbit anti-GBS polyclonal antibodies and anti-rabbit IgG coupled to Alexa 488 respectively. Cellular F-actin was visualized with phalloidin coupled to Alexa 594 and nuclei were stained with DAPI (right panel).

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S. agalactiae is a capsulated bacteria and the size of the capsule is subject to phase variation [46]. By immunogold labeling, we visualized the capsule by electron microscopy and showed that the pilus extends beyond the capsule and thus serve as carrier for surface located adhesive clusters of PilA. Thus, pilus-associated adhesins, as opposed to those directly linked to the peptidoglycan, can overcome masking by the capsule as demonstrated by immunodetection of PilA on capsulated bacterial surface (Figure 3B). In a capsulated strain, PilA is not detected in the absence of PilB or in the srtC3C4 mutant in which pilus polymerization is abrogated (Figure 3B, data not shown). These results indicate that the pilus structure is necessary for optimal display at the bacterial surface of the PilA subunit that is necessary for adherence to epithelial cells. However, the fact that PilA remains a functional adhesin in the absence of a pilus fiber raises question on the role of this appendage. A similar situation was recently reported in Streptococcus pneumoniae in which RrgA, a minor pilus component, is central in pilus-mediated adherence and disease, even in the absence of polymeric pilus production [29]. As mentioned in this work, it is conceivable that the conventional in vitro adherence assays carried out with immortalized cells culture are not adapted to test the functional benefit provided by a pilus fiber. This was recently demonstrated for the pili of S. pyogenes that mediate specific adhesion to human tonsil and skin epithelial cells [47]. The authors showed that pili were not required for S. pyogenes adhesion to immortalized HEp-2 and A549 cell lines but were indispensable for adhesion to ex vivo tissues and primary human keratinocytes highlighting an important limitation of the currently used adhesion models. We reasoned that surface display of PilA adhesin could be important in more stringent conditions, as for example in the presence of liquid flow mimicking the mucociliary movement in the lung. A major limitation of the standard adhesion model is that it neglects the local fluid mechanic environment encountered in the organism. Using a laminar flow chamber system optimized to study the adhesion of Neisseria meningitidis under low shear stress conditions [37], we were able to prove the benefit of the S. agalactiae pilus fiber for adherence to human pulmonary epithelial cells A549 (Figure 4B), thus emphasizing the need of employing models that are more relevant to the infectious process when studying bacterial-host interactions. Closer examination of EM micrographs shows a large heterogeneity in pilus structures in the wild-type strain. The composition, the size, but also the diameter of the individual pilis appears highly variable and, as described in S. pneumoniae, bundles of individual pili could be also observed (data not shown). In agreement with our previous results [22], pili are still formed in the pilA mutant but were longer than those produced by the wild-type strain. This is most probably due to the increased transcription of pilB in the pilA mutant where a 3-fold increase in pilB expression was measured by qRT-PCR [22]. Synthesis of longer pili by mutants overexpressing the major pilin subunit has been demonstrated in C. diphtheriae [48]. Strikingly, we also observed longer and largely extended pili in the pilC mutant and a higher amount of PilB polymers were found shed in the culture medium of this mutant in agreement with a role of PilC as the pilus anchor [43]. Again, similar results were obtained recently in C. diphtheriae [49], a bacterium where the prototype pilus contains a major pilin (SpaA), a tip pilin (SpaC), and a minor pilin (SpaB). Immunoelectron microscopy revealed that when SpaB was absent, the SpaA fibers found in the culture medium and on the bacterial envelope are considerably longer than in the wild-type strain. Incorporation of the SpaB minor pilin in the shaft base serves as the terminal step in pilus polymerization and triggers the concomitant cell wall linkage by sortase A [49].

The von Willebrand Adhesion Domain (VWA) has been identified in several prokaryotic proteins but their function remain unknown [40]. We showed here that the VWA domain of PilA is essential for its adhesive function. S. agalactiae strain NEM316 possesses another pilus locus (PI-I) that is not expressed [22] but displayed a genetic organization similar to that of the PI-2A locus. In particular, the putative pilus associated adhesin (Gbs0632) also contains a central VWA domain surrounded by two Cna-B domains and, interestingly, this central domain structure was also found in the pneumococcal pilus-associated adhesin RrgA [29] and in the minor pilin SpaC of C. diphtheriae [50]. Extracellular matrix proteins constitute good ligand candidates for these adhesins and it was recently shown that RrgA interacts with human fibronectin, collagen I, and laminin [51]. However, sequence comparisons revealed that the VWA domain of RrgA shares 59% identity with that of Gbs0632 but only 37% with that of PilA, suggesting that RrgA and PilA have different VWA-binding ligands. In agreement with this idea of different receptor recognized by different VWA domain, SpaC was shown to promote specific adhesion to human pharyngeal cell line D562 [50].

