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The Extracellular Matrix of *Staphylococcus aureus* Biofilms Comprises Cytoplasmic Proteins That Associate with the Cell Surface in Response to Decreasing pH

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**ABSTRACT** Biofilm formation by *Staphylococcus aureus* involves the formation of an extracellular matrix, but the composition of this matrix has been uncertain. Here we report that the matrix is largely composed of cytoplasmic proteins that reversibly associate with the cell surface in a manner that depends on pH. We propose a model for biofilm formation in which cytoplasmic proteins are released from cells in stationary phase. These proteins associate with the cell surface in response to decreasing pH during biofilm formation. Rather than utilizing a dedicated matrix protein, *S. aureus* appears to recycle cytoplasmic proteins that moonlight as components of the extracellular matrix.

**IMPORTANCE** *Staphylococcus aureus* is a leading cause of multiantibiotic-resistant nosocomial infections and is often found growing as a biofilm in catheters and chronic wounds. Biofilm formation is an important pathogenicity strategy that enhances resistance to antimicrobials, thereby limiting treatment options and ultimately contributing to increased morbidity and mortality. Cells in a biofilm are held together by an extracellular matrix that consists in whole or in part of protein, but the nature of the proteins in the *S. aureus* matrix is not well understood. Here we postulate that *S. aureus* recycles proteins from the cytoplasm to form the extracellular matrix. This strategy, of cytoplasmic proteins moonlighting as matrix proteins, could allow enhanced flexibility and adaptability for *S. aureus* in forming biofilms under infection conditions and could promote the formation of mixed-species biofilms in chronic wounds.

**RESULTS**

Identification of biofilm-associated cell surface proteins. *S. aureus* strain HG003 forms robust biofilms in tryptic soy broth...
(TSB) with 0.5% added glucose (here TSBG) (see Fig. S1A in the supplemental material). Both DNA and protein components of the matrix were found to be important for biofilm formation (see Fig. S1A). However, exopolysaccharide and the biofilm-implicated cell wall proteins, protein A (10) and fibrinogen-binding proteins A and B (11), did not contribute significantly (see Fig. S1B).

We took an unbiased approach to identify cell surface-associated proteins present under biofilm-inducing conditions by adapting a $^{14}$N/$^{15}$N metabolic labeling procedure used in a previous proteomic analysis of *S. aureus* (12, 13). Cell surface-associated proteins were tagged by biotinylation, allowing their specific isolation and identification (13). The ratio of $^{14}$N (biofilm) to $^{15}$N (nonbiofilm) peptides for each protein gave an estimate of the abundance of a given protein under each growth condition (see Table S1 in the supplemental material).

Interestingly, despite our selecting specifically for extracellular proteins by biotinylation, most of the proteins identified were predicted cytoplasmic proteins. We established a cutoff value for proteins considered to be enriched during biofilm formation as those having a $^{14}$N/$^{15}$N ratio greater than 2. These 11 proteins are characterized by roles in central metabolism, such as glycolysis and fermentation (14). Consistent with our findings, several of these proteins (for example, enolase, transketolase, acetolactate synthase, GAPDH [glyceraldehydes-3-phosphate dehydrogenase], and lactate dehydrogenase) were previously found to be abundant under biofilm-inducing conditions and in the biofilm exoprotoeme (15–17) in studies that did not investigate their role in biofilm formation. In addition to identifying proteins that were enriched under biofilm-inducing conditions, we also detected extracellular proteins that were either more abundant under the non-biofilm-inducing conditions or showed equal levels of abundance under the two sets of conditions (having a $^{14}$N/$^{15}$N ratio of between 0.5 and 2) (see Table S1 in the supplemental material).

We focused on two of the identified proteins, enolase and GAPDH, and asked whether we could observe association of these proteins with the cell surface by a method that did not depend on biotinylation. Cells harvested from a mature biofilm were washed in phosphate-buffered saline (PBS) at 4°C for 1 h before the cells were removed by mild centrifugation. Among the proteins that were seen in the buffer fraction, by SDS-PAGE (Fig. 1A, lane 1), were enolase and GAPDH, as judged by immunoblot analysis (Fig. 1B).

