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The Metal Ion-Dependent Adhesion Site Motif of the *Enterococcus faecalis* EbpA Pilin Mediates Pilus Function in Catheter-Associated Urinary Tract Infection

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ABSTRACT

Though the bacterial opportunist *Enterococcus faecalis* causes a myriad of hospital-acquired infections (HAIs), including catheter-associated urinary tract infections (CAUTIs), little is known about the virulence mechanisms that it employs. However, the endocarditis- and biofilm-associated pilus (Ebp), a member of the sortase-assembled pilus family, was shown to play a role in a mouse model of *E. faecalis* ascending UTI. The Ebp pilus comprises the major EbpC shaft subunit and the EbpA and EbpB minor subunits. We investigated the biogenesis and function of Ebp pilus in an experimental model of CAUTI using a panel of chromosomal pilin deletion mutants. A nonpiliated pilus knockout mutant (EbpABC− strain) was severely attenuated compared to its isogenic parent OG1RF in experimental CAUTI. In contrast, a nonpiliated ebpC deletion mutant (EbpC− strain) behaved similarly to OG1RF *in vivo* because it expressed EbpA and EbpB. Deletion of the minor pilin gene ebpA or ebpB perturbed pilus biogenesis and led to defects in experimental CAUTI. We discovered that the function of Ebp pilus *in vivo* depended on a predicted metal ion-dependent adhesion site (MIDAS) motif in EbpA’s von Willebrand factor A domain, a common protein domain among the tip subunits of sortase-assembled pili. Thus, this study identified the Ebp pilus as a virulence factor in *E. faecalis* CAUTI and also defined the molecular basis of this function, critical knowledge for the rational development of targeted therapeutics.

IMPORTANCE

Catheter–associated urinary tract infections (CAUTIs), one of the most common hospital-acquired infections (HAIs), present considerable treatment challenges for physicians. Inherently resistant to several classes of antibiotics and with a propensity to acquire vancomycin resistance, enterococci are particularly worrisome etiologic agents of CAUTI. A detailed understanding of the molecular basis of *Enterococcus faecalis* pathogenesis in CAUTI is necessary for the development of preventative and therapeutic strategies. Our results elucidated the importance of the *E. faecalis* Ebp pilus and its subunits for enterococcal virulence in a mouse model of CAUTI. We further showed that the metal ion-dependent adhesion site (MIDAS) motif in EbpA is necessary for Ebp function *in vivo*. As this motif occurs in other sortase-assembled pili, our results have implications for the molecular basis of virulence not only in *E. faecalis* CAUTI but also in additional infections caused by enterococci and other Gram-positive pathogens.

In recent decades, *Enterococcus faecalis* and *Enterococcus faecium*, commensal gut bacteria, have emerged as human pathogens (1). Enterococci frequently cause hospital-acquired and device-associated infections, including bloodstream infections, infective endocarditis, and catheter-associated urinary tract infections (CAUTIs) (2). As the most common hospital-acquired infection (HAI) (3) and because they are frequently and often unnecessarily treated with antibiotics (4), CAUTIs are a reservoir of nosocomial and antimicrobial-resistant pathogens (5). Due to the tremendous incidence of CAUTI, infrequent but life-threatening sequelae such as bacteremia and urosepsis are significant complications (6). Enterococci are responsible for roughly 15% of CAUTIs (7). The recent surge of enterococcal infections correlates with both their inherent and acquired antimicrobial resistances, including vancomycin resistance (2). Both rising antimicrobial resistance and bacterial biofilm formation on indwelling catheters, biofilm surfaces protected from host immune responses and antibiotics, contribute to the difficulty associated with successful CAUTI treatment. Thus, a better understanding of bacterial pathogenesis in CAUTI is critical for the development of preventative and therapeutic antimicrobial agents.

Among the few described enterococcal virulence factors is the...
endocarditis- and biofilm-associated pilus (Ebp) (8). The Ebp pilus is an example of the sortase-assembled pilus family, whose members are now described among diverse Gram-positive genera, including Corynebacterium (9), Actinomyces (10), Streptococcus (11-13), Bacillus (14), Enterococcus (8, 15, 16), Lactobacillus (17), and Bifidobacterium (18). Sortase-assembled pili consist of a major pilin subunit and up to two minor subunits, each with a C-terminal cell wall sorting signal (CWSS) that includes an LPXTG-like sortase recognition motif (19). One or more genetically linked membrane-associated transpeptidase enzymes, pilus-associated sortases, catalyze the formation of interpilin isopeptide bonds found in mature pili (20). Repeating, covalently linked major pilin subunits comprise the bulk of the pilus fiber. When present, a second minor subunit is proposed to localize to the fiber tip and a third subunit is proposed to localize to the base (19). Respectively, these ancillary pilins may facilitate interaction with host proteins and the anchoring of pilus fibers to the cell wall via processing by the housekeeping sortase. Though dispensable for major pilin polymerization, minor subunits have been shown to affect aspects of pilus biogenesis, including pilus length, thickness, and subcellular compartmentalization in several sortase-assembled pilus systems (21, 22). Characteristically, the E. faecalis ebp operon encodes three structural proteins, EbpA, EbpB, and EbpC, and the pilus-associated sortase SrtC (or Bps) (8). A housekeeping sortase, SrtA, is encoded elsewhere in the genome (23). Ebp pilus fibers consist largely of the major pilin EbpC, whose polymerization is mediated by SrtC. EbpA and EbpB are covalently incorporated into the cell wall-anchored Ebp pilus fibers as minor components (8), but their roles in pilus biogenesis and pilus function have not been directly investigated previously.