Finally, we investigated the possibility that GBS pili could also play a role in bacterial-bacterial interactions, as shown for E. coli.
fusalis, S. pyogenes, and S. pneumoniae [26,31]. We demonstrated that all three individual pilus NEM316 mutants were impaired for bacterial aggregation in liquid culture. Importantly, S. agalactiae strain NEM316 was able to form biofilm in microtiter plates under certain culture conditions. We demonstrated that pili are key surface structures involved in biofilm formation and showed that both PilB and PilA, but not Pic, are essential in this process. Surprisingly, the VWA domain required for adherence to epithelial cells was found to be dispensable for biofilm formation on polystyrene plates. This result indicates that the VWA domain is not required for adherence to abiotic surfaces and suggests that it recognizes specific ligand on epithelial cells. These results revealed GBS pili possess dual and non-overlapping functions in participating in biofilm formation and adherence to host cells. Current work aiming at identifying the epithelial receptor of PilA is ongoing in biofilm formation.

Materials and Methods

Bacterial strains, plasmids, and growth conditions
S. agalactiae NEM316 was responsible for a fatal septicemia and belongs to the capsular serotype III. The complete genome sequence of this strain has been determined [52]. Escherichia coli DH5α (Gibco-BRL) was used for cloning experiments. S. agalactiae was cultured in Todd-Hewitt (TH) broth or agar (Difco Laboratories, Detroit, MI) and E. coli in Luria-Bertani (LB) medium. Unless otherwise specified, antibiotics were used at the following concentrations: for E. coli - ampicillin, 100 μg/ml; erythromycin, 150 μg/ml; for S. agalactiae - erythromycin, 10 μg/ml; kanamycin, 1,000 μg/ml. S. agalactiae liquid cultures were grown at 37°C in standing flasks.

General DNA techniques
Standard recombinant techniques were used for nucleic acid cloning and restriction analysis [53]. Plasmid DNA from E. coli was prepared by rapid alkaline lysis using the Qiaprep Spin Miniprep kit (Qiagen). Genomic DNA from S. agalactiae was prepared using the DNaseasy Blood and Tissue kit (Qiagen). PCR was carried out with Ampli Taq Gold polymerase as described by the manufacturer (Applied Biosystem). Amplification products were purified on Sephadex S-100 columns (Pharmacia) and sequenced with an ABI 310 automated DNA sequencer, using the ABI PRISM dye terminator cycle sequencing kit (Applied Biosystems).

Construction of S. agalactiae mutants
In-frame replacement of the VWA by an HA tag domain in pilA (gbs1478) (O1–O2; O3–O4) and modification of the catalytic sequence signature of sortase A gbs0949 (O5–O6; O7–O8) were constructed by using splicing-by-overlap-extension PCR as previously described [22]. Mutants were confirmed by PCR and sequence analysis.

The sequences (‘5’ to ‘3’) of the primers were: O1, ACCAAT-GAATTCGGGAAAGTACGATCCGACGGGACGTGTCCGGTATCTGGTGGTCTCTTAC; O3, TACCCTTACGACGTCCCGAGTACG-GCTTTGGGCTATCATATGAAAGCCAAATTCTGA; O4, GGAATGCGATCTCCTATCGGCCTGGGTATATACTTACG; O5, TAAACGAAATTCGATCTCTGATTCTTACGG; O6, GCCGGCAGC-CGGGACGCTGCGGCAAGTCTTTGGCTGAGTG; O7, CGGGCAGCAGCAGCGGCTCCGGGAAGGCAG-AAGCTTATTATGTG; and O8, TCTTGGATCCGTA-TAGTCATCGTAAAGCAGATAGGC.

Cell culture and adherence assays
The human cell lines A549 (ATCC CCL-185) from an alveolar epithelial carcinoma and TC7 clone [54] established from the parental colon adenocarcinoma Caco-2, were cultured in Quantum 286 Medium (PAA). Cells were incubated in 15% CO2 at 37°C and were seeded at a density of 2 to 5 × 10^5 cells well in 24-well tissue culture plates. Monolayers were used for 24–48 h of incubation.

Bacterial cultures from overnight cultures OD600 of 2 (approximately 6 × 10^8 CFU/ml) were washed once in PBS and resuspended in DMEM. Cells were infected at a multiplicity of infection (M.O.I) of 10 bacteria per cell for 1 h at 37°C in 10% CO2. The monolayers were then washed four to five times with PBS, and the cells were disrupted by the addition of 1 ml sterile deionized ice-cold water and repeated pipetting. Serial dilutions of the lysate were plated on TH agar for count of viable bacteria. The percent of adherence was calculated as follows: (CFU on plate count/CFU in original inoculum) × 100. Assays were performed in triplicate and were repeated at least three times.