**Do cell surface-associated, cytoplasmic proteins arise from cell lysis during sample preparation?** A possible explanation for the presence of cytoplasmic proteins in our cell surface samples was cell lysis or enhanced permeability during enrichment of surface-associated proteins from cells. We took three approaches to investigate this possibility. First, we asked whether cells become permeable upon harvesting from the biofilm and resuspension in PBS using LIVE/DEAD staining. This analysis indicated that the percentage of permeable cells was unaltered for 1 h in PBS (Fig. 2A).

Second, we monitored cell lysis as the decrease in optical density at 600 nm (OD$_{600}$) with time after resuspension in the presence of Triton X-100, a detergent that stimulates autolytic activity in *S. aureus*. Resuspension of cells from a mature biofilm in PBS had little to no effect on the rate of lysis, whether stimulated by Triton X-100 or not (Fig. 2B). The rate of stimulated autolysis (decrease in OD$_{600}$ with time) was very low in cells harvested from the biofilm compared to cells from exponentially growing cultures in the same medium (Fig. 2B), suggesting that biofilm cells are not significantly prone to lysis and that extraction with PBS does not stimulate lysis.

Finally, we asked whether cell surface-associated cytoplasmic proteins arose during biofilm formation or from lysis in PBS. Proteinase K was added upon inoculation, and then a protease inhibitor was added 1 h prior to harvesting the biofilm cells. The results showed that only a very small amount of protein was detected in the PBS extract of biofilm cells (Fig. 2C). When we “spiked in” a cell lysate after the addition of inhibitor, proteins from the lysate were readily detected in the PBS extract, indicating the effectiveness of the inhibitor. The results are consistent with the idea that the cell surface-associated proteins were released from cells during biofilm formation when they accumulated on the outer side of cells and not during our sample preparation procedure. It should, however, be noted that since treatment with proteinase K inhibits biofilm formation (see Fig. S1A in the supplemental material), presumably through degradation of the protein matrix, it is possible that the treated cells could be physiologically different and thus less prone to subsequent cell lysis.

**Protein retention at the cell surface reversibly depends on pH.** Next, we asked what property of PBS buffer might be responsible for the release of cell surface-associated proteins from cells from the biofilm into the surrounding milieu. We reasoned that a change in pH from the acidic conditions found within biofilms...
to pH 7.5 in PBS might explain our results. In agreement with previous observations, we found that biofilm formation was accompanied by a significant decrease in growth medium pH which was dependent on the availability of glucose (see Fig. S2A in the supplemental material) and that a decrease in pH was required for biofilm formation (see Fig. S2B).

These results led us to hypothesize that cytoplasmic proteins associate with the cell surface to form a protein matrix in a manner that depends on low pH. To investigate this hypothesis further, we incubated biofilm cells in PBS buffer at pH 5. Little to no protein could be detected in the resulting extract, in contrast to the results seen when cells from the same biofilm sample were incubated at pH 7.5 (Fig. 3A, compare lanes 1 and 3). This was not due to protein degradation at pH 5 since proteins could still be extracted at pH 7.5 from biofilm cells that had been incubated at pH 5 for 1 h (Fig. 3A, compare lanes 1 and 2). Proteins remained cell associated at pH 5 but were released upon exposure to higher pH.

We found that proteins added exogenously (not originating from cells within the biofilm itself) could also be “captured” through an interaction with biofilm cells (Fig. 3B). Whole-cell lysate from HG003 cells was added to mature biofilm cell cultures, and the biofilm cells were then incubated at pH 5 and pH 7.5. Proteins (comprising those naturally occurring in the biofilm and those added) (Fig. 3B, compare lanes 4 and 5) were released from the cell surface at pH 7.5 and not at pH 5 (Fig. 3B, compare lanes 2 and 5). Proteins from exogenously added cell-free lysate were equally recovered from PBS at pH 5 and pH 7.5 (Fig. 3B, lanes 3 and 6), indicating that, in the absence of biofilm cells, proteins do not precipitate at pH 5 but remain in aqueous solution. This suggests that, in the presence of cells from a biofilm, exogenously added lysate proteins interact with the cell surface at low pH and are then released at pH 7.5.

