Biofilms are communities of microbes embedded in a microbially extracellular matrix. Their formation is considered the main virulence mechanism enabling the opportunistic bacterial pathogen Staphylococcus epidermidis to cause devastating nosocomial, implant-associated infections. Biofilms often contain proteins, and an 18-kDa protein called small basic protein (Sbp) recently was discovered in the S. epidermidis biofilm matrix and may serve as a scaffolding protein in both polysaccharide intercellular adhesin (PIA)-dependent and accumulation-associated protein (Aap)-dependent biofilm formations. In Aap-mediated biofilm formation, Sbp colocalizes with Domain-B of Aap, implying that Sbp directly interacts with Aap’s Domain-B. However, the structure of Sbp and its interaction with Aap, as well as the molecular mechanism underlying Sbp’s roles in biofilm formation, are incompletely understood. In this work, we used small-angle X-ray scattering (SAXS), NMR, analytical size-exclusion chromatography, and isothermal titration calorimetry analyses to determine the Sbp structure and characterize its interaction with Aap’s Domain-B. We found that Sbp is monomeric and partially folded in solution, and, unexpectedly, we observed no direct interactions between Sbp and Aap Domain-B. Instead, we noted that Sbp forms amyloid fibrils andpartially folded in solution, and, unexpectedly, we observed no direct interactions between Sbp and Aap.

Taken together, the Sbp amyloid fibril structures identified here may account for Sbp’s role as a scaffolding protein in the S. epidermidis biofilm matrix.

Previously regarded as an innocuous human commensal that normally colonized the human skin and mucous membrane, Staphylococcus epidermidis has emerged as a leading opportunistic pathogen causing nosocomial infection associated with permanent or intermittent indwelling medical devices (1, 2). In contrast with its more virulent cousin Staphylococcus aureus, which produces an abundance of toxins, adherence factors, and cytolsins, the main virulence mechanism through which S. epidermidis evade the host immune response relates to its capability of adhering and forming biofilms on biotic and abiotic surfaces (3). Staphylococcal biofilms are adherent multilayered communities of multicellular organisms embedded in self-produced extracellular matrix that are functionally resistant to antibiotics and components of the immune system (4). Therefore, S. epidermidis-mediated infections are difficult to treat and affect millions of patients worldwide annually, causing significant morbidity and mortality and a high burden for the public health system (5). Biofilm formation is a complex, multifactorial process and can be divided into at least three stages: initial attachment to abiotic surfaces, protein-coated materials, and host cells; subsequent cell proliferation and accumulation via cell–cell adhesion leading to a mature biofilm; and finally detachment/dispersal (4, 5). A variety of extracellular matrix components involved in biofilm formation, including proteinaceous adherence factors, polysaccharides, extracellular DNA, teichoic acids, nucleases/proteases, and phenol-soluble modulin factors, have been extensively studied and recently reviewed (6, 7).

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This article contains Figs. S1 and S2 and Tables S1 and S2.

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3 The abbreviations used are: PIA, polysaccharide intercellular adhesin; 2D, two-dimensional; 3D, three-dimensional; Sbp, small basic protein; rSbp, recombinant Sbp; ThT, thioflavin T; BTA-1, 1,2-(4'-methylaminophenyl)benzothiazole; Th3, thioflavin S; SAXS, small-angle X-ray scattering; PDDF, pair distance distribution function; NSD, normalized spatial discrepancy; SEC, size-exclusion chromatography; ITC, isothermal titration calorimetry; AFM, atomic force microscopy; TEM, transmission electron microscopy; \( R_g \), radius of gyration; Aap, accumulation-associated protein; aa, amino acid(s); TEV, tobacco etch virus; \( D_{max} \), maximum distance within the mole-
widely distributed in clinical *S. epidermidis* isolates, has long been identified as a major component of the biofilm matrix and considered as the major intercellular adhesin in staphylococci biofilm formation, not all strains of *S. epidermidis* carry the icaADBC operon, and it is now clear that protein-based matrices also have the ability to mediate biofilm accumulation and cause infection in an ica-independent manner (6, 8). In *S. epidermidis*, among these biofilm matrix proteins, is the well-studied accumulation-associated protein (Aap), which is a cell-wall–anchored, multidomain, multifunctional surface protein with significant roles in biofilm formation (3, 4, 9). Aap consists of an N-terminal Domain-A with a variable number of 16-aa repeats located N terminally and a putative L-type lectin subdomain, a C-terminal Domain-B composed of between 3 and 17 nearly identical 128-aa B repeats terminating in a conserved “half-repeat” motif. Domain-B in Aap is followed by a collagen-like domain and an LPXTG cell wall anchor sequence. Each of these B repeats is composed of two subdomains, G5 and E (10). Recent structural studies revealed that the G5-E repeats can undergo a $\Delta$N$^{2+}$-dependent homophilic interaction, suggesting that two identical Aap molecules expressed on the surface of neighboring cells may interact with each other and promote cell–cell accumulation during biofilm development (11, 12). Besides Aap, specific homophilic interactions have been identified for SrpA (13), IsdC (14), SasG (15), and SdrC (16)–mediated biofilm accumulation; thus, homophilic interactions appear to be a common mechanism by which the matrix proteins promote biofilm formation. Clearly, understanding the molecular mechanism of biofilm matrix proteins involved in biofilm formation is key to combat the significant nosocomial infectious diseases (3, 9).