Laminar flow chamber experiments
Adhesion under flow was performed as previously described [37]. Before the assay, bacteria were grown overnight in TH broth at 37°C, resuspended at OD600 = 0.3, and labeled for 30 min with the fluorescent marker CMFDA (Molecular Probes) at 20 μM on ice. After several washes in PBS, fluorescent bacteria were resuspended in DMEM supplemented with 10% FBS. A549 cells grown on glass slides were placed in the parallel plate flow chamber (3.3 cm×0.6 cm×250 μm, Immunelectics, MA, USA) and sealed with vacuum. About 3×10^7 fluorescent bacteria were introduced in the laminar flow chamber containing the cells at 0.04 dynes/cm2. Experiments were performed in DMEM supplemented with 2% serum and maintained at 37°C with a heated platform (Minitub, Germany). Medium was introduced into the chamber using a syringe pump (Vial Medical, Becton-Dickinson or Harvard Apparatus). Adhesion of bacteria was recorded using an Olympus CKX41 inverted microscope with a 20× objective, a Hamamatsu ORCA255 CCD camera and the Openlab darkroom software (Improvement, UK).

Protein solubility in hot SDS
Cell-wall anchored proteins are insoluble in hot SDS unless the peptidoglycan had been first digested enzymatically with mutanolysin. In contrast, membrane anchored proteins are generally extractable in hot SDS without any prior treatment. The assay described by Garandeau et al. [55] was used to study the solubility of PilB polymers in NEM316 and sortases derivatives. The bacteria in 10 ml overnight culture were collected by centrifugation (6,000 rpm, 4°C, 10 min). Medium corresponds to the supernatant that was filter-sterilized and concentrated 10× by ultra filtration on Sartorius vivaspin 20 devices (cut-off 10 kDa). The bacterial pellet was washed in phosphate-buffered saline (PBS), centrifuged, and resuspended in 500 μl of 4% SDS - 0.5 M Tris-HCl pH 8. The bacterial suspension was boiled for 10 min and then centrifuged at 10,000 rpm for 5 min. Membrane correspond to the SDS-extracted supernatant and cell-wall to the pellet. These different protein fractions were further analyzed by immunoblotting.

Immunoblotting and immunofluorescence analyses
For dot-blot analysis on whole bacteria, late-exponentially growing bacteria were washed in PBS and resuspended in adjusted volumes of PBS to get similar OD600 values. The bacteria were
loaded on nitrocellulose membrane, dried up for 20 min at room temperature, and then blocked in PBS-milk 5% for 30 min. PilB was detected using a specific rabbit polyclonal antibody obtained previously [22] at 1:2000 dilution and the HA epitope was detected using the rat monoclonal antibody (3F10) from Roche at 1:1000 dilution. The secondary horseradish peroxidase (HRP)-coupled anti-rabbit secondary antibody (Zymed) was used at 1:20000 dilution whereas the goat anti-mouse antibody was used at 1:10000 dilution. Detection was performed using the Western pico chemiluminescence kit (Pierce). Image capture and analysis were done on GeneGnome imaging system (Syngene). For Western blotting analysis, proteins were boiled in Laemmli sample buffer, resolved on Tris-Glycine Criterion XT gradient gels 4–12% SDS-PAGE gels and transferred to nitrocellulose membrane (Hybond-C, Amersham). Protein detection was performed as described above.

Immunofluorescence staining of R28/Alp2 and PilB was performed as described [56] using specific rabbit polyclonal antibodies revealed with an anti-IgG coupled to Alexa 488 (Molecular Probes, OR). Microscopic observations were done on a Nikon Eclipse E600 and images acquired with a Nikon Digital Camera DXM1200F.

**Pull-down experiment**

Bacteria (50 ml) were grown in TH medium at 37°C for 18 hours and harvested for preparation of cell wall extracts. Bacteria were washed once in PBS and resuspended in the mutanolysin digestion buffer to get an OD600 of 100 ml^-1 (50 mM Tris-HCl pH 7.3, 20% sucrose and protease inhibitor cocktail (Roche)). Mutanolysin (Sigma) dissolved to 5000 U ml^-1 in potassium buffer (10 mM pH 6.2) was then added to the bacterial suspension to give a final concentration of 200 U ml^-1. The digestion was performed for 2 h at 37°C under gentle rotation. After centrifuging at 12000 g for 15 min at 4°C, supernatants corresponding to the cell wall fractions were transferred to clean tubes. 25 μl of EZview Red anti-HA affinity gel (EZview, Sigma) was added and the samples were rotated overnight at 4°C. Beads were washed five times in solubilization buffer (20 mM Tris-HCl, 137 mM NaCl, 0.25% NonidetP40, 1.5 mM MgCl2, 1 mM EDTA, 10 mM NaF) and resuspended in 20 μl of 2× reducing sample buffer followed by boiling for 5 minutes. Samples were then analyzed by Western blot analysis.

