Abstract

Staphylococcus aureus is an opportunistic pathogen that colonizes the skin and mucosal surfaces of mammals. Persistent staphylococcal infections often involve surface-associated communities called biofilms. Here we report the discovery of a novel extracellular fibril structure that promotes S. aureus biofilm integrity. Biochemical and genetic analysis has revealed that these fibers have amyloid-like properties and consist of small peptides called phenol soluble modulins (PSMs). Mutants unable to produce PSMs were susceptible to biofilm disassembly by matrix degrading enzymes and mechanical stress. Previous work has associated PSMs with biofilm disassembly, and we present data showing that soluble PSM peptides disperse biofilms while polymerized peptides do not. This work suggests the PSMs’ aggregation into amyloid fibers modulates their biological activity and role in biofilms.

Introduction

Staphylococcus aureus is the causative agent of numerous diseases ranging from relatively benign skin conditions to fatal systemic infections. Formation of bacterial biofilms on host tissues and implanted materials contributes to chronic S. aureus infections, as biofilms are exceptionally resistant to host immune response and chemotherapies [1]. Biofilms are multicellular structures encased in a matrix of proteins, polysaccharides, extracellular DNA, and other environmental factors [1,2]. Biomolecules that digest matrix components (e.g., proteases, DNases, and glycoside hydrolases) can disrupt established biofilms and render detached cells susceptible to antimicrobials [3,4,5,6,7].

The precise composition of the biofilm matrix varies greatly by strain, physiological state, and nutrient availability [5,8,9,10,11,12]. In this study, we examined how growth media affects the composition of the biofilm matrix. This led to the discovery of an extracellular fibril structure in S. aureus biofilms grown in a non-standard rich media. These fibers share morphological and biophysical characteristics with functional bacterial amyloids such as curli in Escherichia coli biofilms, TasA of Bacillus subtilis, and the Fap fimbriae in Pseudomonas aeruginosa [13,14,15,16]. Biochemical and genetic analysis revealed that these fibril structures are composed of small peptides called phenol soluble modulins (PSMs). Mutants incapable of producing PSMs formed biofilms that were susceptible to disassembly by enzymatic degradation and mechanical stress.

Previous work has demonstrated that PSMs are surfactant-like peptides that promote biofilm disassembly [17,18,19,20,21]; exhibit antimicrobial activity against niche bacteria [22,23,24]; hinder host immune response by recruiting and lysing neutrophils; and are abundant virulence factors produced by community-associated MRSA strains (CA-MRSA) [18,25,26,27]. The genes encoding the core family of PSM peptides are highly conserved across S. aureus strains: four are expressed from the alpha (psm1–4) operon, two are expressed from the beta (bpsm1&2) operon, and the delta hemolysin (hld) is encoded within the regulatory RNA, RNAIII [28,29,30]. The significance of the PSMs has only recently been investigated because the coding sequences of the xpsm & bpsm peptides are small enough to have eluded detection by conventional gene annotation programs, and they are still poorly annotated in public databases [29,30].

We have found that ordered aggregation of PSM peptides into amyloid-like fibers can abrogate the biofilm disassembly activity ascribed to monomeric PSM peptides [12,17,18,19,20,21]. Our findings suggest that PSMs can modulate biofilm disassembly using amyloid-like aggregation as a control point for their activity. This is the first report to identify and characterize extracellular fibrillae in the S. aureus biofilm, and our research could lead to new approaches in treating persistent biofilm associated infections.

Results

Biofilms grown in PNG media resist biofilm disassembly

Biofilms that persist in the human body are often resistant to conventional antimicrobial treatment prior to dispersal. To gain insight into how the S. aureus biofilm matrix affects disassembly under different growth conditions, we grew S. aureus flow cell biofilms with various lab media. Next we used enzymes known to target primary matrix components in order to test biofilm resistance (Fig. 1A & 1B). These enzymes include protease K (protein), DNaseI (DNA), and dispersin B (polysaccharide). By using a variety of degradative enzymes, we expected to observe complete biofilm eradication.
Biofilms grown in tryptic soy broth supplemented with glucose (TSBg) rapidly disassembled after enzymatic treatment (Fig. 1A).