There is increasing evidence showing that in some bacteria one of the extracellular matrix components includes amyloid fibers (17–19). Amyloids are highly organized protein aggregates that form unbranched protein fibers composed of cross-$\beta$-sheet quaternary structures, which have long been viewed as pathological entities traditionally associated with several incurable neurodegenerative diseases, including Alzheimer’s disease, Parkinson’s disease, Huntington’s disease, and prion diseases (20). Growing evidence, however, indicates that most proteins can adopt the amyloid fold given the correct environment and that some amyloids carry out nonpathological biological roles and are known as functional amyloids (21, 22). Functional amyloids have been found in a wide range of organisms, from bacteria to mammals, with functions as diverse as biofilm formation, development of aerial structures, scaffolding, regulation of melanin synthesis, epigenetic control of polyamines, and information transfer (21). There is a limited but growing number of examples, including the curli homolog in *Salmonella* (23), FapC in many *Pseudomonas* species (24), TasA in *Bacillus subtilis* (25), the phenol-soluble modulins (26) and the biofilm-associated surface protein (27) in *Staphylococcus aureus*, and the adhesin protein P1 in *Streptococcus mutans* (28), found to be able to form amyloid fibrils and that are implicated in biofilm formation (29).

Recently, an 18-kDa extracellular protein called small basic protein (Sbp) was identified in *S. epidermidis* biofilm matrix. Sbp was implicated to serve as a scaffold protein to provide support for cell–cell adhesion and aggregation for both PIA- and Aap-mediated biofilm accumulation (30). Sbp has been found to colocalize with Domain-B of Aap. Besides the probable direct Sbp–Aap Domain-B interactions involved in Aap-mediated biofilm formation, Sbp could also have indirect impact by recruiting additional and yet unknown factors for biofilm formation (30). So far, little is known about the structural function of Sbp and its interaction with Aap, and the precise molecular mechanism underlying its roles in biofilm formation remains largely unexplored.

Here, we report a detailed structural and biophysical characterization of Sbp and its interaction with Aap Domain-B. Sbp is monomeric and partially folded in solution, and unexpectedly, no direct interactions were observed between Sbp and Aap Domain-B under our *in vitro* experimental conditions. Instead, we found that Sbp assembles into amyloid-like fibrils both *in vitro* and *in vivo*. Biochemical, kinetic, and microscopy studies indicate that these fibrils have characteristics of bacterial functional amyloids (18, 31). We propose that amyloid fibril formation of Sbp is responsible for its role as a scaffold protein in *S. epidermidis* biofilm formation.

### Results

**Sbp is monomeric and partially folded in solution**

Previous bioinformatics analysis identified an N-terminal export signal (aa 1–20) for Sbp (30). To study its structure and interaction *in vitro*, a recombinant Sbp (rSbp; aa 21–169) without its export signal was expressed in BL21(DE3) and purified by affinity, ion-exchange, and size-exclusion chromatography. The His$_6$ tag was removed by tobacco etch virus (TEV) protease cleavage. Purified rSbp eluted as a single peak in size-exclusion chromatography (SEC) and was shown as a single band in SDS-PAGE, indicating high homogeneity of the rSbp protein in solution (Fig. S1).

To study the overall conformation of rSbp in solution, we performed small-angle X-ray scattering (SAXS) analysis. The scattering profile, with scattering intensity I(q) plotted against momentum transfer q, and pair distance distribution function (PDDF) transformed from the scattering profile for rSbp are shown in Fig. 1, A and B. The Guinier plot is linear in the Guinier region ($q_{\text{max}} \times R_g < 1.3$) of the scattering profile, indicating that the rSbp is monodisperse and homogeneous in solution. The PDDF is asymmetric with maximum frequency at a distance shorter than half of the maximum distance within the molecule ($D_{\text{max}}$), indicating that rSbp is rather open and elongated in solution. The overall structural parameters, including $R_g$ calculated from Guinier slopes, $R_g$ and $D_{\text{max}}$ from PDDF functions, and molecular weights derived from the power law of volume-of-correlation ($V_r$) (32) and the SAXSMoW package (33), are summarized in Table 1 where the molecular weights from SAXS data are compared with the predicted molecular...
Sbp forms functional amyloid fibrils

- Diagram A: Plot of I(q) vs. q (Å⁻¹) showing the decay of intensity.
- Diagram B: Plot of PDDF vs. r (Å).
- Diagram C: Plot of (qRg)² vs. qRg showing the comparison between Sbp and Lysozyme.
- Diagram D: Plot of q¹(q) vs. q (Å⁻¹) for Sbp and Lysozyme.
- Diagram E: 2D correlation diagram of ¹H (ppm) vs. ¹5N (ppm) for Sbp with a box highlighting the region.
- Diagram F: 2D correlation diagram of ¹H (ppm) vs. ¹5N (ppm) for Lysozyme with a box highlighting the region.

**References:**

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Lysozyme is a well-folded protein serving here as a reference.

A plateau at elevated maximum in the Kratky plot can be observed, and the curve displays shape with a clear maximum. For a completely unfolded protein slightly improves the spectrum (rSbp is only partially folded. Lowering the temperature to 15 °C susceptibility, indicative of unstructured domains (Fig. 1). As shown in Fig. 3A, a minimum at 200 nm and positive ellipticity below 192 nm were observed, suggesting that β-sheets are predominant in Sbp along with regions of low structural complexity. Based on the CD data, an estimation of the secondary structure composition according to Reed’s algorithm (37) is 13% α-helices, 45% β-sheets, and 31% random coil. The tertiary structure of rSbp was further predicted using the I-TASSER web server, resulting in a compact structure with three α-helices packed against a β-sheet (Fig. 3B).