To investigate the specificity of protein interactions with the cell surface, we next determined whether a heterologous protein,
green fluorescent protein (GFP), could interact with the surface of biofilm cells. Cells that had been grown under biofilm-forming conditions were incubated in PBS at pH 5 and pH 7.5 in the presence of lysate proteins isolated from a strain of HG003 that constitutively produces GFP (LCF81). After removal of the cells, GFP was detectable in PBS by immunoblotting with an anti-GFP antibody (Ab) at pH 7.5 but not at pH 5 (Fig. 3C). When cells from PBS at pH 5 were subsequently incubated in PBS at pH 7.5, GFP was released into the higher-pH buffer (Fig. 3C). In contrast, when cells already incubated at pH 7.5 were washed a second time at the same pH, no further GFP could be observed by immunoblot analysis, suggesting that GFP does not associate with cells at pH 7.5 (Fig. 3C). The pH-dependent interaction of GFP with the cell surface was specific to biofilm cells since GFP was detectable in PBS both at pH 5 and at pH 7.5 after incubation with HG003 cells grown to the exponential phase (Fig. 3C). This indicates that a specific feature of cells grown under biofilm-forming conditions is required to mediate the binding of exogenously added proteins to the cell surface.

In the course of conducting these experiments, we observed that cell pellets from biofilm cells formed as diffuse smears (directly from biofilm medium or in PBS at pH 5) at low pH but formed as compact pellets after incubation at pH 7.5 (data not shown). Cell pellets from a biofilm were also more easily resuspended to homogeneity at pH 7.5 than at pH 5. Consequently, we examined cells by phase-contrast microscopy following resuspension. Biofilm cells resuspended at pH 5 were frequently found in large cell clusters that were often associated with other material (Fig. 3D, upper panels). In contrast, cells resuspended at pH 7.5 were largely separate or in small clusters without this associated material (Fig. 3D, lower panels). We propose that low pH causes cytoplasmic proteins to aggregate at the cell surface and that this aggregate is a principal component of the extracellular matrix that holds cells together in the biofilm.

To assess whether the accumulation of cytoplasmic proteins on the cell surface of HG003 biofilm cells is a common feature of S. aureus, we repeated our analysis using four other strains commonly used for the study of biofilm formation. Although they did not make biofilms as robust as those made by HG003, strains Newman, UAMS1, MN8, and RN4220 all produced quantifiable levels of biofilm when grown in 96-well plates in TSBG (see Fig. S3A in the supplemental material); furthermore, the medium pH decreased during growth to around 4.5 to 5 in all cases (data not shown). Cells were harvested from each biofilm and incubated
in PBS at pH 5 and pH 7.5 for 1 h at 4°C. As seen with HG003, it was possible to observe the release of proteins into PBS from three of the four strains at pH 7.5 but not at pH 5 (see Fig. S3B). Interestingly, RN4220 did not appear to release a significant amount of protein from the cell surface into PBS at pH 7.5 despite making a robust biofilm. This might suggest either that proteins adhere more tightly to RN4220 cells or that this strain utilizes a different mechanism for biofilm formation (7, 18). Taken together, these results indicate that accumulation of cytoplasmic proteins on the cell surface might be a frequent feature of biofilm formation in *S. aureus*.