However, biofilms grown in peptone-NaCl-glucose (PNG) media did not disassemble after the same enzymatic treatment (Fig. 1B).

We also assessed the ability of biofilms attached at the air-liquid interface of glass culture tubes to withstand exposure to an anionic surfactant, sodium dodecyl sulfate (SDS). Again, biofilms grown in TSBg were more sensitive to surfactant-mediated disassembly than those grown in PNG (Fig. 1C). We interpreted these results to be an indication that growth in PNG alters the matrix composition, increasing the biofilm’s resistance to enzymatic degradation and surfactant dispersal. We hypothesized that a new, previously unaccounted for matrix component was influencing S. aureus surfactant dispersal. We found that these structures are composed of known virulence factors, which indicates a common pathway between infectious and static lifestyles in the body.

**Biofilms resistant to dispersal contain extracellular fibers**

To investigate how biofilms grown in PNG media are able to resist disassembly, we grew biofilms in drip bioreactors under sensitive (TSBg) or resistant (PNG) conditions for five days. Biofilms were harvested and disrupted by vortexing and sonication. Transmission electron microscope (TEM) imaging of cells revealed the presence of extracellular fibers in enzyme-resistant biofilms (Fig. 2B), but not in enzyme-sensitive biofilms (Fig. 2A). The fibers had a central diameter of ~12 nm and were closely associated with bacterial cells (Fig. 2C). S. aureus has never before been shown to produce large, extracellular structures. Additionally, we observed identical fibers associated with biofilm cells in several lab strains (LAC, UAMS, MN8) and six clinical isolates (three nasal isolates and three blood isolates) by TEM, demonstrating that fiber formation is not specific to strain SH1000. Of note, we found that an agr quorum sensing mutant (SH1001) was unable to produce fibers (Fig. 2D).

**Staphylococcus aureus fibers are composed of phenol soluble modulins (PSMs)**

The novel extracellular fibers isolated from robust biofilm matrices share morphological similarities with the bacterial functional amyloids curli in *E. coli* and TasA in *B. subtilis* [15,16]. Amyloid proteins form highly stable polymerized aggregates that exhibit well-defined biochemical and biophysical characteristics [14,31,32]. We hypothesized that our fibers were also functional bacterial amyloids. To identify the protein composing these fibers, we used two approaches to take advantage of the biophysical characteristics of functional bacterial amyloids.

Amyloid fibrils from bacterial biofilms were previously shown to be poorly soluble in sodium dodecyl sulfate (SDS) and do not migrate through polyacrylamide gels [33]. We therefore employed SDS-PAGE to isolate large insoluble structures. *S. aureus* biofilm
samples were grown in drip bioreactors for five days with PNG media or TSBg. These biofilms were harvested, homogenized, and lysed, and the lysates were run into a 12% SDS-polyacrylamide gel. Lysates from biofilms grown in PNG media retained insoluble material in the wells of the stacking gel while TSBg-grown lysates did not (Fig. 3A). The insoluble material retained within the wells of the stacking gel was recovered, treated with 100% formic acid (FA), then separated once more by SDS-PAGE alongside an untreated control (Fig. 3B). We observed protein enrichment in the FA-treated sample, and the four dominant bands were excised and analyzed via mass spectrometry (MS). Surprisingly, MS analysis identified the same peptides as being abundant in each sample, regardless of the band’s migration through the gel matrix (Fig. 3B & 3D). These proteins were the alpha (αPSM) phenol soluble modulins and the *S. aureus* delta hemolysin (δ-toxin) (Fig. 4A & 4B).