To gain more specific information on the overall 3D structures of rSbp, a low-resolution ab initio shape envelope was built using the program DAMMIN and auxiliary programs. This strategy models a macromolecule as an assembly of scattering beads arranged in space such that a calculated scattering shape envelope was derived using the R_g/V_r power law developed by Rambo and Tainer (32).

No direct interaction between Sbp and Aap Domain-B in vitro

Previously, Sbp was found to be colocalized with Aap Domain-B, suggesting that Sbp may interact with Aap Domain-B directly (30). Domain-B of Aap is a Zn2⁺-binding protein that can mediate intercellular adhesion through Zn2⁺-dependent self-assembly (11, 12). Interestingly, a dose-dependent increase in Sbp binding in the presence of elevated Cl2 concentration has also been observed for immobilized Aap Domain-B in solid-phase binding assays (30). To validate and characterize the interactions between Sbp and Aap Domain-B on a biochemical level, a construct consisting of the terminal intact B repeat along with the C-terminal G5 domain (called Brpt1.5), which represents Aap Domain-B (11), was expressed and purified. The interaction between Brpt1.5 and rSbp was studied in the absence and presence of ZnCl2 using analytical SEC (Fig. 4). Unexpectedly, no interactions between rSbp and peaks disappear possibly due to broadening exchange. The majority of amide proton frequencies are clustered within an area of 1 ppm (8.6–7.6 ppm region) (Fig. S2B), characteristic of intrinsically disordered proteins (IDPs) (35). Increasing the salt concentration up to 400 mM does not improve the spectra significantly; the central region is still heavily overlapped (Fig. S2, C and D), preventing further high-resolution three-dimensional (3D) structure determination of rSbp by NMR.

Table 1

| Data Collection Parameters |
|---------------------------|
| Beam line                 |
| Wavelength (Å)            | 1.280 (APS, ANL) |
| Detector                  | Pilatus 3200 (XAFS) |
| Temperature (K)           | 296 (c)         |
| Exposure time (s)         | 10 (a)          |
| Concentration range (mg/mL) | 1 (b)   |
| Structured Parameters    |
| R_g (Å)                   | 16.9 (a)       |
| I_q (Å⁻²)                 | 17.7 (a)       |
| NSP (a.u.)                | 20.3 (a)       |

Sbp forms functional amyloid fibrils

So far, no experimentally determined high-resolution or homologous 3D structural information is available for Sbp. We therefore first predicted the secondary structure of Sbp (aa 1–169) using the PSIPRED web server (36), which reveals extensive β-sheet regular secondary structure interspersed with several α-helices for Sbp (Fig. 2). To verify the secondary structure composition, a circular dichroism (CD) spectrum was recorded for rSbp in solution. As shown in Fig. 3A, a minimum at 200 nm and positive ellipticity below 192 nm were observed, suggesting that β-sheets are predominant in Sbp along with regions of low structural complexity. Based on the CD data, an estimation of the secondary structure composition according to Reed’s algorithm (37) is 13% α-helices, 45% β-sheets, and 31% random coil. The tertiary structure of rSbp was further predicted using the I-TASSER web server, resulting in a compact structure with three α-helices packed against a β-sheet (Fig. 3B).

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Brpt1.5 were observed either in the absence or presence of Zn²⁺. As shown in Fig. 4A, both protein components in the sample mixtures of Brpt1.5 and rSbp (with molar ratios of 1:2, 2:2, and 4:2, respectively) eluted at their respective retention volumes as for each protein alone through the Superdex™ 75 SEC column. The rSbp binding property of Brpt1.5 was further verified using isothermal titration calorimetry (ITC). Again, as shown in Fig. 4B and C, rSbp does not bind with Brpt1.5 either in the absence or presence of Zn²⁺.

**Sbp forms amyloid fibrils in vitro**

Because no direct interaction between rSbp and Aap Domain-B was observed, Sbp may utilize a different modality to contribute biofilm formation. Given its richness of β-sheet secondary structure and partial folding characteristic, we became interested in its propensity to form amyloid fibrils. A variety of bioinformatics tools have been developed to identify amyloidogenic hot spots within proteins that are prone to aggregation and may induce amyloid formation. The propensity of Sbp to form amyloid fibrils was analyzed using AMYLPRED2 (38, 39), which determines amyloidogenic regions based on the consensus score of 11 such tools and thus provides a reliable estimate of the amyloid propensity. Consensus short peptide segments that might be responsible for amyloid fibril formation were predicted for Sbp (Fig. 2), suggesting that Sbp has high potential to form amyloid fibrils.

The amyloidogenic nature of rSbp was first confirmed using Congo red (CR) binding assays specific for amyloids (Fig. 5A). In *in vitro* assays, fibrillization is routinely triggered by mechanical agitation to increase the probability of the intermolecular collisions necessary to form the initial nucleus. The binding of CR to rSbp that had been agitated at 37 °C for 12 h was evaluated. As shown in Fig. 5A, compared with CR alone, the spectra of the agitated rSbp sample demonstrated dye binding and a shift in the CR absorbance maxima that is characteristic of amyloid fibrils. We speculated that, under this condition, rSbp assembled into an amyloid fibrillar state, hereafter named as rSbpfib (as opposed to the native, soluble Sbpsol). Based on Fou-
rrier transform IR (FTIR) spectroscopy data, distinct structural differences were observed between rSbpsol and rSbpfib (Fig. 5B). The amide I region (1,600–1,700 cm$^{-1}$) reflects the vibrational stretching of the amide backbone and is indicative of secondary structure (40). The shift in the β-sheet peaks to lower wave-number (from 1631 cm$^{-1}$ in Sbpsol to 1624 cm$^{-1}$ in Sbpfib) is indicative of rearrangement of globular β-sheets to extended β-sheets, which is consistent with a previous report of amyloid fibrils (40).

