





TapA acts as specific chaperone in TasA filament formation by strand complementation

Yvette Roske^{a,1}⁽¹⁾, Florian Lindemann^{b,1}⁽¹⁾, Anne Diehl^{b,1}⁽¹⁾, Nils Cremer^b, Victoria A. Higman^c⁽¹⁾, Brigitte Schlegel^b, Martina Leidert^b, Kristina Driller^{d,e}, Kürşad Turgay^{d,e}⁽¹⁾, Peter Schmieder^b⁽¹⁾, Udo Heinemann^{a,f⁽¹⁾}, and Hartmut Oschkinat^{b,f,2}⁽¹⁾

Edited by Robert Tycko, National Institute of Diabetes and Digestive and Kidney Diseases, Bethesda, MD; received October 12, 2022; accepted March 8, 2023

Studying mechanisms of bacterial biofilm generation is of vital importance to understanding bacterial cell-cell communication, multicellular cohabitation principles, and the higher resilience of microorganisms in a biofilm against antibiotics. Biofilms of the nonpathogenic, gram-positive soil bacterium Bacillus subtilis serve as a model system with biotechnological potential toward plant protection. Its major extracellular matrix protein components are TasA and TapA. The nature of TasA filaments has been of debate, and several forms, amyloidic and non-Thioflavin T-stainable have been observed. Here, we present the three-dimensional structure of TapA and uncover the mechanism of TapA-supported growth of nonamyloidic TasA filaments. By analytical ultracentrifugation and NMR, we demonstrate TapA-dependent acceleration of filament formation from solutions of folded TasA. Solid-state NMR revealed intercalation of the N-terminal TasA peptide segment into subsequent protomers to form a filament composed of β -sandwich subunits. The secondary structure around the intercalated N-terminal strand $\beta 0$ is conserved between filamentous TasA and the Fim and Pap proteins, which form bacterial type I pili, demonstrating such construction principles in a gram-positive organism. Analogous to the chaperones of the chaperone-usher pathway, the role of TapA is in donating its N terminus to serve for TasA folding into an Ig domain-similar filament structure by donor-strand complementation. According to NMR and since the V-set Ig fold of TapA is already complete, its participation within a filament beyond initiation is unlikely. Intriguingly, the most conserved residues in TasA-like proteins (camelysines) of Bacillaceae are located within the protomer interface.

Bacillus subtilis | biofilm | TasA | TapA/YqxM | structure

Most microorganisms form sessile multicellular biofilms in which they are protected against stress by a gel-like matrix composed of proteinaceous fibrils, various extracellular polysaccharides and often DNA (1). *Bacillus subtilis* expresses TasA (translocation-dependent antimicrobial spore component) as the major biofilm protein (2), which aided by TapA (TasA anchoring/assembly protein, YqxM) (3) can convert into stable oligomeric structures to form the biofilm matrix (4, 5). Whereas biofilms often contain β -helix type protofibrils that can be stained by Thioflavin T (ThT), e.g., curli of *Escherichia coli* (6), TasA forms different architectures in vitro depending on experimental conditions (2, 7–9). ThT-stainable fibrils were initially generated from acid-treated protein at low pH (2) and recently also from unfolded protein at neutral pH (10), whereas TasA from natural sources was found early on to form stable, nonamyloid oligomers (7, 9).

The importance of TapA (referring to the mature protein, residues 44 to 253, if not otherwise indicated) for robust biofilm formation was already recognized when TasA (mature protein, residues 28 to 261) was identified as the major biofilm protein of *B. subtilis* (2, 11, 12). A role as polymerization nucleator was proposed for TapA (3, 13), whereby its N terminus (14, 15) is considered essential. Outside the cell, TapA is found to colocalize with the membrane (13), and individual observations also identified it in fibrils (13). TapA is present at a much lower abundance than TasA, similar to the signal peptidase SipW (13). The coexpression of TapA and SipW and the overlap of their stop and start codons, respectively, hints at a functional and structural association (16). TapA was characterized as a two-domain protein with an unstructured N terminus (17).

Recently, a structural basis for the different TasA forms became available. A three-dimensional structure of monomeric TasA_{G28-239} was obtained by X-ray crystallography (8). ThT-stainable fibrils were investigated by solid-state NMR (10), and in parallel to this study nonamyloid filaments by cryo-EM (18) were studied at a resolution of 3.5 Å. Furthermore, a structural relationship between archaeal and bacterial biofilms was suggested on the basis of a 4.0 Å cryo-EM structure of *Pyrobaculum calidifontis* bundling pili (19).

Significance

Bacteria on surfaces often protect themselves by forming biofilms, with huge negative impact on health and the economy. Biofilms are stabilized by protein-based super structures in the extracellular matrix and means to suppress or resolve them are urgently needed given the looming health-crisis due to antibioticresistant bacterial strains. Our model organism Bacillus subtilis has disease-causing relatives such Bacillus cereus and Bacillus anthracis, and thus provides a means to investigate common biofilm formation mechanisms. It produces two biofilm proteins, TasA (major component) and TapA (minor component). We have analyzed TapA-triggered filament formation by TasA and find that protomers connect via the donor-strand complementation mechanism of gram-negative bacteria. These results form the basis for the development of effective strategies against biofilm formation.

The authors declare no competing interest.

This article is a PNAS Direct Submission.

Copyright © 2023 the Author(s). Published by PNAS. This open access article is distributed under Creative Commons Attribution License 4.0 (CC BY).

¹Y.R., F.L., and A.D. contributed equally to this work.

 ^2To whom correspondence may be addressed. Email: oschkinat@fmp-berlin.de.

This article contains supporting information online at https://www.pnas.org/lookup/suppl/doi:10.1073/pnas. 2217070120/-/DCSupplemental.

Published April 17, 2023.

Author contributions: Y.R., A.D., K.T., and H.O. designed research; Y.R., F.L., A.D., N.C., V.A.H., B.S., M.L., K.D., and P.S. performed research; Y.R., F.L., A.D., N.C., V.A.H., K.D., K.T., P.S., U.H., and H.O. analyzed data; and Y.R., F.L., A.D., K.T., and H.O. wrote the paper.

TasA homologs (called camelysins, SI Appendix, Fig. S1) occur in various branches of bacteria and archaea (19) but TapA is not even conserved within the Bacillus genus. On the other hand, Bacillus cereus has two camelysins, calY1 and calY2, encoded in one operon together with the specific signal peptidase SipW (sip W-calY1-bc_1280-calY2) instead of the tapA-sip W-tasA combination found in B. subtilis (SI Appendix, Fig. S1, entries for B. cereus in lines 2 and 3). It is likely that one of the two camelysins in *B. cereus* adopts the role as TapA in *B. subtilis* (3). Since bacteria take advantage of mixed biofilms, sometimes even involving fungi (4, 20), conservation of TasA-like proteins is highly informative. B. cereus, for example, induces expression of B. subtilis biofilm genes via the secretion of thiocillins (21). Cross-kingdom biofilm formation of *B. subtilis* involving hyphae of fungi such as *Aspergillus* niger and Agaricus bisporus has also been linked to TasA (22). In this context, it could be useful to identify indicators for joint biofilm formation.

Our aim is to understand the role of TapA in *B. subtilis* biofilm formation and to derive a mechanism for TapA-dependent TasA filament formation. Interactions of TapA with folded TasA are initially monitored by analytical ultracentrifugation (AUC) in a quantitative manner, supplemented by solution NMR data. To provide insight from a structural point, we determined the X-ray structure of a TapA core domain construct and the secondary structure of TasA filaments by solid-state NMR, based on near-complete backbone resonance assignments. Crucial intermolecular distances within the filament were monitored by mixing two TasA proteins with different labelling patterns. Based on our biophysical and structural results, a mechanism of TapA-initiated TasA filament formation is presented. For the gram-positive bacteria system, we demonstrate that TapA initiates TasA oligomerization similar to the chaperones of the chaperone-usher pathway of pilus assembly in gram-negative bacteria.

Results

TapA Enhances TasA Oligomerization. Solutions of monomeric TasA are known to be stable at neutral pH for several days or weeks, but may form non-ThT-stainable filaments when concentrated over a membrane composed of regenerated cellulose (8, 9). By contrast, stainable fibrils, presumably amyloid-like, are formed at low pH from unfolded protein as shown by Romero et al. (2). TapA is known to induce or accelerate the formation of native TasA filaments (15) or fibrils (14). Here, we characterize this process on a molecular level for mature TapA and individual sections of its sequence, mainly using AUC and NMR, to understand whether TapA is merely a catalyst or becomes part of TasA filaments as proposed earlier (3, 10).

Functional units of the TapA sequence were analyzed by employing N- and C-terminal truncations. Secondary structure prediction by Jpred4 (23) and later AlphaFold (24) both indicated a folded domain between residues K72 and D189, including a disulfide bond (Cys92-Cys188) (Fig. 1*A*), and a large unstructured region beyond residue E190. The N terminus was considered flexible. Furthermore, different starting points of the mature protein [A33 (10, 17, 25) or A44 (15)] are reported in the literature, with

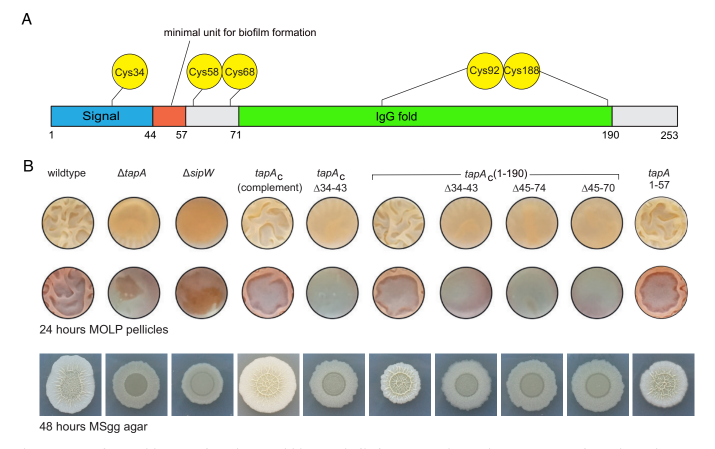


Fig. 1. Structure of TapA and the impact of N- and C-terminal deletions on biofilm formation. (*A*) Schematic domain composition of TapA. The signal sequence (AA 1-43) is shown in blue, the minimal unit (AA 44-57) for biofilm formation (15) in red, the folded domain in green, and the positions of the five cysteines C34, C58, C68, C92, and C188 are indicated. (*B*) Biofilm formation of TapA-modified *B. subtilis* strains grown on MOLP medium as pellicles (two top rows, the second row includes staining by Coomassie blue and Congo red) or on MSgg medium agar plates (bottom row). The strain modifications are indicated, see text for further explanations.