An additional approach to identify the fiber subunit was to isolate fibers from biofilm cells using a tissue homogenizer (Fig. 3C), incubate fiber isolates for 48 hours at pH 2 with pepsin, and analyze the sample with MS. Again, we detected the same αPSM peptides present in the SDS-PAGE isolation plus two beta PSMs (βPSM) (Fig. 3D). αPSM3 was not identified in either preparation, but it should be noted that αPSM3’s sequence contains several trypsin cleavage sites, so it is likely that it would not be detected after extensive digestion. The same fiber isolation procedure revealed no visible fibers by TEM when biofilms were grown in TSBg.

We generated an ΔαβPSM double-knockout mutant and assessed fiber production. TEM analysis of biofilm cells revealed that this mutant did not produce fibers after five days of growth in PNG media (Fig. 5B) compared to the wildtype parent strain grown under the same conditions (Fig. 5A). In addition, fibers could be isolated from wildtype (Fig. 5D) but not mutant biofilms (Fig. 5E). Fiber production was complemented by expression of the *αpsm* and *βpsm* operons in trans (Fig. 5C & 5F).

PSM peptides form fibers similar to bacterial functional amyloids

We assayed synthetic PSMs peptides for their capacity to form fibers in vitro. To minimize the prevalence of polymeric “seed” complexes, all synthetic peptides were treated with HFIP/TFA and dried in vacuo prior to assay [34]. Incubation of the seven previously identified PSM peptides (αPSM1–4, βPSM1–2, and δ-toxin) demonstrated their capacity to self-assemble into fibers (Fig. 6B). We used the amyloid-specific dye thioflavin T (ThT) to observe amyloid formation over time [34, 35]. When we assayed the PSMs for polymerization in the presence of ThT at room temperature, we observed a robust increase in normalized
fluorescence (Fig. 6A). Greater peptide concentration increased ThT fluorescence and showed rapid binding kinetics similar to an amyloid-nucleator system (Fig. 6A) [35,36,37]. PSM fibers exposed to ThT exhibited an emission spike near 490 nm that is also observed in other amyloid fibrils (Fig. 6C) [36,38,39]. Incubation of PSM fibers with the dye Congo red (CR) resulted in a characteristic absorbance “red shift”, indicative of cross β structure conserved in all amyloid fibers (Fig. 6D) [40]. Furthermore, PSM fibers isolated from solution through centrifugation displayed increased β-sheet content (Fig. 6E), which is consistent with data published from other bacterial amyloids [13,16]. These in vitro observations complement our genetic and physiological data, further supporting the notion that PSMs can form amyloid fibrils.

Mutants unable to synthesize PSMs produce biofilms susceptible to matrix-degrading enzymes and mechanical stress

Because biofilms grown in PNG media resist disassembly by matrix-degrading enzymes and surfactants (Fig. 1), we challenged ΔαPSM mutant biofilms grown under the same conditions. In contrast to its isogenic parent strain, an ΔαPSM mutant biofilm readily disassembled after exposure to proteinase K, DNaseI, and dispersin B (Fig. 7A). Complementation of the ΔαPSM mutant in trans restored the resistant biofilm phenotype (Fig. 7B). We also examined the effects of mechanical stress (vortexing) on biofilms attached at the air-liquid interface of glass culture tubes. An ΔαPSM mutant biofilm readily disassembled with exposure to mechanical stress, while biofilms of the isogenic parent and complemented strains both remained intact (Fig. 7C). Taken together these data do suggest that PSM fibers enhance biofilm integrity.

Fibrillation modulates PSM activity

Previous work has demonstrated that soluble PSMs assist biofilm disassembly [19,21,27]. Based on our findings that PSM fibers improve biofilm integrity (Fig. 1 & Fig. 7), we hypothesized that sequestration of PSMs into extracellular fibers could alter their activity. Synthetic αPSM1 peptides readily formed fibers that bind CR and ThT after 24 hours of incubation in solution (Fig. 8C, 8D, 8E). To test whether or not fibrillation alters peptide activity, we exposed 24-hour S. aureus biofilms to either freshly solublized αPSM1 peptides (Fig. 8B) or αPSM1 that had been allowed to polymerize overnight (Fig. 8C). Exposure to soluble αPSM1 significantly reduced the amount of adherent biofilm; however, exposure to αPSM1 fibers had no discernable effect on the biofilms (Fig. 8A). This finding suggests that the aggregation of PSMs into amyloid-like fibers can modulate their ability to disassemble biofilms.