The kinetics of Sbpfib formation was monitored by thioflavin T (ThT) fluorescence at 37 °C. As shown in Fig. 6A, Sbpfib formation follows a typical sigmoidal time profile, suggesting that Sbpfib undergoes a nucleation-dependent polymerization process characterized by an initial lag phase reflecting nucleation and a subsequent elongation/growth phase culminating in a steady plateau (20). We also performed time-resolved atomic force microscopy (AFM) imaging of the system to ascertain that Sbp forms amyloid fibrils. Starting from the homogeneous solution of rSbp monomer, rSbp showed amorphous structure at times 0 and 2 h; isolated protofilament or short fibers were not seen until at 4 h. Upon progression of time, more and longer fibers were formed, and fiber branching was observed at 8 h of incubation (Fig. 6, B–E). The aspect ratio, which was defined as the ratio of fibril length with respect to fibril diameter (41), varies significantly for rSbp fibrils. As shown in Fig. 6E, the diameter of the fibril by AFM is about 3 nm; the aspect ratio of one short and long fibril can be about 84–373. After 12 h of incubation, rSbp formed a mature, intertwined fibril network (Fig. 6F). The formation and morphology of rSbp fibrils were further confirmed with transmission electron microscopy (TEM) analysis of mature rSbp fibrils at 12, 24, and 48 h, respectively (Fig. 6, G–I). The mature fibril structure usually exhibits a width of about 5 nm and a length of 100–5,000 nm.

Sbp undergoes further unfolding in lag phase of fibrillation

To characterize the structural changes of rSbp in lag phase, we performed in situ SAXS measurement of the fibrillation reaction of rSbp at 37 °C. The respective time-dependent scattering profiles and the PDFFs are shown in Fig. 7, A and B. To roughly evaluate changes in the profiles, apparent molecular weights were calculated using the scattering profiles as input by the SAXSMoW package, which reveals gradually increased

Figure 3. Secondary structure and low-resolution 3D structure of rSbp. A, the far UV CD spectrum of rSbp. B, the 3D atomic model of rSbp predicted by the I-TASSER online server, revealing three helix bundles packed against a β-sheet. C, the ab initio shape envelope in two views for rSbp, reconstructed by DAMMIN, is elongated. The atomic model in B was superimposed onto the shape envelope. mdeg, millidegrees.
molecular weight over time but that are still smaller than the molecular weight of a dimer (Table S2), suggesting that the monomeric rSbp dominates during the lag phase. The $D_{\text{max}}$ as well as the dominant peaks in PDDFs shift to larger distances over time, implying obvious structural changes, which were supported by the Kratky plots (Fig. 7C). At the starting point time, the Kratky plot of rSbp has a curving-down feature, suggesting a partial folding structure; however, at a time of 2 h, the Kratky plot has a clear plateau and continues to increase over the $q$ range $0.18-0.3\ \text{Å}^{-1}$, reflecting the predominantly extended conformation characteristic of unfolded rSbp. Therefore, rSbp undergoes further unfolding during the lag phase of fibrillation.

**Sbp forms intracellular amyloid aggregates in Escherichia coli cells**

It has been shown that amyloidogenic proteins from different origins, when expressed in *E. coli*, are accumulated as cytoplasmic inclusions of amyloid nature (42). To investigate whether Sbp can form amyloid aggregates in *vivo*, *E. coli* cells carrying pSbp plasmid were grown at 37 and 16 °C, respectively, until early exponential phase was reached, and then the expression of Sbp was induced. At defined time points (*i.e.* 4, 8, and 12 h), aliquots of induced cells were sampled and fixed with 4% PFA, stained with the probes BTA-1 and thioflavin S (ThS) that specifically bind to amyloid aggregates, and visualized by confocal microscopy. As shown in Figure 8, a variable number of inclusions were observed in cells grown at 37 °C at specific induction time intervals that were recognized by both amloidophilic dyes as intense fluorescent foci. In contrast, cells grown at 16 °C do not accumulate inclusions.

**Sbp is essential for *S. epidermidis* biofilm formation**

Following the observation of amyloid fibril formation of overexpressed Sbp in *E. coli* at 37 °C, to test the relevance of Sbp
Sbp forms functional amyloid fibrils

In this study, we provide a detailed structural characterization of an extracellular protein called Sbp in *S. epidermidis* and demonstrate its ability to form amyloid fibrils using different biophysical techniques. CD, SAXS, and NMR analyses show that Sbp is a β-sheet–rich protein adopting a monomeric and partially folded structure at physiological condition. Alternatively, Sbp can aggregate into amyloid fibrils, which was confirmed and characterized by fluorescence spectroscopy and microscopy both in vitro and in vivo.