a large number of experiments performed on a TapA construct starting with A33, so that we also aimed to investigate more precisely the signal peptide cleavage site leading to mature TapA. The effects of the sequence alterations were examined by growing biofilms on MOLP medium as pellicles or on MSgg medium-containing agar plates (for the wild-type see Fig. 1B, column 1). Deletion of tapA or sipW both corrupt the biofilms (see Fig. 1B, columns 2 and 3). The use of *B. subtilis* $\Delta tasA$ operon strains complemented in *trans* with intact *tapA* in the *lacA* chromosomal locus and the sip W-tasA genes in the amyE locus (tapA complement, Fig. 1B, column 4) restores the wild-type biofilm phenotype, as well as the truncated *tap*A₁₋₁₉₀ (Fig. 1*B*, column 6). Deleting amino acid residues 34 to 43 in both wild-type and $tapA_{1-190}$ ($tapA_C \Delta 34-43$ and $tapA_{1-190}$ $\Delta 34-43$, Fig. 1B, columns 5 and 7) shows no biofilm formation, since signal peptide recognition, processing, and secretion could be affected by this deletion, indicating that the mature protein starts at A44. The deletions of amino acids 45 to 74 and 45 to 70 from the truncated protein TapA₁₋₁₉₀ (Fig. 1*B*, columns 8 and 9) also impairs the biofilm. Furthermore, a strain secreting only residues 44 to 57 of the N terminus after cleavage of the signal peptide ($tapA_{1-57}$, Fig. 1B column 10) showed proper biofilm formation as observed previously (14, 15). These experiments confirm A44 as the N terminus of mature TapA, and the crucial importance of N-terminal residues for biofilm formation. Deleting all residues beyond E190 did not change the appearance of the biofilms compared to wildtype TapA.

To determine conditions for in vitro TapA–TasA interaction experiments, the dependency on pH and/or salt concentration was investigated by isothermal titration calorimetry (ITC), interpreting the occurring heat changes solely as indicators of folding events combined with oligomerization. At pH 3 and 50 mM NaCl heat development was observed for the titration of TapA into TasA (*SI Appendix*, Fig. S2A), whereas the titration at pH 7 and 50 mM NaCl did not yield a response (*SI Appendix*, Fig. S2B). Increasing the salinity to 150 mM NaCl at pH 7.0 resulted again in a heat change (*SI Appendix*, Fig. S2C), demonstrating a considerable influence of salt concentration on TapA-dependent TasA oligomerization at neutral pH.

To ensure purity of the samples investigated below, the nature of the occurring TasA oligomers, potentially consisting of fibrils (ThT-stainable) and/or filaments (non-ThT-stainable) (15) were investigated in ThT-staining experiments (SI Appendix, Fig. S3) under very similar conditions. At pH 7, 20 °C, and 150 mM salt (buffer alone in first column), TasA forms nonstainable filaments (second column). The mixture of $TapA_{44\text{--}190}$ and TasA as well as the functional construct TapA₄₄₋₁₉₀ alone do not show relevant ThT responses either (columns 3 and 4, respectively). At low salt (50 mM), 40 °C, and neutral pH, small amounts of TasA progress toward stainable oligomeric states within 24 h (column 5). At low pH, intense fibril formation takes place within the first four hours (columns 6 and 7). TapA alone does not convert to stainable fibrils at low pH (column 8) and the C-terminally truncated variants of TapA show very similar effects to the full-length, mature protein (columns 9 and 10). In summary, fibrillar states of TasA are not observed under the conditions applied in our experiments at neutral pH but non-ThT-stainable filaments are formed.

TapA-mediated oligomerization of folded, monomeric TasA was then analyzed by sedimentation velocity AUC experiments at 20 °C, 150 mM NaCl, and pH 7 (*SI Appendix*, Table S1) with runs lasting 17 h each. TapA and TasA were first investigated separately, yielding the expected sedimentation coefficient distributions of monomers, with single peaks at an $S_{20,w}$ value of around 1.9 for TapA, and around 3.0 for ²H, ¹³C, ¹⁵N-labeled TasA (Fig. 2*A*) (with labeling, TasA occurs at a larger *s*-value; otherwise it would have occurred at around 2.1 resulting in overlap), and no oligomers were observed. A 1:1 mixture of both, however, revealed the presence of oligomers (Fig. 2A, orange curve) and the peak of monomeric TasA became vanishingly small. The TapA peak area was reduced to 84% +/- 1% as indicated by the comparison of the integrals of its green and orange monomer peaks, revealing a participation of TapA in oligomeric states. The three experiments shown in Fig. 2A were performed in the same run in parallel under the same conditions. The gray columns in SI Appendix, Fig. S3 are obtained after AUC and shaking up the samples. They indicate that amyloid fibers do not contribute to the detected oligomers. To detect an effect of the TasA:TapA ratio, we incubated 100 µM TasA (unlabeled) with 1, 2, 4 and finally 10 μ M TapA (100:1, 50:1, 25:1 and 10:1, respectively) (Fig. 2B). The now overlapping monomer peaks decreased as the oligomer peaks increased in a TapA concentration-dependent manner. Interestingly, the maximum of the high molecular weight species shifts toward smaller S_{20.w} values with increasing TapA concentration (Fig. 2 B, Right, enlarged region) indicating the dominant occurrence of smaller oligomers. In summary, AUC shows that a fraction of TapA becomes part of the oligomers formed, and with higher TapA: TasA ratios on average shorter filament stretches occur.

The relevance of different TapA sections for promoting oligomer formation was investigated by using truncated proteins. Deleting the N terminus resulted in TapA₇₅₋₁₉₀ that yielded only negligible amounts of oligomers in an AUC run with equimolar concentrations (Fig. 2C, turquoise curve; orange curve for comparison). By contrast, deletion of the C-terminal region 191 to 253 showed a very similar response (TapA₄₄₋₁₉₀ in Fig. 2D, orange curve) to mature TapA (red curve in Fig. 2B and orange curve in Fig. 2C). Due to the large excess of TapA in our AUC experiments in comparison to in vivo concentrations [reported molar ratio of 100:1 for TasA to TapA (13)] and the fast depletion of monomeric TasA, we observed relatively "small" oligomers. In Fig. 2 C and D, the $S_{20,w}$ values correspond to a distribution ranging from approximately dimers to octamers. Since a special role in TasA fibrillation was assigned to the N-terminal residues up to T57 of TapA by Earl et al. (15), we also investigated the effect of the adapted peptide C58SF61A-TapA₄₄₋₆₂ on TasA. The mutation C58S was chosen to prevent the occurrence of disulfide linked dipeptides, and F61A to reduce hydrophobicity. A run with 250 µM of peptide (2.5 times molar excess over TasA) yielded a weak response (Fig. 2D, red curve), the effect being comparable to less than $4 \mu M$ mature TapA (Fig. 2D, green curve). This experiment corroborates the crucial importance of the TapA N terminus but also hints at a contribution of the upstream section 63 to 189 at least in vitro, with the subsequent C terminus being dispensable.

A complementary view of the oligomerization process is provided by detecting the disappearance of NMR signals originating from monomeric forms of TapA and TasA. Such signals may vanish due to unfolding or the formation of very large assemblies, which are not observable by solution NMR methods due to broadening of the resonance lines typical for large molecular entities. We followed the TapA-induced intensity change of TasA signals in the ¹H chemical shift range between 0.5 and -0.4 ppm where folded proteins show characteristic, resolved signals (Fig. 2E, see arrows). Representative spectra of a 125 μM TasA/50 μM TapA solution recorded at different time points after mixing are shown in Fig. 2E, with the diagnostic TasA signal decreasing and the TapA signals remaining constant. Without TapA, the signals of TasA alone remain constant over days. A plot of the TasA signal decay at various TasA/TapA ratios, with monomeric TasA initially at 125 μ M, is shown in Fig. 2F. Surprisingly, the intensity of the

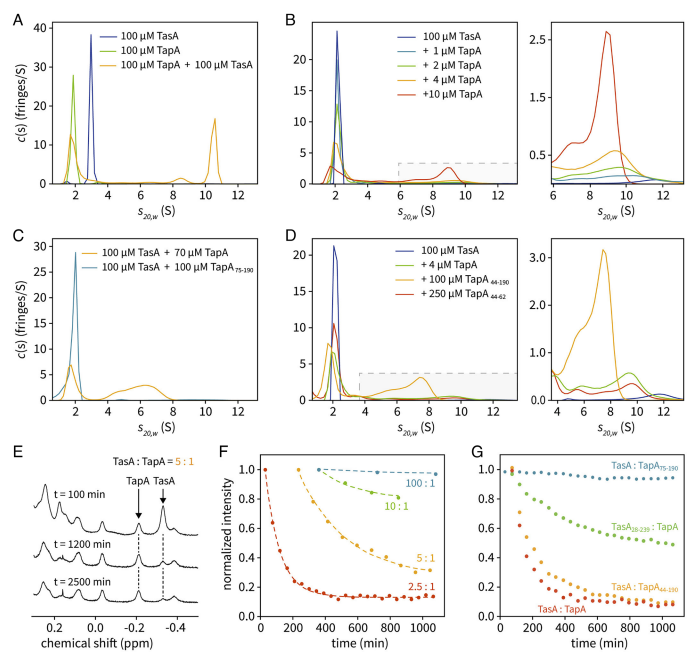


Fig. 2. Interaction of TapA and TasA at 20 °C, pH 7, 150 mM NaCl. (*A–D*) AUC experiments performed with TasA₂₈₋₂₆₁ and TapA₄₄₋₂₅₃, unless otherwise noted. (*A*) TapA and ²H, ¹³C, ¹⁵N-labeled TasA alone (green and blue peaks, respectively), and of a mixture of both consisting of 100 µM TapA and 100 µM TasA (orange). (*B*) 100 µM TasA was analyzed separately or in combination with 1, 2, 4, and 10 µM TapA. The gray-shaded area is shown enlarged to the right. (*C*) AUC of 100 µM TasA with 100 µM TapA₇₅₋₁₉₀ (turquoise) and of 100 µM TasA with 70 µM TapA for comparison (orange). (*D*) Evaluation of the effect of 250 µM C58SF61A-TapA₄₄₋₆₂ on 100 µM TasA (brown curve, see also enlargement to the right). Oligomer formation corresponds to less than 4 µM full-length TapA (green). The variant with deleted C terminus, TapA₄₄₋₁₉₀, shows wildtype behavior (orange). (*E*) ¹H NMR spectra of methyl group region from a 5:1 mixture of TapA remains nearly unaltered, the signal of TapA and TasA signals around –0.2 ppm and –0.35 ppm, respectively, were followed over time. While the signal of TapA remains nearly obtained on the spectra shown in (*E*). Signals are normalized to the first spectrum, time points refer to the time of mixing the solutions. (*G*) Time course of the disappearance of folded, monomeric TasA at different TasA:TapA ratios, with TasA at 125 µM.