Discussion

Biofilm formation and disassembly are carefully coordinated with the production and breakdown of matrix components. The biofilm lifecycle of attachment, maturation, and disassembly participates as a mechanism of virulence in many persistent S. aureus infections [1,2,4,5,11,12,19]. A better understanding of the dynamic S. aureus matrix environment may inspire new, innovative techniques for controlling biofilm infections.

Previous studies have shown that the S. aureus biofilm matrix contains polysaccharides and DNA that interact with structural and enzymatic proteins [3,4,5,7,41]. In this work we demonstrate that under certain growth conditions, S. aureus produces amyloid-like fibers that contribute to biofilm integrity (Fig. 1, 2, 7).
Purification and analysis of fibers revealed that several small peptides of the phenol soluble modulin family were present (Fig. 3 & Fig. 4). Fibers were not detected in biofilms of a ΔABpsm deletion mutant under the same conditions that favor their production in wildtype strains (Fig. 5). ΔABpsm mutant biofilms were further demonstrated to be significantly more susceptible to disassembly with matrix degrading enzymes and mechanical stress than their isogenic parent (Fig. 7). To the best of our knowledge, this is the first report describing an extracellular fibril structure in *S. aureus* biofilms. We refer to Figure 4. Phenol soluble modulins are small peptides expressed from three discrete regions of the *S. aureus* genome. (A) Phenol soluble modulins (PSMs) are encoded in two operons, the alpha (αPSM1–4) and beta (βPSM1–2) operons, and δ-toxin is encoded within the Agr regulatory RNA, RNAIII (hld). (B) PSMs are small hydrophobic peptides with highly similar amino acid content.

doi:10.1371/journal.ppat.1002744.g004

Figure 5. Mutants unable to produce α and βPSMs fail to form fibers during biofilm growth. TEM micrographs of *S. aureus* biofilm cells grown for five days in PNG media. (A) wildtype (strain SH1000), (B) Δαβpsm (strain BB2388), (C) Δαβpsm complemented (strain BB2408). (D–F) TEM micrographs of fiber preparations from wildtype (D), Δαβpsm (E), and Δαβpsm complemented (F). Bars indicate 500 nm.

doi:10.1371/journal.ppat.1002744.g005
these matrix components as being amyloid-like because they possess some of the characteristics often attributed to amyloid proteins: fibril morphology (Fig. 2, 5, 6, 8), relative SDS insolubility (Fig. 1 & Fig. 3), binding to the amyloid-specific dyes thioflavin T and Congo red (Fig. 6 & Fig. 8), and they display β-sheet structure [14,31]. The observation that PSM peptides not only self-assemble, but contribute to the biofilm’s structural integrity is intriguing in light of recent work describing the PSMs’ involvement in biofilm disassembly [11,20,27].

It is well-documented that the PSMs are regulated by the agr quorum-sensing network [11,12,28,42], and we similarly have found that an agr deficient strain did not produce fibers (Fig. 2D). This contributes to a growing body of evidence which implicates the agr system to have wide-ranging effects beyond heightened pathogenicity and biofilm dispersal [4,20,21,43]. It is tempting to speculate that the media-dependent fiber production is somehow influencing agr regulation, perhaps through metabolism or through other signaling cues.