Sortase-assembled pili or their subunits have been reported as virulence factors in vivo in infection models of group B streptococci (GBS; Streptococcus agalactiae) (24-26), group A streptococci (GAS; Streptococcus pyogenes) (27, 28), and the pneumococcus (Streptococcus pneumoniae) (13). A polaron, nonlipil E. faecalis ebp mutant was attenuated in animal models of infective endocarditis (8) and ascending UTI (29), as were E. faecalis mutants lacking SrtC or SrtA (30) and a nonlipil E. faecium ebp mutant (31). A mutant lacking SrtA was also attenuated in the model of CAUTI used in this study (32), but the role of Ebp pilin in CAUTI has not been directly investigated. Though the function of Ebp pilin in pathogenesis is unclear, in vitro studies suggested that Ebp pilin were important for static biofilm formation (8) and adherence to extracellular matrix (ECM) proteins (33) and human platelets (34). However, the unique contributions of EbpA, EbpB, or EbpC to any described Ebp pilus function have not been explored.

In this report, we hypothesized that the Ebp pilus is a virulence factor in CAUTI. To test this hypothesis, we investigated its importance for E. faecalis pathogenesis in a mouse model that mimics many aspects of human CAUTI (32). We evaluated the role of each Ebp structural subunit in pilus biogenesis and in virulence in CAUTI, revealing that the minor pilins mediated pilus function in vivo. Specifically, the metal ion-dependent adhesion site (MIDAS) motif in EbpA’s predicted von Willebrand factor A (VWA) domain, a common domain among sortase-assembled pili (22), was critical for E. faecalis virulence in CAUTI. Thus, this study elucidated the molecular basis of Ebp pilus function in experimental E. faecalis CAUTI.

RESULTS

Ebp pili are important in a mouse model of CAUTI. We used a previously described mouse model of E. faecalis CAUTI (32) and a chromosomal deletion mutant lacking all three Ebp structural pilins (EbpABC− strain) to investigate the importance of Ebp pilin in pathogenesis. Bladders of female mice were implanted with a short segment of silicone tubing (implant) to mimic urinary catheterization and subsequently infected with the E. faecalis OG1RF or EbpABC− strain. Twenty-four hours postinfection (p.i.), the bacterial burdens in the bladders and those associated with the implants were determined as a measure of virulence. Median CFU recovered from the bladders (Fig. 1A) and implants (Fig. 1B) of mice infected with the EbpABC− strain were significantly lower than those recovered from the bladders and implants of mice infected with OG1RF. These results demonstrated that Ebp pili were important for E. faecalis pathogenesis in experimental CAUTI.

To identify the functional subunit critical for pilus-mediated virulence in CAUTI, we created a panel of unmarked, in-frame single and double pilin deletion mutants. In the initial steps of this analysis, we evaluated (i) EbpC assembly and expression in minor pilin deletion mutants (EbpA−, EbpB−, and EbpAB− strains) and (ii) expression of minor pilin(s) in nonpiliated ebpC deletion mutants (EbpC−, EbpBC−, and EbpAC− strains).

Expression of EbpA and EbpB in the absence of pilus fibers. EbpC is the major pilus subunit necessary for fiber polymerization (8). Thus, we evaluated minor pilin expression in the absence of EbpC in Western blot analyses using anti-EbpA and anti-EbpB sera after reducing SDS-PAGE. In EbpC− cell lysates, we observed two EbpA doublets migrating at approximately 140 kDa and 100 kDa (Fig. 2A, open arrowhead and asterisk, respectively). We observed only one EbpB species migrating at a size similar to the larger ~140-kDa EbpA species (Fig. 2B, open arrowhead). Because the ~140-kDa species recognized by both anti-EbpA and anti-EbpB sera was larger than the predicted molecular masses for either EbpA or EbpB monomer after cleavage of its signal sequence and CWSS (~120 and ~46 kDa, respectively), we hypothesized that this species likely represents an EbpA-EbpB heterodimer. As formation of autocatalytic intramolecular isopeptide bonds has been shown to change the mobility of other sortase-assembled pilus subunits on SDS-PAGE (35), we predict that the ~100-kDa EbpA species is the fully processed EbpA monomer.

To test this hypothesis, we analyzed EbpA and EbpB expression in a mutant lacking all sortase genes (SrtC− SrtA− strain) to prevent formation of interpilin isopeptide bonds. In Western blots of SrtC− SrtA− cell lysates, the ~100-kDa EbpA species predominated (Fig. 2A, asterisk). The same result was obtained with our SrtC− mutant (data not shown), consistent with prior reports for OG1RF (8) and E. faecalis OG1X (36). Furthermore, we observed the ~100-kDa EbpA species in cell lysates of the EbpBC− strain (Fig. 2A, asterisk).

The ~140-kDa species was also absent when anti-EbpB sera were used to probe Western blots of SrtC− SrtA− (Fig. 2B and 2C), SrtC− (data not shown), and EbpAC− (Fig. 2B and 2C) strains. In these strains, EbpB migrated just below the 50-kDa protein marker, consistent with its predicted monomer size (Fig. 2B and 2C, hash marks). Thus, formation of the ~140-kDa EbpA and EbpB species observed in the EbpC− strain depended on EbpB and EbpA, respectively, and SrtC, consistent with the formation of an EbpA-EbpB heterodimer. Interestingly, we found EbpB mono-
and EbpAB

Others have observed that deletion of minor subunits from absent EbpC.