TapA signals remained constant, no decay or change of line shape was observed in all recorded series (see indicated TapA signal in Fig. 2*E*). At a TasA concentration of 250 μ M the diagnostic TasA signal disappears within 10 h (Fig. 2*G* red curve) when 50 μ M TapA is added (5:1). This higher sensitivity is exploited to investigate the effect of N-terminal deletions in TapA and TasA. Adding TapA without residues beyond E190 (Fig. 2*G*, orange curve) yields a wild-type-like response (red curve) at the same concentrations. However, deleting the C terminus of TasA (residues 240 to 261) slows the decay of the diagnostic TasA signal considerably (green curve), indicating a stabilizing effect. In agreement with the AUC experiments, the addition of 50 μ M TapA₇₅₋₁₉₀ to 250 μ M TasA does not lead to a reduction of folded TasA (Fig. 2*G*, blue curve), supporting the notion that the N-terminal residues of TapA are critical for stimulating oligomerization.

An interesting picture emerges when looking at the AUC and solution NMR results together. At first sight, it seems surprising that according to AUC a fraction of TapA takes part in filament formation, but all of it remains as dynamic as a small protein since the line width of the TapA ¹H-NMR signals do not broaden during these long experiments, as would be expected in the case of dimer and multimer formation. From this, we conclude that TapA cannot be localized within a filament, which is expected to be rigid, but must be attached in a flexible manner at one or both ends. Furthermore, the influence of the flexible C terminus of TasA (Fig. 2*G*, green curve) is intriguing, since it hints at an importance of this segment in the resulting filament. This and the fact that folded TasA monomers are converted to filaments raises the question as to the thermodynamic stability of the proteins involved.

Monomer and Filament Stability. To identify driving forces for the oligomerization process, we determined the overall stability of the monomeric proteins and of the oligomers formed via thermal shift assays (TSA) on selected TapA and TasA constructs, and on mixtures preincubated for 17 h at 20 °C. The melting of mature TapA is characterized by a biphasic profile with one peak at 50 °C and another at 64 °C (SI Appendix, Fig. S4A). Most stable is TapA₄₄₋₁₉₀ with a $T_{\rm m}$ of 66 °C. In these cases, the TapA response is weak, whereas monomeric TasA shows a strong signal at a comparable concentration, and a melting point of approximately 43 °C (SI Appendix, Fig. S4B, blue curve). The solution of 100 µM TasA with 4 µM TapA (25:1, respectively) still predominantly contained monomeric TasA and TapA according to the AUC experiment (Fig. 2B), thus a melting point (43 °C) typical for monomeric TasA is observed (SI Appendix, Fig. S4*B*, orange curve). With 10 μ M TapA, a shift to appr. 56 °C (SI Appendix, Fig. S4B, red curve) was detected. This increase hints at a higher stability of TasA oligomers (and filaments) than of the monomeric form. The stability of TapA-induced filaments of TasA₂₈₋₂₃₉ is slightly lower, with a melting point of 52 °C (SI Appendix, Fig. S4C, orange curve). In summary, the process of TasA filament formation is thus driven by their higher stability, to which the C-terminal residues of TasA beyond K239 contribute significantly. The different melting points for monomeric TasA and the filaments may indicate structural rearrangements, whereas TapA has a much higher melting point than the filaments and is thus not expected to undergo large structural transitions involving its core domain.

Structural Investigations of TasA Filaments. Further characterization of TapA-induced TasA oligomerization requires structural insight. TasA filaments were prepared by the two different procedures, either concentrating TasA at pH 7 over a membrane or induced by TapA, then harvested by ultracentrifugation. Identical morphologies were observed by negative stain EM, with a width of approximately 3.5 nM and a recurring oscillation of ~25 nM (Fig. 3*A*). A comparison of the ${}^{1}\text{H}-{}^{15}\text{N}$ correlation spectra of both types of filament samples shows almost perfect overlap (SI Appendix, Fig. S5A). Recognizable effects due to TapA are absent, in agreement with the solution NMR results. Backbone assignments (87% coverage, *SI Appendix*, Table S2) were obtained from standard triple resonance experiments (26, 27). The most important assignments are indicated in *SI Appendix*, Fig. S5B, and examples of sequential correlations between residues are given by means of strip plots in SI Appendix, Fig. S6. For sections with peak doubling, we included both chemical shift sets in *SI Appendix*, Table S2.

The assignment provides information on the secondary structure by interpreting chemical shifts. Backbone angles derived with TALOS+ [(28), a dihedral angle prediction program] indicate the overall conservation of the b-sandwich structure of monomeric TasA_{G28-239} (*SI Appendix*, Fig. S7 and Fig. 3*B*). In particular, helices are predicted at the positions where they occur in the X-ray structure, as are most of the β -strands (Fig. 3*B*, blue and yellow, respectively). However, the N-terminal region (residues A28-S41) which shows high B-factors in the crystal structure of the monomer (8) exhibits typical β -strand chemical shifts (*SI Appendix*, Table S2) in the filament. Furthermore, the chemical shifts of β 2 are now typical for random coil arrangements (green in Fig. 3*B* and *SI Appendix*, Fig. S7).

Recording hNHH and hNhhNH experiments with through-space ¹H-¹H radio frequency-driven dipolar recoupling (RFDR) mixing (29) enabled us to observe cross peak patterns characteristic of hydrogen bonding in antiparallel *β*-sheets. Such pairs of hydrogen bonds lead to very strong cross peaks indicating interstrand H^N - H^N contacts due to the short distance of 3.3 Å (SI Appendix, Fig. S8A), much stronger than those observed for parallel β -sheets (30). We detected such interstrand H^N-H^N-contacts involving the originally flexible, N-terminal segment D31 to S41 pairing with A230 to N220 (SI Appendix, Fig. S8B). These interactions are unequivocally assigned since several cross peaks in the 2D ¹H-¹⁵N correlation are resolved with chemical shifts above 9 ppm (K35, A37, S41, I222, L224, A230, see SI Appendix, Fig. S5B where the hydrogen bonded residues are indicated in red, those in between in blue). Furthermore, the backbone signals of both strands are resolved in the 3D spectra used for resonance assignment (SI Appendix, Fig. S6), as are the correlations in the 3D hNHH spectrum (*SI Appendix*, Fig. S8*B*). The newly formed strand, termed β 0, is now aligned with β 9 instead of the segment N51-L57 that was interacting before as $\beta 2$ (Fig. 3 *C* and *D*). Thus, $\beta 1$ and $\beta 2$ of the monomer structure undergo structural changes and the loop between them becomes a strand, now β 1, that associates with β 3, as indicated by β -sheet-typical cross peaks between the H^{N} of K49 and Q71, indicative of hydrogen bonding (Fig. 3*D*). Given the observed hydrogen-bonding pattern between the newly arranged β 1 and β 3 and the overall conservation of most of the TasA structure, the H^N-H^N-pattern involving G42, N59, and L60 (SI Appendix, Fig. S8C) must be intermolecular since G42 and N59/L60 are then situated at distant ends of the β -sandwich. Directly preceding G42 is β 0 whose interactions with β 9 must thus also be intermolecular. To detect this directly, we prepared a sample with two TasA preparations mixed (50/50), one being ²H, ¹⁵N-labelled and ¹H back-exchanged, the other ¹³C-labelled by supplying [2-¹³C]-glycerol to the growth medium. The deuteration of the first sample reduces the excitation of its naturally abundant ¹³C nuclei when cross polarization (CP) from protons is applied, and decreases relaxation of its ¹H^N. This approach reduces the detection of intramolecular H^N-H^N contacts while intermolecular ones are promoted. We recorded 2D ¹H-¹⁵N projections of a hchhNH experiment that first filters protons attached to ¹³C followed by a proton mixing period via RFDR prior to the ¹⁵N-¹H correlation unit that selects protons attached to ¹⁵N. The resulting spectrum represents a selective subset of cross peaks indicative of intermolecular interactions (SI Appendix, Fig. S9A) that does not coincide with the largest cross peaks of the reference (SI Appendix, Figs. S5B and \$9B). A superposition of both shows a large number of signals in the β -sheet region with ${}^{1}H^{N}$ chemical shifts around 9 ppm (SI Appendix, Fig. S9C). A good coverage of the previously established intermolecular contacts is highlighted by the red labels in SI Appendix, Fig. S9D, except for G42 and L236, whereby the cross peaks associated with S41 and N220 are slightly shifted. Of course, more signals occur; however, it can safely be

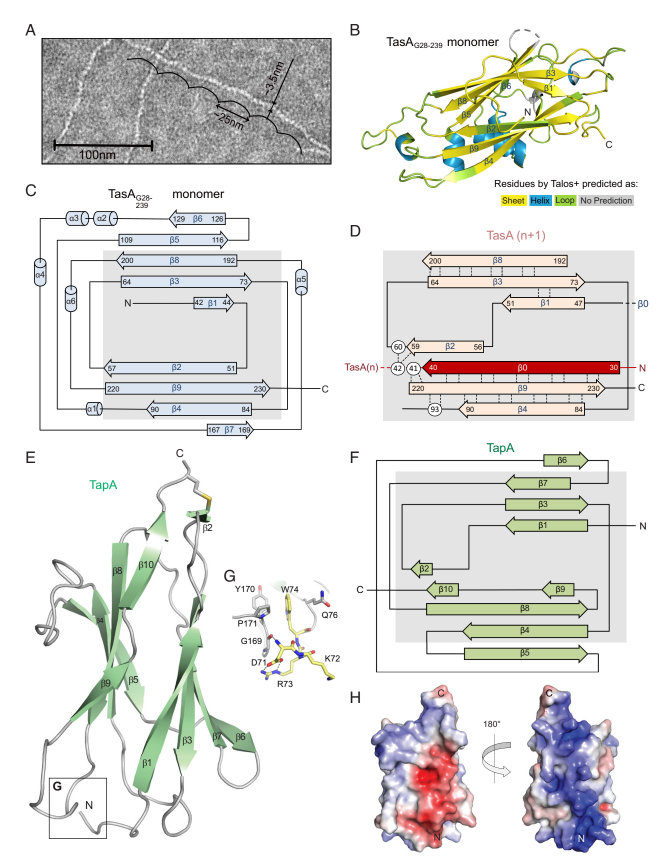


Fig. 3. Comparison of the TasA monomeric and filament secondary structure. (*A*) EM negative stain of TasA filaments. (*B*) TALOS+ predictions of filament secondary structure plotted on the TasA_{G28-239} monomer X-ray structure (PDB 5OF1). (*C*) Secondary structure topology diagram of the TasA monomer structure shown in *B*. The gray box highlights the core region of TasA discussed in the text. (*D*) Hydrogen-bonding pattern identified by solid-state NMR on the basis of H^N-H^N constraints within the TasA filament core region. (*E*) X-ray structure of TapA with disulfide bond (yellow) between C92 and C188. (*F*) Secondary structure topology plot of TapA₇₁₋₁₉₀. The N-terminal residues ₇₁DGRW₇₄ are shown in yellow and hydrogen bonds are indicated by dotted lines. (*H*) Electrostatic surface potential of TapA ranging from -5 kT/e (red) to +5 kT/e (blue) determined by the Pymol Plugin APBS Electrostatics tool. The left orientation is the same as in (*E*).

concluded that strand $\beta 0$ inserts between strands $\beta 9$ and $\beta 2$. This yields the secondary structure diagram shown in Fig. 3*D*, where the interstrand hydrogen bonds detected by hNHH and hNhhNH experiments are indicated by dotted lines.