We initially attempted to determine the high-resolution structure of Sbp using NMR. If well-folded, Sbp should be suitable for NMR study given its medium size. Both secondary and tertiary structure predictions suggest that Sbp could be a well-folded protein consisting of regular secondary structures. However, our NMR and SAXS studies indicate that Sbp is partially folded at near physiological condition. At elevated temperature (37 °C), Sbp shows characteristics of IDPs. This property may relate to its high propensity to form amyloid fibrils at elevated temperature in vitro, which seems relevant to *S. epidermidis* biofilm formation in vivo. Many amyloidogenic proteins such as α-synuclein, Aβ(1–42), SOD1, and TDP-43, which are involved in neurodegenerative diseases, are partially folded or intrinsically disordered proteins that lack definite ordered 3D structure (43). Structural flexibility and plasticity originating from the lack of a definite ordered 3D structure are believed to represent the major functional advantages for these proteins, enabling them to interact with a broad range of binding partners, including self-aggregation to form secondary structures and subsequently fibers with other proteins, membranes, nucleic acids, and various small molecules. One example is the human hormone islet amyloid polypeptide (hIAPP), an IDP that functions in glucose homeostasis (44). Monomeric hIAPP is primarily, but not completely, unstructured. The interaction of hIAPP with membrane promotes formation of secondary structures that nucleate aggregation; this process in turn induces membrane disruption via a two-step mechanism (44, 45). Sbp appears to be a partially unfolded protein with an extended coil-like conformation. As our time-resolved SAXS measurement shows, in the lag phase of fibrillation, Sbp undergoes a further unfolding process, which may expose the amyloidogenic hot spot to form the nucleation sites leading to polymerization.

It was proposed that Sbp fosters PIA- and Aap-mediated biofilm accumulation by forming a biofilm scaffold instead of directly inducing cell aggregation. Recently, Sbp was identified by affinity purification from *S. epidermidis* crude biofilm matrix preparations using recombinant Aap Domain-B as a ligand, suggesting a direct Sbp–Aap Domain-B interaction (30). However, the exact modalities of Sbp–Aap Domain-B interaction remain unclear. It was reported that Aap Domain-B is a zinc-binding protein and undergoes Zn²⁺–induced self-association, suggesting the importance of Zn²⁺ in Aap function. We expressed and purified both Sbp and the Aap Domain-B consisting of 1.5 G5 repeats and characterized the interactions using SEC and ITC both in the absence and presence of Zn²⁺. Unexpectedly, no obvious interaction of Sbp and Aap Domain-B was observed in our condition. Given that Sbp can adopt different folding states, either the partially folded monomer or the amyloid fibril state, there is a possibility that the Sbp monomeric state is not the active form. Notably, PAP(248–286), a peptide fragment of prostatic acid phosphatase, a protein abundant in human semen, is inactive in the monomeric state and must first aggregate to form the active form, semen enhancer of viral infection, with characteristics of amyloid fibrils to promote viral infection. Bacterial amyloid proteins like...
curli can promote the aggregation of PAP(248–286) into semen enhancer of viral infection (46). It is of interest in future studies to determine whether Sbp in crude biofilm matrix preparations forms amyloid fibrils, therefore presenting the active form for Aap Domain-B binding, or whether Aap Domain-B may interact with Sbp by enhancing the aggregation of Sbp into functional amyloid.

Sbp is the first structural S. epidermidis extracellular biofilm matrix protein with significant relevance for both Aap- and PIA-dependent biofilm formation. It is now recognized that several staphylococcal surface proteins, including Aap, can promote biofilm accumulation in a PIA-independent manner through specific homophilic interactions between proteins expressed on different cells (9). Recent studies have recognized amyloid fibers as common functional components in bacterial biofilm matrix. There is a growing number of examples demonstrating the direct implication of amyloids in biofilm formation, including the phenol-soluble modulin and Bap protein from Staphylococcus aureus (29). Bap is a large protein consisting of multiple domains whose function is regulated post-translationally (47). It was found that Bap forms an amyloid scaffold under acidic solution conditions to promote biofilm develop-

Figure 6. Kinetics of amyloid fibrillization of rSbp at 37 °C. A, kinetics of rSbp amyloid fibril formation was monitored by in situ measurement of ThT fluorescence. B–F, kinetics of rSbp amyloid fibril formation probed by AFM. One short (green) and one long (red) fibril in E selected for aspect ratio analysis are marked with arrows. The images are shown as height traces, and the scale bar corresponds to 500 nm. G–I, transmission electron micrographs of negatively stained rSbp amyloid fibrils. Scale bars were set to 100 nm.
To our knowledge, Sbp is the first *S. epidermidis* protein identified to have the propensity to assemble into amyloids at near physiological conditions both *in vitro* and *in vivo*. We propose that the formation of functional amyloid fibrils by Sbp may account for its role as a scaffolding protein in *S. epidermidis* biofilm formation. Amyloid formation may represent another common mechanism in protein-mediated cell–cell accumulation. We expect more such proteins to be identified.

**Experimental procedures**

**Protein expression and purification**

The genomic DNA of *S. epidermidis* 1457 was extracted for gene cloning using the EasyPure Bacteria Genomic DNA kit (TransGen Biotech). The DNA fragments encoding rSbp (aa 21–169) and Brpt1.5 were amplified from the genomic DNA by PCR using the appropriate primers (Table S1). The PCR products were then subcloned into pET28a (for rSbp) and pGEX-4T-2 (for Brpt1.5), respectively. The positive clones were confirmed by DNA sequencing and subsequently transformed into *E. coli* BL21(DE3).