Taken together, these data suggest that an EbpA-EbpB heterodimer is formed by SrtC and anchored to the cell wall in the

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Finally, complementation of the EbpABC− strain with ebpA, ebpB, and ebpC provided in trans resulted in incorporation of EbpA and EbpB into high-molecular-weight pilus ladders (HMWLs) in cell wall fractions, similar to those seen in OG1RF (Fig. 2D and 2E, respectively). However, when the EbpABC− strain was provided with only ebpA and ebpB in trans, EbpA and EbpB expression mirrored that seen in the EbpC− strain: ~140-kDa and ~100-kDa EbpA species (Fig. 2D, open arrowhead and asterisk) and ~140-kDa EbpB species (Fig. 2E, open arrowhead). Taken together, these data suggest that an EbpA-EbpB heterodimer is formed by SrtC and anchored to the cell wall in the absence of EbpC.

Chromosomal deletion of minor pilins perturbs Ebp biogenesis. Others have observed that deletion of minor subunits from other sortase-assembled pilus islands can affect aspects of pilus biogenesis, including morphology and population piliation dynamics. Thus, we analyzed pilus biogenesis in the EbpA−, EbpB−, and EbpAB− minor pilin deletion mutants using anti-EbpC sera. Western blots showed that deletion of either ebpA or ebpB did not prevent expression of EbpC (Fig. 3A and 3C, respectively) or expression of the remaining minor subunit (Fig. 3B and 3D, respectively). However, the EbpC species detected in EbpA− and EbpAB− cell lysates was a compressed band that barely migrated into gels (Fig. 3A). This differed from the EbpC pilus HMWL observed in OG1RF. In contrast, EbpC HMWLs from the EbpB− and OG1RF strains were indistinguishable (Fig. 3C). To test whether ebpA provided in trans could restore OG1RF-like EbpC HMWLs in EbpA− and EbpAB− strains, ebpA was placed under the control of a tetracycline-inducible promoter on a plasmid. High- and low-level EbpA expression was observed from this plasmid with induction (100 μg ml−1 anhydrotetracycline) and no induction, respectively, in the EbpABC− strain. However, EbpA expressed from this plasmid with or without induction was not incorporated into and did not affect compressed EbpC bands in the EbpA− and EbpAB− strains (data not shown). Thus, compressed EbpC bands caused by deletion of ebpA indicated altered pilus morphology that could not be complemented in trans.

We next investigated pilus morphology in minor pilin deletion mutants using deep-etch immunogold electron microscopy (EM). In the OG1RF and EbpB− strains, multiple pilus fibers several hundred nanometers long were observed associated with bacterial cells (Fig. 3E, arrowheads). In contrast, in cultures of the EbpA− and EbpAB− strains, we observed fewer but much longer EbpC fibers (greater than 1 μm) associated with some cells (Fig. 3E, large arrows). An analysis of population piliation dynamics using immunofluorescence microscopy (IFM) also revealed defects of the minor pilin deletion strains in pilus biogenesis. A significantly smaller median proportion of EbpA−, EbpB−, and EbpAB− bacterial cells (1.1%, 33.9%, and 2.8%, respectively) expressed EbpC at their surface than did OG1RF (65.8%) (Fig. 3F).

Minor pilin deletion mutants are attenuated in experimental E. faecalis CAUTI. We then investigated the minor pilin deletion mutants in experimental CAUTI. EbpA−, EbpB−, and EbpAB− strains were all significantly attenuated in bladder colonization compared to OG1RF (Fig. 1A). EbpA− and EbpAB− strains were also attenuated in implant colonization (Fig. 1B). The defect of
these strains in vivo may have been due to the lack of EbpA and/or EbpB or to the effects of the minor pilin deletions on pilus biogenesis described in the studies above.

Expression of EbpA and EbpB in the EbpC− strain is sufficient for pilus-mediated virulence in CAUTI. Interestingly, the nonpiliated EbpC− mutant and OG1RF colonized the bladders and implants of infected mice to similar levels (Fig. 1A and 1B), showing that EbpA and EbpB expression in nonpiliated strains. Western blot analysis described in the studies above.

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by the resultant strain were similar to those observed from the EbpABC\(^{-}\) SrtC\(^{-}\) strain complemented with the unmutated OG1RF locus (p-ebpABC\(_{\text{SrtC}}\)) as determined by negative-stain immunogold EM (Fig. 5A). We next introduced the mutant MIDAS motif into the chromosomal ebpA locus in OG1RF. Pili produced by this MIDAS mutant (EbpAAWAGA\(^{\text{BCsrtC}}\)) and by OG1RF were also similar when examined with negative-stain immunogold EM (Fig. 5B). Similarly, no difference between EbpAAWAGA and OG1RF was observed in an IFM analysis of population pilation dynamics (Fig. 5C). Furthermore, HMWLs of EbpAAWAGA and OG1RF were indistinguishable on Western blots probed with any antipilin immune serum (Fig. 5D). Thus, mutating EbpA’s MIDAS motif did not affect pilus biogenesis.