An additional structural rearrangement involves the C terminus. The signals associated with the amino acids beyond Q232 adopt considerably different chemical shifts in the filament than in the monomer (8) as evident from an extreme H^N chemical shift of 3.18 ppm observed for H256. A further stabilization of the C terminus through hydrogen bonds is indicated by H^{N} - H^{N} cross peaks between S58 and V249 as well as between K65 and L236/ I238 (SI Appendix, Fig. S8C) that are intermolecular since S58 and K65 are very distant to the C-terminal residues in the monomer. Indeed, except for L236 all signals of the C-terminal residues are involved in large intermolecular cross peaks in the mixed sample (SI Appendix, Fig. S9 A and D for assignments). This finding explains the higher stability of filaments formed by mature TasA in comparison to those formed by the shorter Tas A_{28-239} . A summary of detected H^N-H^N contacts, including those observed involving signals of residues located in helices, are given in SI Appendix, Table S4.

Comparing the observed secondary structure and the style of strand insertion with examples in the literature, we found protomer connectivity and secondary structure in TasA filaments (Fig. 3D) to be similar to protein systems of the chaperone-usher pathway forming type 1 pili (31–34) on the surface of gram-negative bacteria. In fact, the unit defined by strands β 0-3, β 8, and β 9 (Fig. 3D) resembles in its secondary structure pattern that of the respective Fim and Pap family proteins in pili, with identical donor strand (β 0) insertion to complete a V-set Ig-like subunit [(35), Fig. 3 therein]. Still, other strand connections are different and reflect the original jellyroll fold of the TasA monomer structure.

TapA structure. The homology of protomer connectivity in TasA filaments with that in type 1 pili systems and our AUC results suggest a possible role of TapA as a chaperone-like facilitator of polymerization and raises interest in its structural properties, including both folded and unfolded areas. We therefore aimed to characterize all sections toward a comprehensive picture. The structure of TapA₄₄₋₁₉₀ was first investigated by NMR, omitting the functionally silent C terminus predicted to be flexible. No long-range NOE pattern was observed involving residues 44 to 71, but a structure of the following Ig domain was obtained (PDB: 6QAY). To characterize the N terminus further, the variants TapA₅₄₋₁₉₀ and TapA₆₀₋₁₉₀ were investigated by 2D ¹H-¹⁵N NMR spectroscopy and their spectra compared to the spectrum of TapA₄₄₋₁₉₀ (*SI Appendix*, Fig. S10) in order to identify its signals. A number of strong cross peaks with random coil chemical shifts are observed in the latter spectrum that are not covered by signals of the shorter forms, corroborating the notion of a disordered N terminus.

Structures of initially TapA₇₅₋₁₉₀ (1.28Å, PDB: 6HQC) and more recently TapA₇₁₋₁₉₀ (1.07Å, PDB: 8AIF) were determined by X-ray crystallography (*SI Appendix*, Table S3). The longer construct was investigated because NMR showed long-range contacts between W74 and the core of the Ig domain. TapA₇₁₋₁₉₀ shows a V-set Ig fold (InterPro code: IPR013106) composed of two antiparallel β -sheets (Fig. 3 *E* and *F*). D71 forms four hydrogen bonds involving residues G169, W74, and R73 (Fig. 3*G*). W74 and D71 together stabilize the loop between β -strands 8 and 9 (Fig. 3*G*). The additional hydrogen bonding suggests a higher stability of TapA₇₁₋₁₉₀ and 46 °C for TapA₇₅₋₁₉₀; see *SI Appendix*, Fig. S4*A*). Structural integrity is supported by a disulfide bond linking C92 and C188. The section connecting the N terminus with the Ig domain contains two further cysteines (C58 and C68) that form an additional disulfide bond according to gel filtration data (*SI Appendix*, Fig. S11). Constructs with odd numbers of cysteines such as TapA₆₀₋₁₉₀ and TapA₆₆₋₁₉₀ showed a tendency to form dimers via disulfide bond formation with their free cysteine C68. Those dimers disappeared upon DTT treatment. TapA₄₄₋₂₅₃, TapA₅₄₋₁₉₀, and TapA₇₅₋₁₉₀, on the other hand, appeared as monomers, since they contain even numbers of cysteines. However, as previously shown deletion of those cysteines has no functional effects on biofilm formation (14).

In summary, TapA consists of an Ig domain in its core (Fig. 3 *E* and *F*) and flexible N and C termini. However, the biphasic melting curve of mature TapA in contrast to the monophasic behavior of TapA₄₄₋₁₉₀ observed in the TSA (*SI Appendix*, Fig. S4*A*) suggests an association of the negatively charged TapA C terminus (191 to 253) with the Ig domain, likely interacting with the prominent, positively charged TapA surface patch (Fig. 3*H*). Intriguingly, the secondary structure pattern of the Ig domain resembles in its core (gray background) that of the TasA filament structure (Fig. 3*D*) and the proteins of the chaperone-usher pathway (32). Fitting to this picture, the first 5 N-terminal residues of TapA,AFHDI, are nearly identical with the TasA N terminus (AFNDI). This gives TapA the potential to take part in the donor-strand mechanism toward the formation of the filaments.

Discussion

Overall, the presented results suggest a mechanism of TapAinduced TasA filament formation in B. subtilis biofilms that resembles in its essence type 1 pili assembly in gram-negative bacteria, with TapA adopting a chaperone-like role reminiscent of the chaperone-usher pathway. AUC enabled us to demonstrate TapA to be necessary for filament formation but included in only small amounts (Fig. 2A). In a seemingly opposite way, solution NMR revealed fast, isotropic motions of the filament-attached TapA Ig domain (Fig. 2*E*) by the unchanged line shape of its narrow NMR signals after participating in oligomerization. Together with the observation of rigid, helical filaments by EM (Fig. 3A) this leads to the conclusion that mobile TapA can only be located at the beginning or end of a filament. Based on the near-complete (87%) assignment of the TasA filament backbone signals (SI Appendix, Figs. S5 and S6) obtained by solid-state NMR methods, the secondary structure of the filaments could be traced (SI Appendix, Fig. S7 and Fig. 3D), including the identification of intermolecular contacts (SI Appendix, Figs. S8 and S9). This revealed donorstrand insertion as the prime mechanism for oligomerization, with each TasA molecule donating its N terminus to become part of an Ig-similar core structure together with the second molecule (Fig. 3D). This process is driven by the higher stability of the filaments, as indicated by a melting point of 56 °C in comparison to 43 °C for monomeric TasA (SI Appendix, Fig. S4B). The X-ray structure of TapA in conjunction with NMR investigations revealed the presence of a complete Ig domain in the center of the TapA sequence and flexible N and C termini. The high melting point of TapA (64 °C) (SI Appendix, Fig. S4A), together with the fact that it already consists of a complete Ig fold, explains its inability to be located within a filament. Knowing the protein's signal peptide cleavage site (Fig. 1) and the observed homology of N-terminal residues between TapA and TasA enables us to propose the mechanism of TapA-induced TasA filament growth, with TapA as an initiator (see below).

In detail, our AUC and NMR experiments demonstrate an acceleration of TasA filament formation by TapA in a concentrationdependent manner (Fig. 2). AUC experiments with similar concentrations of TapA (100 or 70 $\mu M)$ and folded TasA (100 $\mu M,$ Fig. 2 C and D) show multimers that range from approximately dimers up to octamers. At such high TapA concentrations, a high number of oligomers are initiated so that they remain small due to the limited TasA supply. TapA is incorporated into the oligomers generated, as demonstrated by the decrease of the TapA monomer peak in the AUC experiment shown in Fig. 2A, and the concentration-dependency of the oligomer distribution (Fig. 2B) at constant TasA concentration. However, in our time-dependent 1D NMR spectra of samples prepared with a TapA to TasA molar ratio of initially 1:5, TapA signals remain at the same intensity and do not show an increase of line width. Our AUC and NMR data also provide insight into the importance of the TasA C termini for filament formation and stability. The reduced NMR signal decay of TasA₂₈₋₂₃₉ (SI Appendix, Fig. S5A) indicates slower oligomer formation and is paralleled by a decreased melting point of the resulting filaments (52 °C) in comparison to those resulting from wildtype protein (56 °C) (SI Appendix, Fig. S4 B and C). This supports that the stability of the filament affects the kinetics of oligomerization.

Our solid-state NMR data obtained on filament samples allow for distinction of intra- and intermolecular interactions due to the conservation of most of the β -sandwich monomer structure and enable a comparison with an AlphaFold Multimer (24, 36) model of TasA filaments (Fig. 4A). The pitch and diameter as observed by EM (Fig. 3A) is reproduced by the AlphaFold model, which is in good agreement with the recently published cryo-EM structure (18). Our NMR restraints confirm these models (SI Appendix, Figs. S7 and S8), especially the detected hydrogen-bonding patterns (SI Appendix, Table S4) but also the chemical shift data that are in agreement with the secondary structure of the protomers (SI Appendix, Fig. S12). Surprisingly, nearly all of TasA in filaments is ordered and rigid according to the successful CP, whereas only 70+ residues contribute signals in CP spectra when fibrils are investigated (10). The Alphafold model and the EM structure diverge strongly in the region 117 to 126 for which we do not observe NMR signals, presumably due to high flexibility. Similarly, the spatially adjacent residues K176 and T177 discussed to be important for filament bundling (18) are likely flexible, however, G175 shows a ${}^{1}H^{N}$ chemical of 9.6 ppm, which usually indicates an ordered structure. The signals of G175 are at the same position as in solution NMR spectra of the monomeric form (8), indicating a very similar chemical environment.