**Cytoplasmic proteins associate with the outer side of cells in intact biofilms.** If cytoplasmic proteins accumulate on the cell surface under biofilm-inducing conditions and constitute the extracellular matrix, it should be possible to detect them in intact biofilms that have not been manually disrupted. We used immunofluorescence microscopy to visualize only those proteins accessible to antibodies and thus external to cells. To eliminate nonspecific background caused by the IgG-binding activity of protein A, we utilized a protein A deletion mutant of HG003 (Δspa) which is unaffected in its ability to form biofilms (see Fig. S1B in the supplemental material). As expected, cells of the Δspa mutant did not exhibit binding to a non-specific IgG antibody, anti-Halotag (see Fig. S4A).

Fixed biofilms were probed for target proteins using primary rabbit antibodies and visualized using a secondary antibody conjugated to Alexa Fluor 488. The location of individual cells within the biofilm was observed by staining the chromosomal DNA with DAPI (4’,6-diamidino-2-phenylindole). Anti-enolase binding was visualized as a diffuse accumulation of fluorescent signal around and between cells within the biofilm (Fig. 4A). The detected proteins appeared to occupy areas between, and not overlapping with, regions of DAPI staining, suggesting an extracellular location. In contrast, a cytoplasmic protein (represented by fluorescence from GFP expressed constitutively) was seen to colocalize with regions of DAPI staining (see Fig. S4B in the supplemental material). The localization of enolase was distinct from that of a typical covalently attached cell wall protein, protein A, which was seen as clear rings around the cell periphery of wild-type cells (see Fig. S4C). Anti-GAPDH antibodies localized around and between cells within the biofilm in a honeycomb-like pattern, again clearly occupying a space distinct from the nuclei/cytoplasm (Fig. 4A). The proteins detected by the two antibodies were most apparent in certain areas and predominantly closer to the top of the biofilm. Fluorescent signal was not apparent using nonspecific antibodies such as an anti-GFP antibody (with cells not producing GFP) or a rabbit anti-mouse-FITC antibody (data not shown).

Since the genes encoding enolase and GAPDH are essential in *S. aureus*, we were unable to construct deletion mutants that would function as negative controls in immunofluorescence experiments. To confirm that our results were not due to nonspecific binding of anti-enolase and anti-GAPDH antibodies to other cell surface proteins, we constructed Δspa strains in which the native copy of each gene (encoding enolase or GAPDH) was replaced with a copy that would result in a fusion of the HaloTag with the C terminus of the respective protein. The resulting strains (LCF88 and LCF89) were viable and exhibited no significant changes in growth or biofilm formation compared to the Δspa strain (see Fig. S5A in the supplemental material). Furthermore, LCF88 and LCF89 were found to produce bands of the expected size for the fusion products by immunoblotting using an anti-HaloTag antibody in cell lysates and in cell surface extracts (PBS at pH 7.5) of cells grown under biofilm conditions (see Fig. S5B). A commercial anti-HaloTag antibody did not react to proteins on the cell surface of the HG003 Δspa mutant, indicating that the antibody did not display nonspecific binding (Fig. 4B). In contrast, in strains LCF88 and LCF89, respectively, the anti-HaloTag antibody revealed a pattern of protein localization similar to that seen with anti-enolase and anti-GAPDH antibodies (Fig. 4B).

**DISCUSSION**

A hallmark of biofilm formation in certain bacteria is the production of a dedicated protein component of the extracellular matrix, such as Curli in *Escherichia coli* or TasA in *Bacillus subtilis* (19). We...
failed to detect an *S. aureus* protein that is unique to the biofilm matrix, although it is possible that such a dedicated protein(s) contributes to the matrix under other conditions or in other strains. Instead, our evidence suggests a model in which cytoplasmic proteins attach to and assemble on the cell surface to form the extracellular matrix. Formation of the matrix is triggered by a decrease in pH during the post-exponential phase in biofilm-inducing medium. We further propose that the matrix thus formed causes cells in the biofilm to adhere to each other. Interestingly, the assembly process is reversible in that proteins can be solubilized from the cell surface, after manual disruption of the biofilm, by artificially increasing the pH. Finally, our evidence indicates that pH-driven association of proteins on the cell surface is a common feature of protein-dependent biofilm formation in diverse strains of *S. aureus*.