PSMs were first isolated from *Staphylococcus epidermidis* cultures as a polypeptide complex, and have since been shown to interact biochemically [22,29]. We have demonstrated that synthetic *S. aureus* PSM peptides are capable of self-assembling into amyloid-like fibers in vitro (Fig. 6A & Fig. 6B). These fibers demonstrate CR and ThT binding capacities similar to known amyloid proteins. PSMs, including δ-toxin have been previously characterized as amphipathic α-helices [22,30,44,45]. Our data indicate that soluble PSMs have a helical structure in solution, but transition to adopt a more β-rich structure after aggregation (Fig. 6E). The assembly of zPSM1 into fibers prevents the biofilm disassembly activity attributed to soluble peptides species (Fig. 7). We interpret these results as evidence that aggregation into amyloid fibers can regulate PSM activity in the microenvironment of the biofilm.

Our findings demonstrate that *S. aureus* PSMs can be found in biofilms as fibrils, and may implicate fibril formation as a means of altering their activity and function. It is not known at this time what mechanisms influence the PSMs’ ability to switch from a monomeric to fibril state, nor is it clear how this affects the formation and disassembly of biofilms. It is possible that PSM fibrillation is synchronized in *vivo* by a nucleator protein, similar to CsgB in *E. coli* [34]. Formylation may also play a role; the PSMs
and δ-toxin are detected at the protein level both with and without with a formylated methionine modification [29,30,46,47], and the PSMs identified in our MS analysis contained primarily deformylated N-methionines. Recent work demonstrates that non-N-formylated PSMs are strong activators of FPR2 receptors, which also respond to amyloidogenic peptides like Aβ1-42 and serum amyloid A [48,49,50], and may implicate a role for deformylation in fibril construction. We speculate that there are numerous other environmental cues (such as pH and osmolarity) driving the PSMs’ commitment to the fibrillation pathway, and this is currently under investigation.

This study builds upon an emerging paradigm emphasizing that amyloid fibers are common in the biofilm matrices of many bacterial species. Curli fibers produced by pathogenic E. coli and other enterics were the first functional amyloids to be characterized [15,32,33]. The gram-positive bacteria Streptomyces coelicolor produces several small peptide species, which have been shown to polymerize in vitro and in vivo to facilitate sporulation at the air-liquid interface [51,52,54,58]. Recent work in B. subtilis has shown that the antimicrobial and spore coat protein TasA can assemble into amyloid-like fibrils during biofilm growth [16]. Even natural biofilms collected from a variety of environmental niches appear to contain amyloid-like fibers [55], indicating that the production of bacterial amyloids may be a shared feature of biofilm matrices from many different bacterial communities.

We propose that amyloid-like aggregation of toxic proteins is an under appreciated form of posttranslational regulation utilized throughout nature, and even more examples continue to emerge. The antimicrobial activity of the Klebsiella pneumoniae bacteriocin microcin E492 can be turned off through their assembly into amyloid-like fibers [56]. Recent work by Maji et al. has demonstrated that even human peptide hormones can form amyloid-like structures for storage [57]. Likewise, PSMs may be stored as inert fibrils in a sessile biofilm until conditions arise that favor their dissociation to promote biofilm disassembly, antimicrobial activity, or virulence. This work presents evidence that S. aureus PSMs can be found in biofilms as large fibril structures providing new insight into how quorum sensing and virulence play into the complexity of the biofilm lifecycle.

Materials and Methods

Bacterial strains and growth conditions

The bacterial strains and plasmids used in this study are listed in Table 1. All DNA manipulations were performed in Escherichia coli strain DH5α. Oligonucleotides were synthesized by Integrated DNA Technologies (Coralville, IA). Plasmids were transformed into Staphylococcus aureus RN4220 by electroporation, purified, and moved to indicated S. aureus strains by electroporation. Deletion mutants were generated via allelic replacement using the vector pKFC as described previously [58]. To create the αpsm mutation, a region upstream of αpsm was amplified from SH1000 genomic DNA using primers alphaPSM221 (CGC GAG CTC GTT GAG GCA CGC GCC ACT CGC CAG) and alphaPSM162 (GCT AGC GGT ACC ACG CGT GAT GCC AGC GAT ACC CAT TAA) and a downstream region was amplified using alphaPSM163 (ACG CGT GGT ACC CGT GAT GAT ACC ACG CTA TCG CCG CAC G) and alphaPSM164 (TAT CTC GGG GAT GGT GGG GGA CTA TCG CGC ACA G). The resulting PCR products were gel purified, digested with KpnI, ligated with T4 DNA ligase and the ligation was used as a template in a subsequent PCR reaction with the primers alphaPSM221 and alphaPSM164. The resulting PCR product was gel purified and digested with Sall and Xmal and ligated with pKFC plasmid digested with the same enzymes to create pKFC-αpsm. The resulting plasmid construct was used to create an allelic Δαpsm deletion in the SH1000 background following the protocol outline by Kato et al [58].