The difference between the protomer structure and our monomer X-ray structure of TasA_{G28-239} (Fig. 4B) is in β 1 reversing its direction, and $\beta 2$ now being shorter and parallel to $\beta 0,$ see also Fig. 3D. In accordance with the strong H^{N} - H^{N} correlations shown in SI Appendix, Fig. S8, the N terminus of the preceding protomer n runs antiparallel to $\beta 9$ of molecule n+1, $\beta 9^{n+1}$ (Fig. 4 C and D). The model reproduces the detected restraints involving the C-terminal residues well, see orange residues in Fig. 4E. It shows hydrogen-bonding interactions between residues 236 to 238 of protomer n and residues 64 to 66 of $\beta 3^{n+1}$ (Fig. 4 *C*, Right and SI Appendix, Fig. S8C) that manifest a short intermolecular parallel β -sheet, and a cation- π interaction between the W233 six-membered ring and H256 (37) that explains the extreme H^N chemical shift of the latter. Furthermore, the model shows a loop-like pattern for residues 250 to 252 and contacts between S58 of protomer n+1 and V249 of n (Fig. 4 C, Left). Altogether, these data indicate high reliability of the model

concerning the TasA-TasA interface area. This is corroborated by the spectrum indicating intermolecular contacts in SI Appendix, Fig. S9. We have indicated example positions of 2D cross peaks expected from the evaluation of the AlphaFold model in SI Appendix, Fig. S9A, with good agreement. In its core, the interface is made up of residues from three protomers. $\beta 0^n$ spans two protomers, traversing molecule n+1 and touching n+2 via the aromatic ring of F29ⁿ (Fig. 4 D and E). By binding $\beta 0^n$, a protomer n+1 is able to present an oligomerization-competent TasA-TasA interface to protomer n+2. In this context, F29ⁿ, $Q232^{n+1}$ and $F200^{n+2}$ form a perfect binding site for P62ⁿ⁺², a "proline box" as known from the analysis of domains that bind proline-rich peptides (38) (Fig. 4 D and E). There, Q232 is involved in three hydrogen bonds with the P62 and F29 car-bonyls, and F29 H^N. Indeed, a ¹³C-¹³C correlation spectrum of a ¹³C, ¹⁵N-labeled sample produced with [2-¹³C]-glycerol (39) shows a separated signal for the CS of P62 (SI Appendix, Fig. S13A), and cross peaks involving the Ca of A28 and F29 that indicate short distances (SI Appendix, Fig. S13B). Strikingly, this part of the protomer interface, the proline box, is the most conserved in TasA and camelysin alignments (see residues highlighted as most conserved in SI Appendix, Fig. S1), even though the identity of the compared sequences may be as low as 26%. The side chains of residues with highest conservation, together with a comparison of secondary structure, are shown as spheres in SI Appendix, Fig. S14, all appearing at the interface between protomers. As is common for β -sheet structures, strand $\beta 0$ is not strictly conserved except for its beginning. The high conservation of the TasA-TasA contact area potentially enables the formation of joint filaments with TasA homologs from other species. This is in line with the observation that *B. cereus* is able to stimulate TasA production in B. subtilis (21).

With the data presented, a possible mechanism for B. subtilis biofilm formation emerges. According to earlier reports (2, 6, 13, 15) TapA is associated with the membrane. Analogous to the chaperoning of pili proteins (31-34), TasA molecules arrive unfolded through SecYEG and may start forming filaments with the help of the presented, homologous (Fig. 4F) TapA N terminus, leading to an initial heterodimer, then binding the next TasA molecule to become a trimer (Fig. 4G). The N-terminal residues of TapA (Fig. 4F) bind in a very similar way as TasA to protomerⁿ⁺¹, whereby H46 of TapA in position of N30 in TasA serves in the same manner to establish hydrogen bonds to the TasA residues E229ⁿ⁺¹ and N215ⁿ⁺² (Fig. 4D). Since folded, monomeric TasA supports biofilm formation when supplied to $\Delta tasA^{-}$ B. subtilis cultures (8), a scenario is required that explains the mechanism of its inclusion into the growing chain, driven by the higher thermodynamic stability of the filament (SI Appendix, Fig. S3B). Two pathways may be envisaged: unfolding of TasA or rearrangement of β 1-2. The first option is supported by our observation that in filaments made from deuterated protein all ${}^{2}H^{N}$ were always replaced by ¹H^N after oligomerization, especially in the core of the protein (see assignment table in SI Appendix, Table S2). TapA thus functions as a chaperone in analogy to the chaperone-usher pathway since it already has a complete and especially stable (SI Appendix, Fig. S4A) V-set Ig fold and has the ability to complement TasA in the process of filament formation by donating its previously unstructured N-terminal strand. As a result, structurally similar TapA and TasA filament core units are observed (SI Appendix, Fig. S14 B-D). Owing to the stability of the TapA Ig domain, it is not able to open a cleft to receive a donor strand, and thus can only be located at the beginning of a filament, acting as filament inducer.

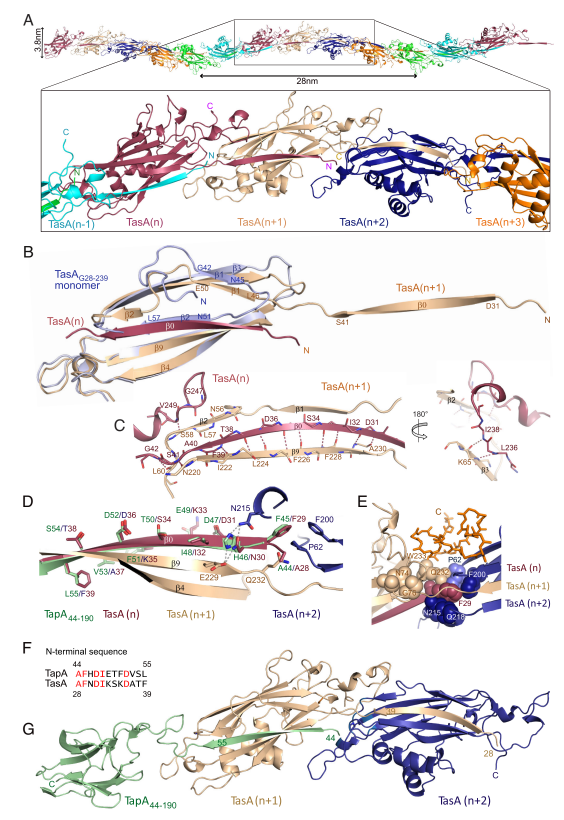


Fig. 4. TapA–TasA interactions. (*A*) TasA filament AlphaFold model with close-up view onto five consecutive protomers. (*B*) Superimposition of the monomer X-ray structure (light blue) obtained by crystallization of TasA_{G28-239} with a protomer structure of the filament [beige (n+1)]. The N terminus of the preceding TasA molecule (n) is shown in red. (*C*) Hydrogen-bonding pattern identified by solid-state NMR confirming the β 0-strand interaction. (*D*) TapA N terminus (green) superimposed onto the TasA β 0 strand (red) of the TasA filament (*A*). Binding site residues F29, Q232, and F200 (TasA counting) form an ideal cavity for P62, all of them belonging to the most conserved residues. Side chain residues are presented as sticks and hydrogen bonds as gray dashed lines. (*E*) Cose-up view on the TasA–TasA interaction site composed of the most conserved residues (shown as spheres), highlighting the binding pocket of TasA (n) F29 (red) surrounded by residues of TasA (n+1) and TasA (n+2) shown in beige and blue, respectively. (*F*) Sequence alignment of the N-terminal TapA and TasA residues. Identical residues are depicted in red. (*G*) AlphaFold model of TapA (green)–TasA (beige and blue) interaction.

Materials and Methods

Constructs for Protein Expression. Primers for cloning of different TapA constructs are summarized in *SI Appendix*, Table S5. Chromosomal DNA of *B. subtilis* 168 and primer pair 1 were used to amplify the gene for TapA₄₄₋₂₅₃ (Uniprot-AOA6M4JJ23) by PCR. The PCR product was cut by XhoI and BsmI and cloned into pCA528 (40) (a modified pET24a vector with Kanamycin resistance) to generate pCA528_His_ Sumo_TapA₄₄₋₂₅₃. Further variants of TapA were generated by a modified QuikChange protocol for site-directed mutagenesis. For the C-terminally truncated version pCA528_His_Sumo_TapA₄₄₋₁₉₀, a stop codon was introduced at position 191 with primer pair 2. Primer pairs 3 to 7 were used to generate additional N-terminally truncated constructs pCA528_His_Sumo_TapA_{54;60;66;71 and 75-190}, respectively. TasA constructs were prepared as previously described in ref. (8).

Protein Expression and Purification. Methods published previously for TasA (8) were applied to TapA.

Prediction Methods. Jpred4 was used to predict secondary structure (http:// www.compbio.dundee.ac.uk/jpred4) (23). The probability for disulfide bridges was assessed by Disulfind (41). The ProteinDataBase (PDB, https://www.rcsb.org/) was screened for structures of proteins with similar amino acid sequence. Threedimensional structure predictions based on the AlphaFold (24) algorithm were run via the publicly available ColabFold (https://github.com/sokrypton/ColabFold) (42) infrastructure through Google Colaboratory. For TasA oligomer structure predictions, the multiple sequence alignment was conducted with MMseqs2 (UniRef+Environmental) and model_type was set to AlphaFold2-multimer. Default settings were used for all remaining options.

Strain Construction for Biofilm Experiments. All primers and plasmids used are listed in the tables (*SIAppendix*, Tables S6 and S7). For the complementation of the P_{tapA}-tapA Δ 13-234-*sipW*-tasA-kan in the *amyE* locus, the region was amplified by PCR from the genomic DNA of *u* 3610 coml^{O12L} and cloned into the pBS1K plasmid by restriction digest with Xbal and Spel, subsequent treatment with alkaline phosphatase and ligation. For the complementation of the different *tapA* constructs in trans in the *lacA* locus (plasmids pKXD2 and pKXD7), the full-length *tapA* as well as the truncated *tapA*₁₋₁₉₀ were amplified by PCR and cloned into pBS2E by restriction digest using Xmal and Pael. The *tapA* constructs containing different deletions were constructed in a 2-step PCR and subsequently digested using Xmal and Pael and cloned into pBS2E. The truncated TapA₁₋₅₇ was constructed from pKXD7 using PCR amplification and KLD enzyme mix from NEB for circularization.