EbpA MIDAS motif mutants are attenuated in experimental CAUTI. We next examined MIDAS motif mutants in experimental *E. faecalis* CAUTI. There were no significant differences between the median 24-h bladder or implant bacterial titers of mice infected with strains expressing pili with unmodified MIDAS motifs: OG1RF/pGCP123 (empty vector) and EbpABC\(^{-}\) SrtC\(-/\) p-ebpABC\(_{\text{SrtC}}\) strain. Twenty-four-hour bacterial titers from mice infected with the nonpiliated EbpABC\(^{-}\) SrtC\(-/\)pGCP123 strain and with the EbpABC\(^{-}\) SrtC\(-/\)p-ebpAAWAGA\(_{\text{BCsrtC}}\) MIDAS motif mutant also did not differ significantly from each other. However, infections with either of these strains resulted in significantly lower bladder and implant bacterial burdens 24 h p.i. than those of OG1RF/pGCP123 or the EbpABC\(^{-}\) SrtC\(-/\) p-ebpABC\(_{\text{SrtC}}\) strain (Fig. 6A and 6B). Thus, supplying the OG1RF ebpABC\(_{\text{SrtC}}\) locus to the EbpABC\(^{-}\) SrtC\(^{-}\) strain in *trans* fully complemented its virulence defect in CAUTI, while provision of an ebpAAWAGA\(_{\text{BCsrtC}}\) mutant MIDAS motif locus in *trans* resulted in attenuation in CAUTI similar to that seen with provision of the empty vector alone.

Similarly, when we implanted and infected mice with the chromosomal MIDAS motif mutant (EbpAAWAGA\(^{\text{BCsrtC}}\)) and monitored the bacterial burdens both 24 h (Fig. 7A and 7B) and 7 days (Fig. 7C and 7D) p.i., the EbpAAWAGA and EbpABC\(^{-}\) strains were similarly attenuated in both bladder (Fig. 7A and 7C) and implant (Fig. 7B and 7D) colonization at each time point tested compared to OG1RF, revealing that EbpA’s MIDAS motif is necessary for Ebp pilus function in experimental *E. faecalis* CAUTI.

**Ebp pilus importance in UTI is tissue and model specific.** There were no significant differences between the median bacterial kidney titer of mice infected with any pilin mutant and that of mice infected with OG1RF at any time point in experimental CAUTI (see Fig. S1 in the supplemental material). Thus, the importance of Ebp pili in this model was specific to implants and bladder infection.

![FIG 4](https://mbio.asm.org) The SrtC\(^{-}\) mutant is attenuated in experimental CAUTI. Mice were infected with ~2 \times 10^6 CFU of OG1RF (closed circles) or the nonpiliated SrtC\(^{-}\) mutant (open hexagons). Bacterial titers 24 h p.i. in the bladders (A) and implants (B) from 2 independent experiments are shown. Each shape corresponds to one mouse. Median titers (CFU/bladder, CFU/implant) are shown with a bar: OG1RF (2.16 \times 10^7, 4.90 \times 10^7) and SrtC\(^{-}\) strain (1.18 \times 10^7, 20). Dashed lines are limits of detection (40 CFU/bladder; 20 CFU/implant). Statistically significant differences between OG1RF and SrtC\(^{-}\) titers are shown (***, \(P < 0.001\)).

![FIG 5](https://mbio.asm.org) Mutation of EbpA’s MIDAS motif does not affect pilus biogenesis. Mouse anti-EbpC polyclonal sera were used to assess pilus biogenesis by negative-stain immunogold EM (A and B), IFM (C), and Western blot analysis (D) after SDS-PAGE of the indicated cell fractions. (A and B) Piliation of the MIDAS motif mutant strains (EbpABC\(^{-}\) SrtC\(-/\)p-ebpAAWAGA\(_{\text{BCsrtC}}\) strain and EbpAAWAGA\(^{\text{BCsrtC}}\)) was similar to that of control strains (EbpABC\(^{-}\) SrtC\(-/\)p-ebpABC\(_{\text{SrtC}}\) strain and OG1RF, respectively). Bars, 500 nm. (C) Comparison of the median percentages of EbpC\(^{-}\) bacterial cells in OG1RF (50%) and EbpAAWAGA (38%) from 2 independent experiments revealed no significant differences in population pilation dynamics. Whiskers show the 10th and 90th percentiles; dots show outliers (ns, not significant). (D) Pilus HMWLs on Western blots of OG1RF and EbpAAWAGA probed with anti-EbpC (left), anti-EbpB (middle), and anti-EbpA (right) sera were indistinguishable.
implanted bladders. To further investigate the tissue specificity of Ebp pili, we compared the EbpABC− and EbpC− strains to OG1RF in a previously described mouse model of ascending UTI (38) in which *E. faecalis* displays kidney tropism (see Text S1 for methods). Since growth in serum was reported to increase expression of Ebp pili (8), we tested two inoculum preparation methods: bacterial subculture in brain heart infusion (BHI) broth alone or with 40% horse serum. Each mutant fared as well as or better than OG1RF in bladders and kidneys at both 6 h and 48 h p.i. using either inoculum preparation method (see Fig. S2 in the supplemental material). Thus, Ebp pili were dispensable for *E. faecalis* virulence in our model of ascending UTI using mice of the same age and genetic background as those used in our model of CAUTI (7- to 8-week-old C57BL/6 mice). Consistent with the model specificity of Ebp pili in UTI pathogenesis observed here, Singh et al. previously showed a defect for a polar, nonpiliated ebp mutant in a different model of ascending UTI using younger, outbred mice, different bacterial inoculum preparation methods, and distinct virulence metrics (29).