For purification of rSbp, cells were grown in LB culture medium at 37 °C until *A*₆₀₀nm reached ~0.8. Expression of the rSbp recombinant protein was induced with 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) at 16 °C for another 20 h before harvesting. Bacteria were collected by centrifugation at 4,500 g for 30 min. The cell pellet was suspended in lysis buffer (20 mM Tris-HCl, pH 7.4, 500 mM NaCl) and then homogenized by a high-pressure cell disruptor system. After centrifugation at 18,500 × g for 30 min, the supernatant was loaded onto a nickel-nitrilotriacetic acid affinity resin that was pre-equilibrated with binding buffer (20 mM Tris-HCl, pH 7.4, 500 mM NaCl). The resin was extensively washed by two different buffers (20 mM Tris-HCl, pH 7.4, 1 M NaCl, 20 mM imidazole and 20 mM Tris-HCl, pH 7.4, 200 mM NaCl, 20 mM imidazole) sequentially to remove nucleic acid contamination. The Sbp protein was then eluted with 20 mM Tris-HCl, pH 7.4, 200 mM NaCl, 250 mM imidazole. The protein fractions were mixed and digested by TEV protease to remove the His tag. Next, the protein was further purified using an ion-exchange HiTrap S column and a HiLoad16/600 Superdex 75 PG column pre-equilibrated with gel filtration buffer (20 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1 mM DTT). The final sample purity was checked by SDS-PAGE and MS (MALDI-TOF).

For purification of Brpt1.5, cells were grown in LB culture medium supplemented with 100 µg/ml ampicillin at 37 °C until *A*₆₀₀nm reached ~0.8. Expression of Brpt1.5 was induced with 0.5 mM IPTG at 37 °C for 5 h. The cells were harvested, suspended, and lysed using a similar protocol as described above. After centrifugation at 18,500 × g for 30 min, the supernatant was loaded onto a GST column. The resin was extensively washed with binding buffer. The target protein was eluted with 20 mM Tris-HCl, pH 7.4, 200 mM NaCl, 25 mM GSH. The GST tag was removed with TEV protease treatment. After cleavage, the protein was loaded onto a GST column to remove uncleaved protein and the free cleaved GST tag. The flow-through was collected, concentrated, and injected onto a HiLoad16/600 Superdex 75 PG column pre-equilibrated with gel filtration buffer. The final sample purity was checked by SDS-PAGE.
Structural analysis by small angle X-ray scattering

Conventional and in situ time-resolved SAXS measurements were carried out at room temperature and 37 °C, respectively, at beamline 12 ID-B of the Advanced Photon Source, Argonne National Laboratory, equipped with a temperature controller. The scattered X-ray photons were recorded with a PILATUS 1M detector (Dectris). The setups were adjusted to achieve scattering $q$ values of $0.005 < q < 0.89 \text{Å}^{-1}$ where $q = (4\pi/\lambda) \sin \theta$ and $2\theta$ is the scattering angle. Thirty two-dimensional images were recorded for each buffer or sample solution using a flow cell with an exposure time of 0.5–2 s to minimize radiation damage and obtain good signal-to-noise ratio. No radiation damage was observed as confirmed by the absence of systematic signal changes in sequentially collected X-ray scattering images. The 2D images were reduced to one-dimensional scattering profiles using Matlab scripts on site. Scattering profiles of the protein were calculated by subtracting the background buffer contribution from the sample buffer profile using the program PRIMUS \cite{48} following standard procedures \cite{49}. Concentration series measurements (4- and 2-fold dilution and stock solutions) for the same sample were carried out to remove the scattering contribution due to interparticle interactions and to extrapolate the data to infinite dilution. The forward scattering intensity $I(0)$ and the radius of gyration ($R_g$) were calculated from the data of infinite dilution at low $q$ values in the range of $q R_g < 1.3$ using the Guinier approximation: $\ln I(q) = \ln I(0) - R_g^2 q^2/3$. These parameters were also estimated from the scattering profile with a broader $q$ range of $0.006 - 0.30 \text{Å}^{-1}$ using the indirect Fourier transform method implemented in the program GNOM \cite{50} along with the PDDF, $p(r)$, and $D_{\text{max}}$. The parameter $D_{\text{max}}$ (the upper end of distance $r$) was chosen so that the resulting PDDF has a short, near-zero-value tail to avoid underestimation of the molecular dimension and consequent distortion in low-resolution structural reconstruction. The $V_c$

![Figure 8. Formation of intracellular amyloid inclusions in E. coli cells expressing rSbp at 16 and 37 °C, respectively. Cells sampled after 4, 8, and 12 h of induction, respectively, were fixed with PFA, then stained with either BTA-1 or ThS, and visualized by confocal microscopy. BF, bright field. Scale bar, 1 μm.](image)
values were calculated using the program Scatter, and the molecular weights of solutes were calculated on a relative scale using the \( R_g/V_c \) power law developed by Rambo and Tainer (32), independent of protein concentration and with minimal user bias.

Low-resolution ab initio bead models were built up with the program DAMMIN, which generates models represented by an ensemble of densely packed beads (51), using scattering data within the \( q \) range of 0.006 – 0.30 Å\(^{-1}\). Thirty-two independent runs were performed, and the resulting models were subjected to averaging by DAMAVER (52), superimposed by SUPCOMB (53) based on the normalized spatial discrepancy (NSD) criteria, and filtered using DAMFILT to generate the final model. NSD is a measure of quantitative similarity between sets of three-dimensional models: if two models systematically differ from each other, their NSD exceeds 1; for identical objects, it is 0.