How are cytoplasmic proteins released into the surrounding milieu during biofilm growth? It has previously been reported that a variety of species of Gram-positive bacteria, including *S. aureus*, release cytoplasmic proteins into the external environment during stationary phase (20, 21). However, the mechanism by which proteins lacking signal peptides are exported from cells is unclear, and both specific (e.g., secretion) and nonspecific (e.g., cell lysis) mechanisms are possible (20–23). Biofilm formation by *S. aureus* has previously been suggested to be dependent on regulated autolysis for the accumulation of extracellular DNA (24). Conceivably, regulated autolysis may also be the basis for the extracellular presence of cytoplasmic proteins.

The idea that cytoplasmic proteins might accumulate extracellularly and perform a novel function in that location is not new (22, 25). Such “moonlighting” functions of proteins appear to be widespread. In particular, enzymes of glycolysis are commonly present in the microenvironment of a wound or in-dwelling medical-device infection. However, many sites of inflammation in the context of this model for extracellular DNA (eDNA), which, along with protein, is required for biofilm formation. We suggest that eDNA might serve as an electrostatic net linking protein-coated cells in the biofilm. We note that at pH 4.5 to 5, these surface-associated, cytoplasmic proteins are likely to carry a net positive charge, facilitating their interaction with the proposed eDNA net. Finally, our findings are of course limited to an *in vitro* static biofilm model at a specific biofilm growth stage. It remains to be seen whether these findings will be applicable to other *in vitro* biofilm models (e.g., flow cells) and to physiologically relevant biofilms.

The repurposing of abundant cytoplasmic proteins as matrix components offers *S. aureus* the advantage of not investing in the production of dedicated matrix proteins during biofilm formation. Conceivably, other Gram-positive bacteria employ a similar moonlighting strategy in making biofilms, and the use of conserved cytoplasmic proteins in the biofilm matrix might facilitate interspecies interactions. *S. aureus* has been found in chronic wounds (such as diabetic ulcers) with multiple other bacterial species, and 60% of such wounds were observed to contain biofilms (4). *S. aureus* might form multispecies biofilms without the necessity to specifically recognize the dedicated matrix components of the other species. The recycling of cytoplasmic proteins could therefore represent an efficient and versatile strategy for building multicellular communities. It will be interesting to see in future work whether cytoplasmic proteins that moonlight as matrix components represent a widespread strategy for biofilm formation.

**Materials and Methods**

**Strains and growth conditions.** *S. aureus* strains are listed in Table S2 in the supplemental material. *S. aureus* was cultured in tryptic soy broth (TSB; EMD Millipore) and on LB agar at 37°C. Other growth media were brain heart infusion (BHI; EMD Millipore), BioExpress 1000 (Cambridge Isotope Laboratories), and TSB without glucose (peptone from casein [BD], 17 g/liter; peptone from soymeal [Amresco], 3 g/liter; NaCl [Sigma], 5 g/liter; dipotassium hydrogen phosphate [Maccon], 2.5 g/liter). *E. coli* DH5α was cultured in LB and on LB agar. Where appropriate, *S. aureus* was selected on 3 μg/ml tetracycline and *E. coli* on 50 μg/ml ampicillin. The pH of biofilm cultures was measured by spotting 20 to 30 μl on pH strips (range, 4.5 to 10; VWR).