To create the βpsm mutation, a region upstream of βpsm was amplified from SH1000 genomic DNA using primers BetaUpF

![Image](58x24 to 76x41)

**Figure 7. An αpsm mutant forms biofilms susceptible to disassembly by matrix degrading enzymes and mechanical stress.** Confocal micrographs of Δαpsm mutant (A) (strain BB2388) versus complemented mutant expressing α and βpsm operons in trans (B) (strain BB2408) flow cell biofilms grown for 30 hours prior to proteinase K, dispersin B, and DNase exposure (at 0.2 μg/mL each). Images are representative of three separate experiments and each side of a grid square represents 20 μm. (C) Analysis of biofilm development at the air-liquid interface of test tube cultures in PNG media after vortexing. Graph shows quantification of biofilm biomass (OD A595) and images below show stained test tube biofilms. * P<0.005 compared to wildtype.

doi:10.1371/journal.ppat.1002744.g007
(CCC GGA TCC GGT GTA GTG TAG TTC AGG) and BetaUpR (ACG CGT GGT ACC GCT AGC GCG TTA AAT AAA CCT TCC ATT G) and a downstream region was amplified using primers BetaDownF (5’ GCT AGC GGT ACC ACG CGT GGC ACA AGT ATC GTA GAC ATC G) and BetaDownR (5’ GCG GTC GAC GGC GTC TGA TTT AAC CTT CTC). The resulting PCR products were gel purified and used as a template in a subsequent PCR reaction with the BetaUpF and BetaDownR primers. The resulting PCR product was gel purified and digested with BamHI and SalI and ligated with pKFC plasmid digested with the same enzymes to create pKFC-bpsm. The resulting plasmid was used to create an allelic Dbpsm deletion in the SH1000 DapscM background following the protocol outline by Kato et al. to create the double knockout Abpsm mutant [58].

Complementation vectors were created as follows: the bpsm locus with its native promoter was amplified from S. aureus SH1000 genomic DNA using primers GAC GAA TTC AGG CAA CTT AAT TGT G and GAC AAG CTT GCT TCC CAA TGT TGG TG. The resulting PCR product was digested with HindIII and EcoRI and ligated with pAH8 [4], that had been digested with the same enzymes to create pRSbpsm. The apscM locus was amplified from SH1000 genomic DNA using primers ACT GAG GTA CCA GAA TTC AAT AA and ACT AGG AGC TCC AAA GGA GGA AAT CTT AAT GGG T. The resulting PCR product was digested with KpnI and SacI and ligated with pALC2073 [59], then digested with the same enzymes to create pALC2073aPSM.

Biofilm experiments
Flow cell and drip biofilms were grown as previously described [5,60]. Biofilm growth medium was either 0.6 g/L tryptic soy broth and 1.5 g/L glucose (TSBg) or 3.3 g/L peptone, 2.6 g/L NaCl, 3.3 g/L glucose (PNG).

For biofilm disassembly experiments performed in flow cells, enzymes proteinase K, DNaseI and dispersin B were suspended in water and added to the media reservoir at a final concentration of 0.2 μg/mL. Confocal scanning laser microscopy and image analysis was performed as described previously [5]. Strains contained pAH9 expressing mCherry or were stained with propidium iodide as previously described [5].