For construction of strain BKD30, the plasmid pKXD13 was transformed into BRK49. In short, the receptor strain was grown in competence medium (17.5 g/L K₂HPO₄, 7.5 g/L KH₂PO₄, 2.5 g/L (NH₄)₂SO₄, 1.25 g/L tri-Sodium citrate x2 H₂O, 0.5% glucose, 7 mM MgSO₄, 0.1 mg/mL L-tryptophan, 0.02% casamino acids, 0.22 g/mL ammonium iron citrate) to OD₆₀₀ 0.6 to 0.8, starvation medium (17.5 g/L K₂HPO₄, 7.5 g/L KH₂PO₄, 2.5 g/L (NH₄)₂SO₄, 1.25 g/L tri-sodium citrate x2 H₂O, 0.5% glucose, 7 mM MgSO₄, 0.25 g/L (NH₄)₂SO₄, 1.25 g/L tri-sodium citrate x2 H₂O, 0.5% glucose, 7 mM MgSO₄, 0.1 mg/mL L-tryptophan, 0.02% casamino acids, 0.22 g/mL ammonium iron citrate) to OD₆₀₀ 0.6 to 0.8, starvation medium (17.5 g/L K₂HPO₄, 7.5 g/L KH₂PO₄, 2.5 g/L (NH₄)₂SO₄, 1.25 g/L tri-sodium citrate x2 H₂O, 0.5% glucose, 7 mM MgSO₄) was added for 1 h and then 1 mL competent culture was incubated with the DNA to be transformed for 2 h. Positive clones were selected on LB agar containing the antibiotics required and the integration verified by PCR and DNA sequencing. The different *tapA* constructs were subsequently transformed in BKD30 as described above.

Biofilm Experiments. Overnight cultures were prepared in LB-medium containing the required antibiotics (spectinomycin 150 µg/mL, kanamycin 10 µg/mL, erythromycin 1 µg/mL and lincomycin 25 µg/mL). The overnight cultures were diluted 1:100 in 10 mLLB-medium without antibiotics in 100 mL Erlenmeyer flask and grown at 37 °C, 180 rpm until OD₆₀₀ reached 0.5. The cultures were diluted as described above in fresh LB-medium and grown again to OD₆₀₀ 0.5. In a 24-well plate, 1.4 mL of medium optimal for lipopeptide production (MOLP) (30 g/L peptone, 20 g/L saccharose, 7 g/L yeast extract, 1.9 g/L KH₂PO₄, 0.001 mg/L CuSO₄, 0.005 mg/L, FeCl₃ × 6 H₂O, 0.004 mg/L, Na₂MoO₄, 0.002 mg/L KI, 3.6 mg/L, MnSO₄ × H2O, 0.45 g/L MgSO₄, 0.14 mg/L, ZnSO₄ × 7 H₂O, 0.01 mg/L H₃BO₃, and 10 mg/L C₆H₈O₇) (43) was inoculated with 14 µL culture and optionally 14 µL biofilm dye (final concentration 0.02 mg/mL Congo Red, 0.01 mg/mL Coomassie Brilliant Blue G250) (2) was added.

For biofilms on minimal salts glutamate glycerol (MSgg) agar plates, 10 μ L culture were spotted on MSgg medium [5 mM potassium phosphate (pH 7), 100 mM Mops (pH 7), 2 mM MgCl₂, 700 μ M CaCl₂, 50 μ M MnCl₂, 50 μ M FeCl₃,

1 μ M ZnCl₂, 2 μ M thiamine, 0.5% glycerol, 0.5% glutamate, 50 μ g/mL tryptophan, 50 μ g/mL phenylalanine] (44) solidified with 1.5% agar.

The Biofilms shown were incubated at 30 °C 24 h for MOLP and 48 h for MSgg medium.

TSA. TSA was done in 50 μ L setups in 96-well PCR plates (Biorad MLP9601) covered with a nonfluorescent sealing foil (Biorad INN120411) in a real-time PCR machine IQ5 (BioRAD) with a nonstandard filter combination (excitation at 438 nm, emission at 495 nm) to measure SYPRO Orange (5,000-fold stock from Invitrogen) fluorescence. A mixture of onefold SYPRO orange and protein in 20 mM phosphate buffer pH 7.0, 50 mM NaCl was heated 1 degree per minute between 25 and 90 °C. With unfolding the fluorescence increases and the negative first derivative shows the melting point as minimum (IQ5 software). The protein amount had to be optimized in advance; typically, the range from 5 to 50 μ M was checked, depending on the size and structure of the protein. For comparable melting curves, 10 μ M TasA₂₈₋₂₆₁, but 50 μ M TapA₄₄₋₂₅₁ was needed.

ThT Assay (ThT assay). If not otherwise mentioned 100 μ L 20 μ M protein solution and 100 μ L 40 μ M ThT, both in 20 mM phosphate buffer pH 7.0, 50 mM NaCl, were mixed in a black 96-well plate with clear bottom (Corning 3615), and fluorescence was monitored using the bottom mode (excitation at 438 nm emission at 495 nm; 5 nm band width, gain 80) in a Safire microplate reader (Tecan).

Solution NMR. All NMR spectra were recorded on Bruker AV-III spectrometer operating at 600 and 750 MHz ('H frequency) at a temperature of 27 °C. In both spectrometers a "TCI"-type cryoprobe equipped with one-axis, shielded gradients was used. The spectrometers were operated using TopSpin 3.5. For the determination of the structure of TapA_{\rm 44-190} a sample labeled with ^{13}C and ^{15}N was used, the concentration was 540 μ M in 20 mM phosphate buffer pH 7.0, 50 mM NaCl. An extensive set of experiments was recorded [¹⁵N-HSQC, ¹³C-HSQC, HNCA, HNCACB, HN(CO)CACB, HNCO, HN(CA)CO, HBHA(CO)NH, H(CCO)NH, CC(CO)NH, HCCH-COSY, HCCH-TOCSY] resulting in the assignment of almost all resonances of aa 75 to 190. Distance information was extracted for that region from 3D¹⁵Nand ¹³C-NOESY spectra. In all spectra, water-suppression was accomplished using a Watergate scheme. Data were processed using topspin and analyzed using the program from the Collaborative Computing Project for the NMR community (CCPN), CcpNmr Analysis 2.4.2 (45). The structures were calculated using ARIA (46). For the analysis of the dynamics of the protein ¹⁵N-T1, T2, and hetNOE spectra were recorded using a ¹⁵N-labeled sample of a concentration of 1 mM, again using Watergate for water suppression. The assignments are deposited in the Biological Magnetic Resonance Data Bank (BMRB) (entry 34341), the structure can be found in the PDB (code 6QAY). To check samples and to compare protein constructs of different length, ¹⁵N-HSQC spectra were recorded using Waterqate water-suppression. Experiments to follow the effect of TapA on TasA were recorded as one-dimensional spectra using Watergate with excitation sculpting. To follow the development over a longer time period, pseudo two-dimensional spectra were recorded. The data were processed using topspin 3.5, and decay curves were extracted using in-house scripts.

Filament Formation. Monomeric TasA separated by gel filtration in 20 mM phosphate buffer pH 7.0 with 150 mM NaCl was slowly concentrated over 2 to 3 d in an Amicon stirring device equipped with a 10 kDa regenerated cellulose membrane at 10 °C. As a result, a turbid solution was obtained. Prior to sample transfer into the NMR rotor, the filaments were sedimented by ultracentrifugation for 1 h at 8 °C and 130,000 g, using a TLA110 fixed angle rotor.

Alternatively, growth of TasA filaments (e.g., 250 μ M) was achieved in the presence of TapA (e.g., 50 μ M) overnight at 20 °C in 20 mM phosphate buffer, pH 7.0 with 150 mM NaCl. The deuterated sample used for establishing resonance assignments was prepared as described below under "Analytical and Preparative Ultracentrifugation", last paragraph.

Solid-State NMR. Assignment spectra were recorded on a Bruker Avance III spectrometer operating at 600 MHz ¹H Larmor frequency. For studies on TapA-induced filamentous TasA, fully ²H, ¹³C, ¹⁵N labeled and 100% ¹H back-exchanged protein in 20 mM phosphate buffer, pH 7.0, and 150 mM NaCl, washed with the same buffer but 50 mM NaCl, was spun at a 60 kHz magic angle spinning (MAS) rate using a 1.3 mm rotor. Temperature control was achieved with a BCU II, a gas flow of 1,200 L/h, and a cooling gas temperature of 252 K. For the assignment spectra, manufacturer

provided pulse programs hNH2D.dcp, hCaNH3D.tcp, hCONH3D tcp, hcaCbcaNH3D. tcp, hCOcaNH3D.tcp, hcaCbcacoNH3D.tcp, and hcoCacoNH3D.tcp with CP (47) for heteronuclear magnetization transfer and a J-coupling based scheme for homonuclear (^{13}C - ^{13}C) transfer (48) were employed (26). Parameter recommendations provided in the pulse programs were closely followed and subsequently optimized for existing conditions. Spatial information was extracted from in-house programmed hNHH and hNhNH pulse sequences. The hNHH consists of an hNH CP step yielding the two indirect dimensions followed by a 1.5 ms ^{1}H - ^{1}H RFDR step (29) prior to acquisition. The hNhNH pulse includes an additional hNH CP step.

For the sample with mixed labeling pattern, we prepared TasA monomer solutions that were either ²H, ¹⁵N labeled or ¹³C labeled by supplying solely [2-¹³C]-glycerol as carbon source during expression. These solutions were mixed 50/50 and filaments created by concentrating the sample in aqueous buffer with 20 mM phosphate buffer, pH 7.0, and 150 mM NaCl. Filaments were washed with the same buffer, however, containing only 50 mM NaCl, and transferred into a 1.3 mm rotor. For the measurements, NMR spectrometer and settings were as above. A ¹H-¹⁵N spectrum (hNH2D.dcp) for comparison was acquired optimizing the previously determined parameters. A hchhNH pulse sequence was used to monitor intermolecular contacts. It consisted of two CP blocks (hch and hNH, respectively) in between which a 3 ms RFDR mixing period was introduced (180 rotor cycles at 60 kHz MAS, depiction in *SI Appendix*, Fig. S9*E*).

For ¹³C - ¹³C correlation measurements by dipolar-assisted rotational resonance (DARR) either uniformly ¹³C, ¹⁵N- or sparsely [2 - ¹³C]-glycerol, ¹⁵N-labeled TasA was purified and subjected to filament formation through concentration as mentioned previously. TasA filaments were filled into a 3.2 mm rotor and measured at a Bruker Avance III spectrometer operating at 800 MHz ¹H Larmor frequency. For uniformly labeled TasA, a short mixing time (20 ms) was used and for the sparsely labeled protein a long one (400 ms).