**S. aureus quantitative biofilm assays.** For biofilm growth, an overnight culture of HG003 was diluted at 1 in 1,000 into fresh medium (typically TSB supplemented with 0.5% glucose [TSBG] or as specified in the text) and 200 μl was divided into aliquots and introduced into a Nunc MicroWell 96-well microplate (catalog. no. 167008; Thermo/Fisher Sci-
entific). The starting OD_{600} was recorded using a Bio-Tek Synergy II plate reader (BioTek Instruments). Plates were incubated statically at 37°C for 24 h. The medium in each well was removed to a new 96-well plate. The biofilms were washed twice with 100 μl phosphate-buffered saline (PBS) (pH 7.5), and the two washes were amalgamated into a new 96-well microtiter plate. The biofilms were resuspended in 200 μl PBS. The OD_{600} of each fraction was recorded using a Bio-Tek Synergy II plate reader. The starting OD_{600} was subtracted from that for each medium sample, and the absorbance of PBS was subtracted from the values for the wash and biofilm samples. Results from replicate wells (≥4) were averaged, and a standard deviation was calculated. Proteinkinase Α (Omega Bio-Tek) was used at 0.1 mg/ml. DNase I (Qiagen) was used at 28 U/ml.

Construction of unmarked deletion mutants in HG003. In-frame, unmarked deletions of specific regions of the HG003 genome were contructed essentially as described in reference 34. Deletion constructs were generated in pKFT using Gibson assembly (35) and primers shown in Tables S3 and S4. Constructs were transferred to HG003 and double-crossover recombinants selected as described in reference 34. For a detailed description, see the supplemental material.

Enrichment for cell-associated proteins from biofilm cells. HG003 was inoculated at 1 in 1,000 into TSBG and BioExpress 10015N in 6-well tissue culture plates (Falcon 353046) for each of three biological replicates. Plates were incubated statically at 37°C for 24 h. Cells and growth media were harvested using a well scraper at 4°C, and cells were pelleted by centrifugation at 7,500 × g for 5 min. Cells grown in TSBG for each replicate were resuspended in 5 ml PBS (pH 7.5 to 8) containing 1 mM phenylmethylsulfonyl fluoride (PMSF), and this was used to reuspend cells grown in BioExpress (equal numbers of OD_{600} units). Surface-associated proteins were enriched by biotinylation as described in reference 13. Proteins were separated by SDS-PAGE and stained with Coomassie brilliant blue R-250. A complete protein separation lane was cut into 5 equal gel slices. In-gel digestion was performed with trypsin (Promega) (36).

All peptides obtained from an in-gel digestion were separated by liquid chromatography and measured online by electrospray ionization (ESI) mass spectrometry. Liquid chromatography-tandem mass spectrometry (LC-MS/MS) analyses were performed using an Easy-nLC II high-pressure liquid chromatography (HPLC) system (Thermo/Fisher Scientific) coupled directly to an LTQ Orbitrap Velos mass spectrometer (Thermo/Fisher Scientific). All MS analysis and protein identification and quantification procedures were performed at the Harvard Mass Spectrometry and Proteomics Resource Laboratory, FAS Center for Systems Biology, Cambridge, MA.

Analysis of proteins released into PBS. Biofilm cells grown in 6-well plates were pelleted by centrifugation at 5,500 × g for 10 min. Proteins were precipitated from growth medium using trichloroacetic acid (TCA) precipitation. Biofilm cells were gently resuspended in 5 ml PBS (pH 7.5 to 8 or pH 5 [pH decreased by the addition of HCl]) containing cOmplete Protease Inhibitor Cocktail (Roche) (according to manufacturer’s instructions) by gently swirling the 50-ml tube. Cells were incubated at 4°C with gentle rotation for 1 h. Cells were pelleted by centrifugation at 5,500 × g for 10 min, and the supernatant was passed over a 0.2-μm-pore-size filter. Proteins were concentrated for gel electrophoresis by TCA precipitation. Whole-cell lysates were produced by FastPrep disruption (MP Biomedicals) (speed, 6.5; three 40-s steps) followed by centrifugation at 18,000 × g and 4°C for 15 min to pellet cell debris. Proteins were routinely separated on 12.5% SDS-PAGE gels and stained with GelCode Blue Stain reagent (Thermo/Fisher Scientific).