Test tube biofilms forming at the air-liquid interface of glass culture tubes were grown in 3 mL of TSBg or PNG for 2 days at 37°C shaking at 200 rpm. Liquid media was removed and exchanged with either 10 mL sterile ddH2O containing 1% SDS or sterile ddH2O alone. Tubes were vortexed for 5 seconds and all liquid was removed. The remaining biofilm biomass was visualized by staining with 0.1% crystal violet and quantified by solubilizing the stain in acidified ethanol and measuring the optical density at A595.

Transmission electron microscopy (TEM) was performed using a Philips CM12 transmission electron microscope. Samples prepared for TEM imaging were spotted onto formvar-coated copper grids, incubated for 5 minutes, washed with sterile ddH2O, and negatively stained with 2% uranyl acetate for 60 seconds.

Isolation of fibers from biofilm cultures
Drip bioreactor biofilms were grown as previously described [60]. Fibers were collected after 5 days growth by scraping biofilms into 3 mL of potassium phosphate buffer (50 mM, pH 7). The biofilm suspensions were homogenized using a tissue homogenizer (TissueMiser, Fisher) to shear fibers free from the cell walls. Supernatants were clarified by repeated centrifugation at 13,000 RPM for 2 minutes to remove cells. The cell-free
supernatant was incubated in 200 mM NaCl and the fibers were isolated using Millipore Amicon Ultra Centrifugal Filter Units with a pore size of 100 kDa. Presence of fibers was confirmed via TEM imaging.

Identification of aggregative peptides

Fibril subunits were identified by harvesting drip biofilms after 5 days of growth in PNG and suspending them in 15 mL 10 mM Tris HCl, pH 8.0 (Tris buffer), supplemented with 0.1 mg of RNase A (bovine pancreas; Sigma Chemical Co., St. Louis, Mo.) and 0.1 mg of DNaseI (bovine pancreas; Boehringer, Mannheim, Germany) per mL. Cells were lysed by repeated sonication and the addition of lysostaphin (1 mg/mL, Sigma) and 1 mM MgCl₂ prior to incubation at 37°C for 20 min. Lysozyme (Sigma) was added to 1 mg/mL, and the samples were incubated with shaking for 40 min at 37°C, after which they were adjusted to 1% sodium dodecyl sulfate (SDS) and incubated further (30 min, 37°C). The remaining insoluble material was collected by centrifugation (12,100xg, 15 min, 25°C), washed and suspended in 10 mL 10 mM Tris buffer. The pellet was digested again with RNase, DNaseI, and lysozyme as described above, washed twice with Tris buffer, and suspended in 2 mL SDS-polyacrylamide gel electrophoresis (PAGE) sample buffer (10% glycerol, 5%-mercaptoethanol, 1% SDS, 62.5 mM Tris HCl [pH 6.8]). The sample was boiled for 5 min, loaded onto a 12% polyacrylamide gel (3% stacking gel), and subjected to electrophoresis at 20 mA for 5 h. The material retained in the stacking gel was excised, washed three times in ddH₂O, extracted twice with 95% ethanol, and dried in a speedvac. The desiccated sample was resuspended in ddH₂O and sonicated to break up any clumps. Half of this material was incubated with formic acid (90%) for 20 min then dried in a speedvac. Both the formic acid treated and untreated samples were resuspended in SDS-PAGE sample buffer and run into a fresh 12% PAGE gel. Bands that appeared in the formic acid treated sample were excised and analyzed via LC-MS/MS (MS Bioworks, Ann Arbor, MI).

Fiber protein components were also identified by incubating fiber isolates in pepsin for 24 h before subsequent LC-MS/MS analysis (MS Bioworks, Ann Arbor, MI). The value for the abundance measurement is the Normalized Spectral Abundance Factor (NSAF).