Data acquisition and processing was conducted using TopSpin 3.5pl7 (Bruker). Peak picking and protein assignment was conducted with CcpNmr Analysis 2.4.2 (45).

Crystallization and Structure Determination. Tap $A_{75\cdot190}$ and Tap $A_{71\cdot190}$ in 20 mM Tris buffer pH 7.0 50 mM NaCl were crystallized using the sitting-drop vapor-diffusion method. Crystallization setups were performed by using a Gryphon pipetting robot (Matrix Technologies Co.) for pipetting 200 nL protein with a concentration of 23 to 26 mg/mL to an equal volume of precipitant solution. The Rock Imager 1000 storage system (Formulatrix) was used for storing and imaging of the experiments. Crystals appeared within 1 to 3 d for both TapA variants and were flash-frozen in liquid nitrogen in the presence of 20% ethylene glycol. For crystallographic phase determination, SeMet-incorporated TapA75-190 was crystallized in a crystallization condition containing 20% MPD, 0.1 M Bicine pH 9.0, whereas native TapA75-190 was crystallized in a 20% PEGMME 550, 0.1 M NaCl, 0.1 M Bicine pH 9.5 condition. The N-terminally elongated TapA_{71.190} yielded crystals in 24% PEG3350, 0.35 M LiNO₃ All diffraction data were recorded on BL14.1 at BESSY II (Helmholtz-Zentrum Berlin, HZB), processed and scaled using XDSapp (49). The crystallographic phase problem for the SeMet derivative of TapA₇₅₋₁₉₀ (1.3 Å data, anomalous signal up to 1.6 Å) was solved by using the automatic crystal structure determination platform Auto-Rickshaw (50). The structure of native TapA_{75.190} (1.28 A) was solved by molecular replacement with Phaser (51) using the SeMet-TapA₇₅₋₁₉₀ Auto-Rickshaw structure as the search model. The TapA₇₁₋₁₉₀ structure was also solved by molecular replacement with Phaser, but here the TapA75-190 structure was used as the search model. Finally, both protein models were manually built using COOT(52) and iteratively refined using Refmac (53).

All residues of the TapA₇₅₋₁₉₀ structure are nicely explained in the electron density. Crystals of TapA₇₁₋₁₉₀ contain 3 molecules (chains A-C) per asymmetric unit. Chain A covers residues 72 to 189, chain C residues 71 to 188, and chain B shows the complete molecule. The missing residues in chain A and C were disordered and therefore not visible in the electron density.

All structure statistics is given in *SI Appendix*, Table S3. 95.6% of the residues in the TapA_{75.190} structure were in the allowed regions of the Ramachandran map and 96.6% for the TapA_{71.190} structure. The Ramachandran statistics of both structures were analyzed using Molprobity (54). Figures and structure superimpositions were generated with PyMol (http://www.pymol.org). The atomic coordinates of TapA_{75.190} and TapA_{71.190} have been deposited in the Protein Data Bank (PDB ID codes 6HQC and 8AIF, respectively).

Analytical and Preparative Ultracentrifugation. Sedimentation velocity experiments were performed with a Beckman Optima XL-I analytical ultracentrifuge at 20 °C and a rotor speed of 40,000 rpm (129,000 g at cell bottom). The samples indicated in *SI Appendix*, Table S1 were measured in dialysis buffer (20 mM Na phosphate pH 7.0, 150 mM NaCl) separately or in combination at a final concentration of 100 μ M unless otherwise stated. In titration experiments, the TapA concentration was reduced to 10, 4, 2 and 1 μ M. Interference data with a total of 200 scans were recorded every 5 min. Sedimentation coefficient distributions c(s) were analyzed with the program Sedfit (55). The protein partial-specific volume and the buffer physical constants were calculated from amino acid and buffer composition, respectively, using SEDNTERP (56). Figures were created with GUSSI (57). After the run the samples have been recovered, mixed, and applied to ThT assay and TSA as described there.

To obtain samples for NMR and ThT assays (Fig. 24), the mixtures of 1.5 mL of 100 μ M ²H, ¹³C, ¹⁵N-TasA, and 1.5 mL of 100 μ M unlabeled TapA were ultracentrifuged (Beckman Optima Max) under the same g-force as mentioned above overnight. The resulting pellet was washed once with 20 mM Na phosphate pH 7.0, 50 mM NaCl and filled into a 1.3 mm MAS rotor for NMR. A small amount of sample was analyzed by negative stain EM.

ITC. ITC experiments were performed using a PEAQ-ITC microcalorimeter (Malvern). All titrations were performed at 18 °C with 200 μ M TapA variants (500 μ M for TapA peptides) in the syringe and 20 μ M TasA in the reaction chamber. The protein as well as the titration components were dissolved in buffer containing 20 mM Tris/HCl and 50 mM or 150 mM NaCl, pH 7.0. Malvern software was used to visualize the binding process.

Sequence Alignment. Additional protein sequences homologous to TasA were located by supplying diverse sequences annotated as camelysines to the program blast; evolutionary relevant sequences were chosen from the result. Alignment was then performed with MEGA11 (58) and the ClustalW algorithm (59) with default settings. The resulting alignment was then exported to .fasta format and visualized using the strap (60) web application (https://www.bioinformatics.org/strap/aa/).

Data, Materials, and Software Availability. TapA structures are available in the PDB under accession codes 6QAY (61) for NMR, and 6HQC (62) (TapA₇₅₋₁₉₀) and 8AIF (TapA₇₁₋₁₉₀) for the X-ray structures. Solution NMR chemical shifts of TapA are deposited in the BMRB under entry 34341 (63). Solid-state NMR raw data, chemical shift and peak lists of TasA filaments are deposited in the BMRB as entry 51785 (64). Processed TasA filament spectra and the CcpNmr Analysis 2.4.2 (45) assignment project are uploaded to zenodo (10.5281/zenodo.7534572) (65). All study data are included in the article and/or *SI Appendix*.

ACKNOWLEDGMENTS. We thank Regina Alver for initial cloning of TapA and TasA and construction of *Bacillus* strains. Further strains were provided by Åkos Kovács (Technical University of Denmark, Copenhagen). Thanks to Natalja Erdmann, Daniel Friedrich, and Matthias Herrera-Glomm (all Leibniz-Forschungsinstitut für Molekulare Pharmakologie, FMP) for excellent technical assistance, and to Ines Kretzschmar (FMP) for the synthesis of peptides. We acknowledge Daniel Roderer for initial electron microscopy investigations. We are grateful to Barth van Rossum for providing figures. H.O. acknowledges funding by the German Research Foundation (Deutsche Forschungsgemeinschaft, DFG), grant OS 106/17-1. K.T. was supported by the DFG (Tu106/6 & SPP1879) and the Max Planck Society. We acknowledge access to beamlines of the BESSY II storage ring via the Joint Berlin MX-Laboratory sponsored by the Helmholtz Zentrum Berlin für Materialien und Energie, the Freie Universität Berlin, the Humboldt-Universität zu Berlin, the Max-Delbrück-Centrum, the Leibniz-Institut für Molekulare Pharmakologie, and Charité–Universitätsmedizin Berlin.

Author affiliations: ^aStructural Biology, Max Delbrück Center for Molecular Medicine, 13125 Berlin, Germany; ^bLeibniz-Forschungsinstitut für Molekulare Pharmakologie, 13125 Berlin, Germany; ^bDepartment for Molecular and Cell Biology, Leicester Institute of Structural and Chemical Biology, University of Leicester, Leicester LET 7HB, United Kingdom; ^dMax Planck Unit for the Science of Pathogens, 10117 Berlin, Germany; ^eInstitute of Microbiology, Leibniz Universität Hannover, 30419 Hannover, Germany; and ^fInstitut für Chemie und Biochemie, Freie Universität Berlin, 14195 Berlin, Germany