**DISCUSSION**

In this study, we demonstrated a specific role for Ebp pili in a newly characterized mouse model of *E. faecalis* CAUTI. To investigate the molecular basis of pilus function *in vivo*, we described the effects of an extensive panel of pilus structural subunit deletions on both pilus biogenesis and pilus function in experimental CAUTI. Finally, we showed that EbpA and the MIDAS motif encoded by its predicted VWA domain were critical for pilus-mediated virulence in *E. faecalis* CAUTI.

The EbpABC− mutant lacking all structural subunits was severely attenuated in bladder and implant colonization in experimental CAUTI, showing that the Ebp pilus was an important virulence factor in this model. The residual bladder and implant colonization by the EbpABC− strain suggests that additional bacterial factors may play a role in *E. faecalis* CAUTI. To investigate

**FIG 6** Transformation of EbpABC− SrtC− strain with p-ebpABCsrtC or pGCP123 complemented its virulence defect in experimental CAUTI. Bacterial titers 24 h p.i. of the bladders (A) and implants (B) from 2 independent experiments are shown. Each shape corresponds to one mouse; open shapes represent a nonpiliated bacterial strain. Median titers (CFU/bladder, CFU/implant) are shown with a bar: OG1RF/pGCP123 (5.92 × 10², 1.96 × 10³), EbpABC− SrtC−/pGCP123 strain (1.94 × 10⁴, 1.72 × 10⁴), EbpABC− SrtC−/p-ebpABCsrtC strain (2.60 × 10⁴, 1.20 × 10⁴), and EbpABC− SrtC−/p-ebpAWAGAABC strain (1.20 × 10⁴, 5.80 × 10³). Dashed lines are the limits of detection (10 CFU/bladder, 5 CFU/implant). All possible strain pairs were compared statistically; P values were adjusted for 6 comparisons. Significant differences are shown (*, P < 0.05; **, P < 0.01; ***, P < 0.001).

**FIG 7** The chromosomal MIDAS mutant (EbpAWAGA) was as attenuated as the EbpABC− strain in experimental CAUTI. Twenty-four-hour (A and B) or 7-day (C and D) viable bacterial counts from the bladders (A and C) and implants (B and D) of mice infected with *E. faecalis* were pooled from 2 to 3 independent experiments for each strain. Each shape corresponds to one mouse; open shapes represent a nonpiliated bacterial strain. Median bacterial titers (CFU/bladder, CFU/implant) were determined at 24 h p.i. for OG1RF (3.16 × 10², 3.24 × 10³), EbpAWAGA (2.40 × 10⁴, 1.80 × 10³), and EbpABC− (3.20 × 10⁴, 5) strains. Bars are medians; dashed lines are the limits of detection (10 CFU/bladder, 5 CFU/implant). All possible strain combinations were compared statistically; P values were adjusted for 3 comparisons. Significant differences are shown (*, P < 0.05; ***, P < 0.001).
the molecular basis of pilus function in vivo, we characterized a panel of pilin deletion mutants. We found that deletion of ebpA altered pilus morphology, leading to extended EbpC fibers. Similarly, deletion of either the pilA or pilC minor pilin gene from the GBS NEM316 pilus island 2A (PI-2A) led to longer pilus fibers (22). In Corynebacterium diphtheriae, mutants lacking the minor anchor pilin (SpaB) produced longer pilus fibers, presumably because cell wall anchoring achieved via SpaB processing by the housekeeping sortase prevents further polymerization by the pilus-associated sortase (21, 39). However, pili of our mutant lacking EbpB, the predicted Ebp base pilin, were not appreciably different from pili of OG1RF in our study. As overexpression of the major pilus has also been shown to increase pilus length (40), it is possible that the ebpA deletion affected relative EbpC levels.

We demonstrated that the deletion of either or both minor pilins reduced the proportion of pilated E. faecalis cells in a population by an unknown mechanism. Many sortase-assembled pilus islands include a divergently transcribed, upstream positive regulator, ebpR in the case of the E. faecalis ebp operon (41). Additionally, deletion of rnrB, a putative RNase J2, reduced pilus expression and levels of ebpABC mRNA transcript (42). Interestingly, it was recently reported that RrgA, the tip pilin of the S. pneumoniae rlrA pilus islet, interacts with RlrA, the upstream positive regulator of pilus expression, to exert a negative effect on population piliation dynamics of S. pneumoniae (43), presenting a mechanism whereby a structural pilin affected population piliation dynamics. Future studies will determine how the ebpA and ebpB deletions affected pilus biogenesis and whether they interacted with ebpR or rnrB to do so. Not surprisingly, the minor pilin deletion mutants (EbpA−, EbpAB−, and EbpB− strains), which all exhibited perturbed pilus biogenesis, were attenuated in experimental CAUTI.

Interestingly, the nonpiliated EbpC− mutant behaved similarly to OG1RF in experimental CAUTI, showing that the major polymerizing EbpC subunit and pilus fibers were dispensable for E. faecalis virulence. Similarly, the major subunits, but not the minor RrgA and Cpa tip pilins, were dispensable for pneumococcal mouse upper airway colonization (44) and GAS skin colonization in a humanized mouse model (27), respectively. Indeed, minor pilins expressed in the absence of pilus fibers have been shown to govern several sortase-assembled pilus functions in vitro, including adherence to cell lines and static biofilm formation (22, 44, 45). It has thus been suggested that pilus fibers serve to extend a minor functional pilin beyond the bacterial capsule where it can interact with host molecules (22). In this case, since OG1RF does not produce the E. faecalis capsular polysaccharide (46), EbpC pilus fibers, but not a functional minor subunit, would be dispensable for pilus function in OG1RF, just as we observed in experimental CAUTI.