PSM polymerization experiments

Non-formylated PSM peptides were synthesized by Peptide 2.0 and assayed to be >90% pure by HPLC. Synthetic peptides were prepared and assayed as previously described [15,34] to eliminate
large aggregates from lyophilization prior to assay. Each dry peptide stock was dissolved to a concentration of 0.5 mg/mL in a 1:1 mixture of trifluoroacetic acid (TFA) and hexafluoroisopropanol (HFIP). Peptides were then sonicated for 10 minutes and incubated at room temperature for 1 h. Solvent TFA/HFIP was removed by speedvac at room temperature. Dried peptide stocks were stored at −80°C. All assays were performed with equal stoichiometric ratios of 0.1 mg/mL peptide unless otherwise noted.

All polymerization assays were performed in 96-well black opaque, polystyrene, TC-treated plates (Corning). Prior to assay, treated peptides were thawed and dissolved in dimethyl sulfoxide (DMSO) to a concentration of 10 mg/mL immediately prior to assay. Freshly dissolved peptides were diluted into sterile ddH₂O containing 0.2 mM thioflavin T (ThT) and assayed at room temperature. Fluorescence was measured every 10 minutes after shaking by a Tecan Infinite M200 plate reader at 438 nm excitation and 495 nm emission. ThT fluorescence during polymerization was corrected by subtracting the background intensity of an identical sample without ThT.

Additionally, ThT fluorescence and Congo red (CR) absorbance scans were performed on polymerized peptides that were allowed to polymerize for 48 h in ddH₂O. Samples were incubated in 0.2 mM ThT or 0.001% (w/v) CR in ddH₂O for 15 minutes prior to assay on the Tecan plate reader. CR and ThT absorbance scans were corrected by subtracting the background intensity of an identical sample without dye.

Circular dichroism spectroscopy
Treated peptide stocks were thawed and dissolved in hexafluoroisopropanol (HFIP) to a concentration of 10 mg/mL immediately prior to assay. Triplicate samples consisting of 0.1 mg/mL of each freshly dissolved peptide diluted together in 500 μL sterile ddH₂O were incubated with shaking at room temperature for 48 h. Samples were then pelleted at 15,000 RPM for 30 minutes to isolate any aggregated species. The supernatant was carefully removed from the pellet by aspiration and transferred to a clean, sterile eppendorf tube. The remaining pellet was resuspended in 200 μL ddH₂O by bath sonication for 10 minutes. The supernatant and pellets of each sample were assayed separately. Far UV circular dichroism (CD) measurements were performed with a Jasco-J715 spectropolarimeter using quartz cells with 0.1 cm path length. CD spectra between 190 and 250 nm were recorded in millidegrees and converted to molar ellipticity using an average MRW of 113 for PSM1–4, βPSM1&2, and δ-toxin. The average of five scans was recorded at 25°C using a 2 nm bandwidth with a 20 nm min⁻¹ scanning speed. All triplicate samples showed similar ellipticity patterns.

Biofilm dispersal assay
Synthetic PSM peptides were allowed to polymerize overnight, and fibril formation was verified by TEM imaging. Equivalent concentrations of either polymerized or freshly diluted peptides were added to 24 hours SH1000 biofilms grown in 66% TSB+0.2% glucose and incubated at 37°C for 6 hours. Biofilms were washed to remove non-adherent cells then stained with 0.1% crystal violet, dried, and solubilized with acidified ethanol and spectrophotometrically quantitated at A595.

Statistics were performed using a 1-way analysis of variance (ANOVA). Results are expressed as mean ± standard deviation.

Acknowledgments
We would like to extend our gratitude to members of the Chapman laboratory for their helpful discussions and suggestions. Additionally, we acknowledge the contributions of Gregg Sobocinski, Dr. Ursula Jakob and the Jakob lab, Dr. Ari Gafni and Dr. Joseph Schauerte to this work for their technical assistance.

Author Contributions
Conceived and designed the experiments: KS AKS RES BRB. Performed the experiments: KS AKS RES AHR BRB. Analyzed the data: KS AKS RES BRB. Contributed reagents/materials/analysis tools: AH R. Wrote the paper: KS BRB.
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