For proteinkinase K experiments, 0.1 mg/ml proteinkinase K was added at the time of inoculation to appropriate wells. Protease activity was inhibited by adding ProteaseInhibit solution (G-Biosciences) to biofilm wells at a 3× final concentration and incubating for 1 h at 37°C. Where applicable, whole-cell lysate from HG003 or LC8F1 grown to the stationary phase in TSB (passed over a 0.2-μm-pore-size filter to remove intact cells) (approximately 25 μg total) was added either directly to biofilm cultures after removal of cells from the biofilm growth plate or in the PBS extraction buffer, as indicated.

Western blot analysis. Proteins separated by SDS-PAGE were transferred to a polyvinylidene difluoride (PVDF) membrane. The membrane was blocked for 2 h at room temperature in 5% milk powder–5% goat serum (Sigma-Aldrich)–Tris-buffered saline–Tween 20 (TBST; 150 mM NaCl, 50 mM Tris-HCl, 0.05% Tween 20). Membranes were probed with primary antibodies diluted in 5% milk–TBST. Rabbit anti-endoase antibody (against B. subtilis enolase; kind gift of Jørg Stülke) was used at a dilution of 1:5,000, rabbit anti-GAPDH (37) was used at a dilution of 1:10,000, rabbit anti-GFP (38) was used at a dilution of 1:4,000, and rabbit anti-HaloTag polyclonal Ab (pAb) (Promega) was used at a dilution of 1:5,000. Antibody binding was detected with a goat anti-rabbit antibody conjugated to horseradish peroxidase (Bio-Rad Laboratories) and SuperSignal West Dura chemiluminescent substrate (Thermo/Fisher Scientific).

LIVE/DEAD staining. A 1-μl volume of propidium iodide (diluted 1/10 in PBS) and a 1-μl volume of SYTO 9 (1/10 in PBS) (LIVE/DEAD BaCl2) were added per ml of cells. Cells were incubated with the dyes at room temperature for 15 min. Cells were washed once in PBS, resuspended in 150 μl PBS, and observed on a phase-contrast microscope fitted with a 63× objective. Images were taken using phase (500-ms exposure), GFP (100-ms exposure), and mCherry (100-ms exposure) filters. Live (green) and dead (red) cells were counted manually, and data were averaged from a minimum of 1,000 cells across at least three fields of view in three replicate experiments.

Autolysis assay. Triton X-100 induced autolysis assays were performed essentially as described in reference 39. Briefly, cells were grown in TSBG under biofilm-inducing conditions for 24 h or under shaking conditions to the exponential phase (2 h). Cells were harvested by centrifugation as described and then resuspended in PBS to an OD600 of 2. An equal volume of PBS or PBS containing 0.1% Triton X-100 was then added, and 100 μl of cells was introduced into eight replicate wells in a 96-well microtiter plate. The OD_{600} was monitored every 10 min for 1.5 h and every 30 min thereafter. The OD_{600} at each time point was plotted as a percentage of the starting OD_{600}.

Immunofluorescence. To carry out immunostaining of S. aureus biofilms, we adapted a method for visualizing cell surface proteins on planktonic cells (10) (for details, see the supplemental material). To detect antibody binding, samples were exposed to goat anti-rabbit IgG conjugated to Alexa Fluor 488 dye (Life Technologies) diluted 1:1,000 in PBS and chromosomal DNA was stained with DAPI (7.5 μg/ml final concentration). Biofilms were visualized by confocal laser scanning microscopy on a Zeiss LSM 700 microscope fitted with a 63× oil objective. Alexa Fluor 488 was excited with a 488-nm laser and DAPI with a 405-nm laser. Images were processed and analyzed using ZEN lite 2012 (Blue edition; Zeiss).

Construction of gene-HaloTag fusions at native loci. Replacement of the eno and gnp genes, at their native loci in the genome, with copies of the respective ORFs fused to a sequence encoding the HaloTag (Promega) was carried out as described in the supplemental material. Subsequently, to allow the use of these strains for immunostaining, the spa gene was deleted by homologous recombination using construct pLF048.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01667-14/-/DCSupplemental.
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