Our analysis of minor pilin expression in ebpC and sortase mutants argued that a sortase-assembled EbpA−EbpB heterodimer was expressed in the absence of pilus fibers in the EbpC− strain. Similarly, the C. diphtheriae SpaC and SpaB minor pilins heterodimerized and anchored to the cell wall in the absence of the major pilin in a sortase-dependent fashion (47). The nonpiliated EbpAC−, EbpBC−, and SrtC− mutants that expressed mainly EbpA and/or EbpB monomers, but not the putative EbpA-EbpB heterodimer, were severely attenuated in experimental UTI, suggesting that sortase-assembled EbpA and/or EbpB mediated Ebp pilus function.

To directly test the importance of the minor pilins in CAUTI, we sought to create mutations in functional domains that did not affect pilus biogenesis. EbpA contains a predicted Cna B domain, not investigated here, and a VWA domain. VWA domains, named only for their role in platelet adhesion to damaged vascular endothelium by the human plasma protein von Willebrand factor (48), are widely distributed among archaea, bacteria, and eukaryotes. Well-studied examples occur in some integrins, ECM proteins, and magnesium (Mg) chelatases and perform diverse functions, usually protein-protein interaction or cell adhesion (49). Coordination of a divalent cation by a VWA motif, present in almost all of all VWA domains, is critical for the function of some VWA domain-containing proteins (49). Most prokaryotic VWA domains have not been investigated in detail. However, Konto-Ghiorgi et al. showed that the GBS PilA tip pilin VWA domain was important for pilus-mediated bacterial adhesion to human alveolar and intestinal epithelial cells in vitro (22). Furthermore, the crystal structure of the pneumococcal RrgA tip pilin modeled an Mg2+ ion coordinated by the VWA motif of its VWA domain (50). We therefore hypothesized that EbpA’s MIDAS motif would be important for Ebp pilus function in our model of CAUTI. Indeed, MIDAS motif mutants were as attenuated in vivo as were the relevant nonpiliated control strains, showing that an intact MIDAS motif is necessary for Ebp pilus function in bladder and implant colonization in experimental E. faecalis CAUTI. To our knowledge, this is the first study ascribing a sortase-assembled pilus function in vivo in a disease model to a specific protein domain. The importance of a MIDAS motif for the function of a prokaryotic VWA domain-containing protein has otherwise been shown only for the Rhodotorula capsulatus Mg chelatase BchD subunit (51), a member of an evolutionarily distinct family of VWA domain-containing proteins (49).

The functional role of the Ebp pilus governed by EbpA’s VWA domain and MIDAS motif in bladder and implant colonization in experimental CAUTI remains to be determined. However, this function was tissue and model specific since pilin mutants colonized kidneys similarly to OG1RF in experimental CAUTI and both kidneys and bladders in our mouse model of ascending UTI. MIDAS motifs in the integrin beta and some alpha subunits are involved in integrin binding to ECM proteins (49). The VWA domain- and MIDAS motif-containing tip pilins RrgA and PilA have both been reported to bind ECM proteins such as collagen (26, 52). Furthermore, PilA’s interaction with collagen is a critical component of GBS virulence in a mouse model of hemorrhagic meningitis (26). Crude cell wall extracts of a distinct nonpiliated E. faecalis mutant demonstrated reduced adherence to purified human collagens and fibrinogen compared to those of OG1RF, implicating Ebp pilin in adhesion to these ECM molecules (33). Binding of bacteria to ECM proteins exposed by damage to vascular endothelium initiates infective endocarditis, an enterococcal disease in which Ebp pili are also implicated (8). In our CAUTI model, implantation leads to physiological changes in the bladder epithelium and induction of inflammation (32), potentially revealing host binding partners, such as ECM proteins, for recognition by EbpA. Colonization of implants may proceed by the same mechanism, as urinary catheters become coated with host proteins and components (53). Alternatively, EbpA’s VWA domain may perform a distinct behavior that facilitates in vivo biofilm formation on implants. Polar ebp disruption mutants showed reduced static biofilm formation in vitro (8), suggesting that Ebp pili...
may be involved in adherence to abiotic surfaces or bacterial surface components.

The diversity of bacterial species with sortase-assembled pili is matched only by the variety of potential niches and disease processes in which these pili function. However, a particular pilin or pilin domain is implicated for only a few of these behaviors. By mutating the predicted metal ion-coordinating amino acids of the MIDAS motif in EbpA’s VWA domain, we preserved pilus biogenesis and showed a clear role for this motif in the function of the Ebp pilus in vivo. Future studies will determine whether the VWA domains of EbpA and other sortase-assembled tip pilins function similarly to those of the integrin subunits or other characterized MIDAS motif-containing VWA domains, allowing the development of structure-function correlates for pilus-mediated virulence in a wide variety of diseases.