**Immunogold electron microscopy**

For scanning electron microscopy analysis, bacteria were applied to polysine coated glass coverslips, and fixed with 0.1% glutaraldehyde/4% paraformaldehyde in 0.1 M Sorensen buffer (pH 7.2) for 30 min. Fixed bacteria were incubated in PBS supplemented with 0.25% NH4Cl for 20 min then washed extensively with PBS. Samples were incubated in incubated in PBS/BSA 1% for 10 min. Following incubation for 30 min with the primary antibody, samples were washed and incubated for 10 min with the secondary antibody conjugated to colloidal gold. Preparations were washed with PBS and fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.2) overnight at 4°C, then washed three times for 5 min (each time) in 0.2 M cacodylate buffer, post-fixed for 1 h in 1% osmium in 0.2 M cacodylate buffer and rinsed with distilled water. Bacteria were dehydrated through a graded series of ethanol (25, 50, 75, 95 and 100%) followed by critical point drying with CO2. Dried specimens were sputter coated twice with carbon, with a with a GUN ionic evaporator PEC 682 and were examined and photographed with a JEOL JSM 6700F field emission scanning electron microscope operating at 5 kV. Images were acquired from the YAG BSE detector. For transmission electron microscopy, samples were processed as above. After dehydratation in ethanol the samples were embedded in epoxy resina and 70 nm thin sections were prepared and examined using a JEOL JSM1010 microscope operating at 80 kV.

For double and triple labeling experiments, the same procedure was applied using the following antibodies: the mouse monoclonal S9 anti-type III capsule (1/5), the rat monoclonal anti HA antibody (clone 3F9 from Roche at 1/100), the rabbit polyclonal α-PilB (1/100) and the rabbit polyclonal α-PilA (1/10). The secondary antibodies were goat anti mouse or goat anti rabbit conjugated to 20 nm-, 10 nm- or 5 nm gold beads.

**Biofilm formation assays**

Bacterial attachment and surface growth on polystyrene microtiter plates were studied during growth of S. agalactiae in LB medium supplemented with 1% glucose. Overnight cultures grown in TH were used to inoculate LB glucose medium at OD600 0.1, were vortexed briefly and 180 μl volumes were dispensed into 96-wells plate (Costar 3799; Corning, Inc., NY) followed by incubation at 37°C for 24 h. The OD600 of each culture was measured to ensure that all cells had reached stationary phase with a similar OD600, and the wells were washed twice in PBS and air-dried for 15 min. Biofilms were stained with 0.1% crystal violet for 30 min (100 μl per well) and the wells were washed twice with PBS and air-dried. The stained biomass was resuspended for quantification in ethanol/acetone (30:20) and A595 was measured. The assay was performed in quadruplet.

**Supporting Information**

Figure S1  IEM analysis of the capsular type III polysaccharide. S. agalactiae wild-type strain NEM316 was incubated with a mouse monoclonal antibody raised against the type III capsular polysaccharide (mAb S9) and rabbit polyclonal antibody raised against PilA and PilB, and. Antibodies were conjugated to 5 nm gold particles for capsule, 10 nm for pilB and 20 nm for PilA. The outer layer of the capsule is marked by black arrows. Scale bar is shown for each panel.

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Figure S2  Visualization of pili by immunofluorescence. Visualization of pili by immunofluorescence using polyclonal anti-PilB antibody. (A) on whole bacteria- (B) detached pili found in the extracellular medium.

Found at: doi:10.1371/journal.ppat.1000422.s002 (3.00 MB TIF)

Figure S3  Adhesion under flow conditions. A monolayer of A549 cells was cultivated and placed in a flow chamber. The same amount of fluorescently labeled strains were introduced under flow and adherent bacteria were detected by fluorescent microscopy. Representative fields are presented: A549 cells as seen by phase contrast (A) and adherence of the wild type strain NEM316 and isogenic mutant derivatives (B-F).

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Figure S4  Multiple sequence alignments of procaryotic and eucaryotic VWA-domains. The alignment was generated by comparing PilA (GBS1478) to the SMART database using Profile hidden Markov models (HMMER). Computations were made online at the following URL (http://smart.embl-heidelberg.de/). Note the presence of a VWA-domain in the pili adhesin subunit Gbs0632 from Streptococcus agalactiae (PI-1 pili operon), RggA from Streptococcus pneumoniae, and SpaC from Corynebacterium diphteriae. Explanation of codes used in CHROMA coloured alignments is as described in http://smart.embl-heidelberg.de/help/chroma.shtml.
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Author Contributions

Conceived and designed the experiments: GD PTC SD. Performed the experiments: YKG EM AM SD. Analyzed the data: EM AM GD PTC SD. Contributed reagents/materials/analysis tools: EC. Wrote the paper: GD PTC SD.

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