NMR spectroscopy

A similar preparation protocol as above was followed to prepare \(^{13}\)C,\(^{15}\)N-labeled rSbp except that cells were grown and induced in M9 minimal medium with \(^{15}\)NH\(_4\)Cl and \([^{13}\text{C}_6]\)glucose as the sole nitrogen and carbon sources, respectively. The samples used for NMR experiment were as follows: 0.5 mM \(^{13}\)C,\(^{15}\)N-labeled-rSbp in 90% H\(_2\)O, 10% D\(_2\)O containing 20 mM K\(_2\)HPO\(_4\)/NaH\(_2\)PO\(_4\) \(\times 1\), 100 mM NaCl, 50 \(\mu\)M NaN\(_3\), pH 7.20, and additives. 2D \(^1\)H-\(^{15}\)N HSQC spectra were recorded on Bruker Avance 600- or 800-MHz spectrometer at variable temperature, all equipped with four radio frequency channels and a triple-resonance cryoprobe with pulsed field gradients. Each spectrum comprised 1,024 complex points in the \( t_2 \) dimension and 160 complex points in the \( t_1 \) dimension. All spectra were processed using the software package NMRPipe (54) and analyzed with NMRView (55).
Sbp forms functional amyloid fibrils

CD spectroscopy

The secondary structure of rSbp in solution was checked by CD on an Applied Photophysics Chirascan-plus (Leatherhead) controlled by Pro-Data Chirascan v4 software. Far-UV CD spectra were recorded between 190 and 260 nm with a step resolution of 1 nm, a slit width of 0.6 nm, and an integration time of 5 s. Acquisition was performed at 25 °C using a 0.2-mm-path length cuvette with a protein concentration of ~0.3 mg/ml. The spectra were averaged over three scans and corrected by subtraction of the buffer signal.

Secondary and tertiary structure prediction

The secondary structure of Sbp was predicted using the PSIPRED web server (36). The tertiary structure of Sbp was predicted using the I-TASSER online server (56).

Analytical size-exclusion chromatography

Samples of Sbp alone, Brpt1.5 alone, and the mixture of Sbp and Brpt1.5 with molar ratios of 1:1 and 2:1 were injected onto a Superdex 75 10/300 GL column (24.0 ml; GE Healthcare) pre-equilibrated with binding buffer (20 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1 mM DTT or 20 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1 mM DTT, 1 mM ZnCl₂, respectively). Concentrations of Brpt1.5 protein were kept constant at 30 μM.

Isothermal titration calorimetry

Purified Sbp and Brpt1.5 samples were extensively exchanged into buffer (20 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1 mM DTT or 20 mM Tris-HCl, pH 7.4, 100 mM NaCl, 1 mM DTT, 1 mM ZnCl₂, respectively) using a Superdex 75 10/300 GL column and then concentrated to 30 μM for Brpt1.5 and 300 μM for Sbp. The ITC measurements were performed with a MicroCal iTC 200 calorimeter (GE Healthcare) at 25 °C. The background data obtained from the buffer sample were subtracted before the data analysis. The data were fitted using the Origin7 software package (MicroCal). Measurements were repeated twice, and similar results were obtained.

In vitro amyloid fibril formation

For amyloid fibril formation, rSbp at a concentration of 250 μM in 20 mM Tris-HCl, pH 7.2, containing varied concentrations of NaCl and 0.01% sodium azide was incubated at 37 °C with rotary agitation at 220 rpm. The rSbp solution after salt-induced fibrillation was subjected to CR assay, FTIR characterizations, CD spectroscopy, ThT fluorescence assay, AFM, and TEM characterization at certain time points.

CR assay

The assay was performed with an Implen UV-visible nanophotometer (München, Germany). The concentration of CR solution was 5 μM. The buffer was 10 mM phosphate saline buffer, pH 7.4. The assay was performed ex situ. 20 μl of rSbp solution was added into 980 μl of CR solution in a 1.0-cm quartz cuvette. The solution was incubated in the dark for 30 min. The solution in the cuvette was shaken first before each spectral acquisition.

FTIR spectroscopy

The FTIR spectra were recorded using a Bruker Vertex 70 FTIR spectrometer equipped with a DLaTGS detector. A Bruker-made sample shuttle was installed inside the sample compartment, which has two sample holders: one for sample and the other for reference. The FTIR spectra of the samples were collected with 4 cm⁻¹ resolution and 32 scans. 100 μl of solution was deposited onto the CaF₂ windows, and then the CaF₂ window surface was dried with a dryer.

ThT binding assay

The ThT binding assay was performed with a Hitachi F-7000 fluorescence spectrophotometer (Tokyo, Japan). An excitation wavelength of 450 nm with a slit width of 5 nm was used. The emission spectrum in the region of 460–600 nm was measured using a slit width of 10 nm. The concentration of ThT solution was 10 μM. The buffer was 20 mM Tris-HCl, 100 mM NaCl, pH 7.2. The assay was performed ex situ. For each measurement, 1 μl of rSbp solution was added into 1 ml of ThT solution in a 1.0-cm quartz cuvette. The solution in the cuvette was shaken first before each spectral acquisition.

Atomic force microscopy

Time-dependent morphological changes during rSbp aggregation were measured using an atomic force microscope (SPM P47 Solver, NT-MDT Co., Zelenograd, Russia) in tapping mode. At regular time intervals during aggregation, small aliquots of the incubated rSbp samples were taken out and diluted with deionized water to a final concentration of 40 μM. A 50-μl aliquot of the diluted solution was deposited onto a freshly cleaved mica surface followed by incubation at room temperature for 5 min. Washing steps were carried out twice with deionized water to remove unbound proteins/aggregates followed by drying in a desiccator. All AFM images were taken on dried samples in air, and 5 × 5 μm scanning was used throughout the AFM experiment.