MATERIALS AND METHODS

Bacterial strains and growth conditions. Bacterial strains are listed in Table S1 in the supplemental material. Unless otherwise noted, we grew *Escherichia coli* in Difco LB (Luria-Bertani) broth at 37°C with agitation and *E. faecalis* OG1RF (ATCC 47077) and its derivative strains statically at 37°C in Bacto BHI broth with rifampin (Rif; 25 to 100 μg ml⁻¹). Plasmid-containing strains were grown with appropriate antibiotics (see Table S2 in the supplemental material). For *E. coli*, erythromycin (Erm) was added at 500 μg ml⁻¹, kanamycin (Kan) at 50 μg ml⁻¹ (25 μg ml⁻¹ for pREP4), and ampicillin (Amp) at 100 μg ml⁻¹. For *E. faecalis*, Erm was added at 25 μg ml⁻¹ and Kan at 500 μg ml⁻¹. All media were purchased from BD (Becton, Dickinson and Company, Franklin Lakes, NJ). Antibiotics were purchased from Sigma-Aldrich Corporation (St. Louis, MO).

General cloning techniques. DNA and amino acid sequences were retrieved and analyzed as described in Text S1 in the supplemental material. Bacterial genomic DNA (gDNA) was isolated with the Wizard Genomic DNA purification kit (Promega Corp., Madison, WI). Plasmids are listed in Table S2 in the supplemental material. Aside from pABG5, pGCP123, and their derivatives, which were purified with the Hurricane Maxi Prep kit (Gerard Biotech LLC, Oxford, OH), plasmid DNA was isolated with the Wizard Plus SV Miniprep DNA purification system (Promega Corp.). Primers are listed in Table S3 in the supplemental material. PCR was performed with Phusion DNA polymerase from Finnzymes (Thermo Fisher Scientific, Inc., Rockford, IL). Site-directed mutagenesis (SDM) was carried out with QuickChange or QuickChange II kits (Stratagene, La Jolla, CA). Restriction endonucleases and T4 DNA ligase were purchased from New England Biolabs (Ipswich, MA). Ligation reactions were transformed into *E. coli* TOP10, NovaBlue, or XL1-Blue cells. Plasmids derived here were confirmed by sequencing of inserts.

Generation of chromosomal deletion strains. Deletion of pilin coding sequences from the OG1RF chromosome was accomplished by allelic replacement as described previously using the temperature-sensitive shuttle vector pJR233 (36). The ΔebpA, ΔebpB, ΔebpC, ΔebpAB, ΔebpABC, ΔebpABCstrC, and ΔstrC deletion alleles comprised upstream (US) and downstream (DS) DNA fragments (~1,000 bp each) of the region to be deleted that were spliced by overlap extension PCR (SOE-PCR) (54). Single gene deletion alleles removed protein-coding open reading frame (ORF) sequences of the deleted gene only, preserving any overlapping ORF or intergenic sequence. Alleles for multiple gene deletions preserved any overlapping and intergenic sequence between remaining ORFs and deleted ORFs but removed any intergenic sequence between consecutively deleted ORFs. Table S4-A in the supplemental material details allele construction. Each allele was cloned into pJR233 as described for each resultant plasmid listed in Table S2-G. When noted in Table S4-A, alleles were first blunt end ligated into a commercial vector. The *EbpA*⁺, *EbpB*⁺, *EbpC*⁺, *EbpAB*, *EbpABC*, and *SrtC*⁺ strains were created by transformation of OG1RF with pSJH-529, pSJH-530, pSJH-523, pSJH-531, pSJH-532, and pSJH-189, respectively, by electroporation. The *EbpAC*⁻, *EbpABC*⁻ *SrtC*⁻, and *SrtC*⁻ *SrtA*⁻ strains were created by transformation of EbpC⁺*, EbpB⁺*, and EbpA⁺* strains with pSJH-529, pSJH-279, and pSJH-189, respectively. Transformants were selected with Erm at 30°C and then passaged at the nonpermissive temperature (42°C) with Erm to select for chromosomal integration of the plasmid. Subsequent plasmid excision was allowed by passage at 30°C without Erm. Erm-sensitive colonies with double-crossover events that integrated deletion alleles were selected by PCR screening. Deletions were confirmed by PCR with gDNA.

Chromosomal MIDAS motif mutant construction. Mutation of EbpA’s MIDAS motif on the OG1RF chromosome was accomplished by allelic replacement with the *ebpAAWAGA* allele (coding for Ala315-Trp-Ala317-Gly-Ala319 in EbpA; see Table S4-B in the supplemental material) using pGCP213 instead of pJR233. Creation of the pcGCP213 temperature-sensitive shuttle vector is described in Text S1 in the supplemental material. The *ebpAAWAGA* allele was introduced into pGCP213 as described for pSJH-509 in Table S2-H. An extraneous nucleotide added by primers HVN228 and HVN229 was deleted by SDM using primers HVN241 and HVN242, resulting in pSJH-509. OG1RF was transformed with pSJH-509 and passaged as described above. We screened for incorporation of the mutant allele by BspEI digestion of internal ebpA colony PCR products. A BspEI restriction site in the OIG1RF allele is absent in the *ebpAAWAGA* allele. The resultant *EbpAAWAGA* chromosomal MIDAS motif mutant was confirmed by gDNA sequencing.

Generation of *E. faecalis* expression strains. Plasmids were created for expression of ebp genes in trans in *E. faecalis* using the Gram-positive expression vector pGCP123 (derived here as described in Text S1 in the supplemental material). All pGCP123 derivatives included the region 500 bp upstream of EbpA’s translational start codon as the putative ebpA promoter (ebpAp). See Table S2-I for construction details of the following plasmids: pSJH-491 (p-ebpABC) encodes all structural pilins, pSJH-492 (p-ebpAB) encodes just EbpA and EbpB, pSJH-496 (p-ebpABCstrC) encodes all structural pilins and StrtC, and pSJH-559 (p-ebpA*AWAGA*B-CartC) encodes the *ebpA*AWAGA allele of EbpA and OIG1RF alleles of EbpB, EbpC, and StrtC. EbpABC⁺ and EbpABC⁻ StrtC⁻ strains were transformed with empty pGCP123 (see Table S1-C) and the plasmids derived above (see Table S1-D).