- H.-C. Flemming et al., Biofilms: An emergent form of bacterial life. Nat. Rev. Microbiol. 14, 563–575 (2016).
- D. Romero, C. Aguilar, R. Losick, R. Kolter, Amyloid fibers provide structural integrity to Bacillus subtilis biofilms. Proc. Natl. Acad. Sci. U.S.A. 107, 2230–2234 (2010).
- A. Driks, Tapping into the biofilm: Insights into assembly and disassembly of a novel amyloid fibre in *Bacillus subtilis*: Role of TapA in TasA fibre assembly and disassembly. *Mol. Microbiol.* 80, 1133–1136 (2011).
- S. Arnaouteli, N. C. Bamford, N. R. Stanley-Wall, Á. T. Kovács, Bacillus subtilis biofilm formation and social interactions. Nat. Rev. Microbiol. 19, 600–614 (2021).
- H. Vlamakis, Y. Chai, P. Beauregard, R. Losick, R. Kolter, Sticking together: Building a biofilm the Bacillus subtilis way. Nat. Rev. Microbiol. 11, 157–168 (2013).
- L. Hobley, C. Harkins, C. E. MacPhee, N. R. Stanley-Wall, Giving structure to the biofilm matrix: An overview of individual strategies and emerging common themes. *FEMS Microbiol. Rev.* 39, 649–669 (2015).
- L. Chai et al., Isolation, characterization, and aggregation of a structured bacterial matrix precursor. J. Biol. Chem. 288, 17559–17568 (2013).
- A. Diehl et al., Structural changes of TasA in biofilm formation of Bacillus subtilis. Proc. Natl. Acad. Sci. U.S.A. 115, 3237–3242 (2018).
- E. Erskine *et al.*, Formation of functional, non-amyloidogenic fibres by recombinant *Bacillus subtilis* TasA: Functional recombinant non-amyloidogenic TasA fibres. *Mol. Microbiol.* **110**, 897–913 (2018).
- N. E. Mammeri et al., Molecular architecture of bacterial amyloids in *Bacillus* biofilms. FASEB J. 33, 12146–12163 (2019).
- A. G. Stöver, A. Driks, Control of synthesis and secretion of the *Bacillus subtilis* protein YqxM. J. Bacteriol. 181, 7065–7069 (1999).
- 12. F. Chu, D. B. Kearns, S. S. Branda, R. Kolter, R. Losick, Targets of the master regulator of biofilm formation in *Bacillus subtilis. Mol. Microbiol.* **59**, 1216–1228 (2006).
- D. Romero, H. Vlamakis, R. Losick, R. Kolter, An accessory protein required for anchoring and assembly of amyloid fibres in *B. subtilis* biofilms: *B. subtilis* amyloid fibre accessory protein. *Mol. Microbiol.* **80**, 1155–1168 (2011).
- D. Romero, H. Vlamakis, R. Losick, R. Kolter, Functional analysis of the accessory protein TapA in Bacillus subtilis amyloid fiber assembly. J. Bacteriol. 196, 1505–1513 (2014).
- C. Earl et al., The majority of the matrix protein TapA is dispensable for Bacillus subtilis colony biofilm architecture. Mol. Microbiol. 114, 920–933 (2020).
- M. Bertolini et al., Interactions between nascent proteins translated by adjacent ribosomes drive homomer assembly. Science 371, 57-64 (2021).
- R. Abbasi et al., The bacterial extracellular matrix protein TapA is a two-domain partially disordered protein. ChemBioChem 20, 355-359 (2019).
- J. Böhning et al., Donor-strand exchange drives assembly of the TasA scaffold in Bacillus subtilis biofilms. Nat. Commun. 13, 7082 (2022).
- F. Wang, V. Cvirkaite-Krupovic, M. Krupovic, E. H. Egelman, Archaeal bundling pili of *Pyrobaculum calidifontis* reveal similarities between archaeal and bacterial biofilms. *Proc. Natl. Acad. Sci. U.S.A.* 119, e2207037119 (2022).
- S. Andrić, T. Meyer, M. Ongena, Bacillus responses to plant-associated fungal and bacterial communities. Front. Microbiol. 11, 1350 (2020).
- R. Bleich, J. D. Watrous, P. C. Dorrestein, A. A. Bowers, E. A. Shank, Thiopeptide antibiotics stimulate biofilm formation in *Bacillus subtilis. Proc. Natl. Acad. Sci. U.S.A.* **112**, 3086–3091 (2015).
- B. Kjeldgaard *et al.*, Fungal hyphae colonization by *Bacillus subtilis* relies on biofilm matrix components. *Biofilm* 1, 100007 (2019).
- A. Drozdetskiy, C. Cole, J. Procter, G. J. Barton, JPred4: A protein secondary structure prediction server. Nucleic Acids Res. 43, W389–W394 (2015).
- J. Jumper et al., Highly accurate protein structure prediction with AlphaFold. Nature 596, 583–589 (2021).
- N. Verma *et al.*, Computational investigation for modeling the protein–protein interaction of TasA(28-261)–TapA(33-253): A decisive process in biofilm formation by *Bacillus subtilis*. J. Mol. Model. 26, 226 (2020).
- E. Barbet-Massin et al., Rapid proton-detected NMR assignment for proteins with fast magic angle spinning. J. Am. Chem. Soc. 136, 12489–12497 (2014).
- P. Fricke et al., Backbone assignment of perdeuterated proteins by solid-state NMR using proton detection and ultrafast magic-angle spinning. Nat. Protoc. 12, 764–782 (2017).
- Y. Shen, F. Delaglio, G. Corrilescu, A. Bax, TALOS+: A hybrid method for predicting protein backbone torsion angles from NMR chemical shifts. J. Biomol. NMR 44, 213–223 (2009).
- A. E. Bennett, R. G. Griffin, J. H. Ok, S. Vega, Chemical shift correlation spectroscopy in rotating solids: Radio frequency-driven dipolar recoupling and longitudinal exchange. *J. Chem. Phys.* 96, 8624–8627 (1992).
- 30. K. Wüthrich, M. Billeter, W. Braun, Polypeptide secondary structure determination by nuclear
- magnetic resonance observation of short proton-proton distances. J. Mol. Biol. **180**, 715–740 (1984). 31. W. J. Allen, G. Phan, G. Waksman, Pilus biogenesis at the outer membrane of Gram-negative
- bacterial pathogens. *Curr. Opin. Struct. Biol.* 22, 500–506 (2012).
 G. Waksman, S. J. Hultgren, Structural biology of the chaperone-usher pathway of pilus biogenesis. *Nat. Rev. Microbiol.* 7, 765–774 (2009).

- M. Vetsch et al., Mechanism of fibre assembly through the chaperone–usher pathway. EMBO Rep. 7, 734–738 (2006).
- M. K. Hospenthal *et al.*, The cryoelectron microscopy structure of the type 1 chaperone-usher pilus rod. *Structure* 25, 1829–1838.e4 (2017).
- T. M. Pyburn et al., A structural model for binding of the serine-rich repeat adhesin GspB to host carbohydrate receptors. PLoS Pathog. 7, e1002112 (2011).
- R. Evans et al., Protein complex prediction with AlphaFold-Multimer. bioRxiv [Preprint] (2022). https://doi.org/10.1101/2021.10.04.463034 (Accessed 5 January 2022).
- K. Baskaran et al., Anomalous amide proton chemical shifts as signatures of hydrogen bonding to aromatic sidechains. Magn. Reson. 2, 765–775 (2021).
- L. J. Ball, R. Kühne, J. Schneider-Mergener, H. Oschkinat, Recognition of proline-rich motifs by protein-protein-interaction domains. *Angew. Chem. Int. Ed. Engl.* 44, 2852–2869 (2005).
 F. Castellani *et al.*, Structure of a protein determined by solid-state magic-angle-spinning NMR
- F. Castellani et al., Structure of a protein determined by solid-state magic-angle-spinning NMR spectroscopy. Nature 420, 99–102 (2002).
- C. Andréasson, J. Fiaux, H. Rampelt, M. P. Mayer, B. Bukau, Hsp110 is a nucleotide-activated exchange factor for Hsp70. J. Biol. Chem. 283, 8877–8884 (2008).
- A. Ceroni, A. Passerini, A. Vullo, P. Frasconi, DISULFIND: A disulfide bonding state and cysteine connectivity prediction server. *Nucleic Acids Res.* 34, W177–W181 (2006).
- M. Mirdita et al., ColabFold: Making protein folding accessible to all. Nat. Methods 19, 679–682 (2022).
- F. Ahimou, P. Jacques, M. Deleu, Surfactin and iturin A effects on *Bacillus subtilis* surface hydrophobicity. *Enzyme Microb. Technol.* 27, 749–754 (2000).
- S. S. Branda, J. E. González-Pastor, S. Ben-Yehuda, R. Losick, R. Kolter, Fruiting body formation by Bacillus subtilis. Proc. Natl. Acad. Sci. U.S.A. 98, 11621–11626 (2001).
- W. F. Vranken et al., The CCPN data model for NMR spectroscopy: Development of a software pipeline. Proteins. 59, 687-696 (2005).
- W. Rieping et al., ARIA2: Automated NOE assignment and data integration in NMR structure calculation. Bioinformatics 23, 381–382 (2007).
- A. Pines, M. G. Gibby, J. S. Waugh, Proton-enhanced nuclear induction spectroscopy. A method for high resolution NMR of dilute spins in solids. J. Chem. Phys. 56, 1776-1777 (1972).
- E. Barbet-Massin et al., Out-and-back 13C–13C scalar transfers in protein resonance assignment by proton-detected solid-state NMR under ultra-fast MAS. J. Biomol. NMR 56, 379–386 (2013).
- M. Krug, M. S. Weiss, U. Heinemann, U. Mueller, XDSAPP: A graphical user interface for the convenient processing of diffraction data using XDS. J. Appl. Crystallogr. 45, 568–572 (2012).
- S. Panjikar, V. Parthasarathy, V. S. Lamzin, M. S. Weiss, P. A. Tucker, *Auto-Rickshaw*: An automated crystal structure determination platform as an efficient tool for the validation of an X-ray diffraction experiment. *Acta Crystallogr. D Biol. Crystallogr.* 61, 449–457 (2005).
- 51. A. J. McCoy et al., Phaser crystallographic software. J. Appl. Crystallogr. 40, 658–674 (2007).
- P. Emsley, B. Lohkamp, W. G. Scott, K. Cowtan, Features and development of *Coot. Acta Crystallogr.* D Biol. Crystallogr. 66, 486–501 (2010).
- G. N. Murshudov, A. A. Vagin, E. J. Dodson, Refinement of macromolecular structures by the maximum-likelihood method. *Acta Crystallogr. D Biol. Crystallogr.* 53, 240–255 (1997).
- V. B. Chen et al., MolProbity: All-atom structure validation for macromolecular crystallography. Acta Crystallogr. D Biol. Crystallogr. 66, 12–21 (2010).
- P. Schuck, Size-distribution analysis of macromolecules by sedimentation velocity ultracentrifugation and lamm equation modeling. *Biophys. J.* 78, 1606–1619 (2000).
- T. M. Laue, B. D. Shah, T. M. Ridgeway, S. L. Pelletier, "Computer-aided interpretation of sedimentation data for proteins" in *Analytical Ultracentrifugation in Biochemistry and Polymer Science*, S. E. Harding, A. J. Rowe, J. C. Horton, Eds. (Royal Society of Chemistry, 1992), pp. 90–125.
- C. A. Brautigam, "Calculations and publication-quality illustrations for analytical ultracentrifugation data" in *Methods in Enzymology* (Elsevier, 2015), pp. 109–133.
- K. Tamura, G. Stecher, S. Kumar, MEGA11: Molecular evolutionary genetics analysis version 11. Mol. Biol. Evol. 38, 3022–3027 (2021).
- 59. M. A. Larkin et al., Clustal W and Clustal X version 2.0. Bioinformatics 23, 2947–2948 (2007).
- C. Gille, M. Fähling, B. Weyand, T. Wieland, A. Gille, Alignment-annotator web server: Rendering and annotating sequence alignments. *Nucleic Acids Res.* 42, W3–W6 (2014).
- V. A. Higman, P. Schmieder, A. Diehl, H. Oschkinat, Structural investigation of the TasA anchoring protein TapA from Bacillus subtilis. Protein Data Bank. https://www.rcsb.org/structure/6QAY. Deposited 20 December 2018.
- Y. Roske, U. Heinemann, Structural investigation of the TasA anchoring protein TapA from Bacillus subtilis. Protein Data Bank. https://www.rcsb.org/structure/6HQC. Deposited 24 September 2018.
- V. A. Higman, P. Schmieder, A. Diehl, H. Oschkinat, Structural investigation of the TasA anchoring protein TapA from Bacillus subtilis. *Biological Magnetic Resonance* Data Bank. https://bmrb.io/ data_library/summary/index.php?bmrbId=34341. Deposited 20 December 2022.
- F. Lindemann, M. Hiller, H. Oschkinat, Solid-state NMR assignment data of TasA filaments. Biological Magnetic Resonance Data Bank (BMRB). https://bmrb.io/data_library/summary/index. php?bmrbld=51785. Deposited 17 January 2023.
- F. Lindemann, M. H. Glomm, H. Oschkinat, Solid-state NMR assignment data of TasA filaments. Zenodo. https://zenodo.org/record/7534572#.ZCo9YntBw2w. Deposited 31 March 2023.