Transmission electron microscopy

The formation and morphology of amyloid fibrils by Sbp were confirmed by negatively staining samples. rSbp fibrils were diluted to 0.5 mg/ml, and 3 μl of sample was directly deposited onto a glow-discharged 300 mesh carbon-coated copper grid (BZ10023a, Zhongjingkeyi, China) followed by staining with 2% (w/v) uranyl acetate and blotted dry with filter paper three times. The stained samples were imaged using a Tecnai Spirit with iCorr D1319 (FEI) operating at 120 kV at a magnification of ×52,000.

Knockout of Sbp by Gateway system in S. epidermidis 1457

An allele replacement strategy was used for construction of a specific Sbp knockout mutant in S. epidermidis 1457 as reported previously (30). Briefly, two primers flanking regions upstream (sbp_ko_att, 5'-GGGGACAGTTTTGTACAAAAAAGCAGGCTTATCTCTTGACTGTTG-3' and sbp_eco_r, 5'-ACCGCGGAATCTCACTCTTCCCCATTCAAGCTCTTGTCTTCTG-3') and reverse primers covering downstream regions (sbp_ko_eco, 5'-ACCGCGGAATTCAAGATAAAA-
ATGTGAAGTTATATCATA-3' and sbp_att_r, 5'-GGGGA-
CCACCTTGTGACAAGAAGCTGGATGTCGACG-
TAAACGT-3') of the Sbp-coding region were amplified fol-
lowing a standard PCR protocol. Purified PCR products were
cloned using EcoRI, and the resulting, sbp-flanking fragments
were ligated with T4 DNA ligase. The 5' and 3' attB site ligation
products were introduced into pKOR1 using BP clones (Invit-
rogen, Karlsruhe, Germany) and transformed into E. coli
DC10B. Recombinant plasmid pKOR1Δsbp was verified by
PCR and DNA sequencing.

For electrocompetent cells, S. epidermidis 1457 were grown
in tryptic soy agar medium until A_{600 nm} reached 0.8. Cells
were harvested and washed three times with double-distilled H_2O
by centrifugation at 3,000 \times g for 10 min at room temperature.
Cells were resuspended and washed with 2 ml of 2% glycerol
twice and finally resuspended with 10% glycerol. Electropora-
tion was performed using a Bio-Rad Gene Pulser and Pulse
Controller using a standard protocol (57). Briefly, 10 \mu g of
pKOR1 was added into 50 \mu l of competent cells and incubated
on ice for 30 min before transfer to a prechilled 0.2-cm-gap
electroporation cuvette. The cells were immediately pulsed
with 2.1-kV voltage and diluted in 800 \mu l of B2 culture medium
followed by shaking at 30 °C for 3 h. Transformation products
were plated on a TSB_{Kan} plate and grown at 37 °C for 48 h.
Clones verified correct by PCR were transferred into 3 ml of
TSB_{m10} medium at 30 °C overnight followed by dilution at a
1:100 volume ratio into 50 ml of TSB_{m10} medium and shaking
at 42 °C overnight. Cells were diluted at a 1:100 volume ratio
into 5 ml of TSB_{m5} medium and shaken at 42 °C overnight,
twice followed by plating on a TSB_{m5} plate, and then incubated at 2 °
overnight. Single clones were picked into 5 ml of TSB medium
and shaken at 30 °C overnight. Cell were diluted 1:10,000,
plated on a TSB_{ATP} plate, and grown at 37 °C overnight. Ten
cloned were picked and inoculated into both a TSB plate and a
TSB_{m10} plate. Three clones that only grew on the TSB plate
were picked, and genomic DNA was extracted for PCR verifi-
cation and sequencing.

**Cell fixation and staining for confocal microscopy**

Cells were grown at 37 or 16 °C with shaking at 220 rpm until
A_{600 nm} reached 0.8 and then induced with 0.5 mM IPTG for
another 12 h. At regular time intervals, 500 \mu l of cell cultures
was sampled and centrifuged at 3,000 \times g for 4 min at room
temperature, and the pellets were washed twice with the same
volume of PBS buffer for the following fixation and staining
steps. For BTA-1 staining, the washed cells were resuspended
in 250 \mu l of 4% PFA, incubated for 30 min at room temperature,
then washed twice with 250 \mu l of PBS buffer, resuspended in 1
\text{mm} BTA-1 prepared in 100% ethanol, and incubated for 30 min
at room temperature. Finally, the stained cells were washed
twice with PBS again. For ThS staining, the washed cells were
suspended in 250 \mu l of 0.05% (w/v) ThS in 12.5% ethanol,
icubated for 1 h at room temperature, and then washed
three times with 250 \mu l of PBS buffer. Stained cells were
suspended in the appropriate volume of PBS buffer and
observed under confocal microscopy.

**Confocal microscopy**

Microscopy observations were performed using an LSM 780
(Leica) inversion confocal microscope with a 63×/numerical
aperture 1.40 oil immersion objective. BTA-1 fluorescence was
excited with a 405-nm laser diode, and the emission was regis-
tered between 410 and 495 nm. ThS fluorescence was excited
using a 488-nm argon laser, and the emission was registered in
a range from 493 to 630 nm.

**Structural illustration**

All of the structural illustrations were generated using the
PyMOL Molecular Graphics System, version 1.3 (Schrödinger,
LLC).

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