Generation of polyclonal antisera. Expression constructs for each Ebp pilin lacking the signal sequence and CWSS (EbpA-X, EbpB-X, and EbpCA-X) were created as described in Table S4-D in the supplemental material and cloned into the isopropyl-d-1-thiogalactopyranoside (IPTG)-inducible expression vector pQE-30Xa (Qiagen Inc., Valencia, CA), resulting in the addition of an N-terminal G5XHis tag. The Ebp-X construct comprised roughly the C-terminal half of EbpA. When noted in Table S4-D, constructs were first blunt end ligated into a commercial vector. The resultant pQE-30Xa-derived plasmids encoding recombinant EbpA, EbpB, and EbpC (pSJH-541, pSJH-547, and p-SJH-550, respectively) were used to transform the *E. coli* expression strain M15/ pREP4 or SG13009/pREP4, resulting in strains SJH1987, SJH1988, and SJH1985. Text S1 describes the purification of recombinant EbpA, EbpB, and EbpC. Polyclonal antisera were generated commercially by immunization of New Zealand White rabbits with purified, recombinant EbpA or EbpB (New England Peptide, Gardner, MA) and by immunization of mice with purified, recombinant EbpC (Agro-Bio, La Ferté Saint-Aubin, France). Specificities of the immune sera were confirmed by a lack of signal on Western blots of cell lysates from the appropriate deletion mutant strains.

Western blots. Bacterial cell fractions (prepared as described in Text S1 in the supplemental material) were boiled for at least 10 min in β-mercaptoethanol-containing loading buffer, and SDS-PAGE was performed with NuPAGE Novex 3 to 8% Tris-acetate gels in NuPAGE Tris-acetate SDS running buffer (Life Technologies Corp., Carlsbad, CA). Proteins were transferred to nitrocellulose membranes (Bio-Rad, Hercules, CA) and incubated with 5% nonfat dry milk in Tris-buffered saline containing 0.05% Tween 20. Membranes were probed with antipilin sera as indicated and Pierce stabilized horseradish peroxidase-conjugated goat anti-rabbit or goat anti-

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mouse IgG (Thermo Fisher Scientific, Inc.). Blots were developed with SuperSignal West Femto chemiluminescent substrate (Thermo Fisher Scientific, Inc.); film was processed with a Kodak X-Omat processor. Precision Plus Protein Kaleidoscope Standards (Bio-Rad Laboratories, Inc., Hercules, CA) are indicated. Distinct blots or exposures are separated by white space in the figures; lines represent an irrelevant gel lane that was removed using Adobe Photoshop CS2 (Adobe Systems Inc., Mountain View, CA).

IFM. IFM was performed as described previously (36). Slides were labeled with mouse anti-EbpC sera and 18-nm gold bead-conjugated goat anti-mouse secondary antibody (Jackson ImmunoResearch Laboratories Inc., West Grove, PA). Samples were freeze-dried and imaged as described elsewhere (55, 56). The Shadows/Highlights function of Adobe Photoshop CS2 (Adobe Systems) was applied to Fig. 3E as a whole.

Negative-stain immunogold EM. Bacteria grown overnight were diluted 1:1,000 into TSBG (BBL Trypticase soy broth with 0.25% glucose) were deposited onto glass slides, fixed, and labeled as described previously (55) using mouse anti-EbpC sera and 18-nm gold-bead-conjugated goat-anti-mouse secondary antibody (Jackson ImmunoResearch Laboratories Inc., West Grove, PA). Images were analyzed using Axiovision software and a Zeiss Axioskop 2 MOT Plus wide-field fluorescence microscope at the Department of Molecular Microbiology Imaging Facility of Washington University in St. Louis, MO. Quantification of EbpC-expressing cells is described in Text S1 in the supplemental material.

Statistical analyses. Data from multiple experiments were pooled. Two-tailed Mann-Whitney U tests were performed with GraphPad Prism 5 software (GraphPad Software, San Diego, CA) for all comparisons described in CAUTI, ascending UTI, and IFM experiments. When noted in the figure legends, Bonferroni’s adjustment for multiple comparisons was performed manually. An adjusted P value of <0.05 was considered statistically significant.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.orglookup/suppl/doi:10.1128/mBio.00177-12/-/DCSupplemental

Table S3, PDF file, 0.2 MB. Table S4, PDF file, 0.1 MB.

ACKNOWLEDGMENTS

This work was supported by the NIDDK grant DK51406 and its ARRA supplement DK51406-12S1 awarded to S.J.H., the American Heart Association Midwest Affiliate Predoctoral Fellowship 10PRE2640099 to H.V.N., an ASM Robert D. Watkins Graduate Research Fellowship Award to P.S.G., and a W. M. Keck Postdoctoral Fellowship to G.C.P.

We thank D. K. Robertson, R. Roth, and J. E. Heuser for assistance with EM studies. We are grateful to K. W. Dodson for critical reading of the manuscript and to members of the Hultgren, Caparon, and Normark laboratories for helpful insight and discussion.

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