Architecture of the flexible tail tube of bacteriophage SPP1

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Bacteriophage SPP1 is a double-stranded DNA virus of the Siphoviridae family that infects the bacterium Bacillus subtilis. This family of phages features a long, flexible, non-contractile tail that has been difficult to characterize structurally. Here, we present the atomic structure of the tail tube of phage SPP1. Our hybrid structure is based on the integration of structural restraints from solid-state nuclear magnetic resonance (NMR) and a density map from cryo-EM. We show that the tail tube protein gp17.1 organizes into hexameric rings that are stacked by flexible linker domains and, thus, form a hollow flexible tube with a negatively charged lumen suitable for the transport of DNA. Additionally, we assess the dynamics of the system by combining relaxation measurements with variances in density maps.
tailed bacteriophages—order Caulovirales—comprise the prevailing majority of known phages and are subdivided into three families based on their tail morphology. Podoviridae feature a short tail, Myoviridae a long, contractile tail and Siphoviridae a long noncontractile, flexible tail, respectively. The latter two possess a helical tail tube assembled around a tape measure protein that is tapered by a tail completion protein. Contractile tail tubes are furthermore environed by a sheath. These tail-structures are crucial for host-cell recognition, membrane penetration, and DNA transport into the host. Recently, a high-resolution cryo-EM structure of the short fiber-less tail from the Podoviridae T7 phage was reported at a resolution of 3.3 Å. Also, a cryo-EM structure of the prehost attachment baseplate including two rings of the tail tube and sheath proteins from the Myoviridae T4 phage was solved at a resolution of 3.8–4.1 Å, and a cryo-EM reconstruction focused solely on the tail tube was obtained from the same images at a resolution of 3.4 Å. Structural analysis of the tube arrangement of Siphoviridae phages was for long hindered by the variable tail bending that results from its flexibility (see Fig. 1 of Tavares et al.13). Structural information was limited to pseudo-atomic models, which were generated for SPP16 based on solution nuclear magnetic resonance (NMR) structures of monomeric tail tube proteins (TTPs), and for phages T5 and λ by fitting structures of monomeric TTPs19 into a 6 Å cryo-EM density map. In 2020, a cryo-EM model of the baseplate of the Staphylococcus aureus 80a phage was reported, which includes two rings of the tail tube that are anchored within the baseplate and are, thus, not part of the flexible tube region10. Also, cryo-EM models of the tails of the flagellotropic tailed bacteriophage YSD111 and the Siphoviridae-like gene transfer agent of Rhodobacter capsulatus12 were reported recently. All models show a striking structural homology between TTPs from Myoviridae and Siphoviridae; as well as between these phage TTPs and the tube-forming proteins from other injection systems, like the bacterial type VI secretion system13 and the extracellular injection system from bacteria and archaea14. The tube-forming proteins share a common fold composed of two orthogonally packed β-sheets that hexamericize through the formation of an inner β-barrel that defines the lumen of the tube. However, variable elements such as loops, N-arms and C-arms are critical to mediate intermonomer contacts driving tube assembly in these systems, and sparse high resolution data are available on the position of these elements within the long-tailed phage tubes6-12.

In the Siphoviridae SPP1 phage, the tail tube consists of the TTPs gp17.1 and gp17.1* in a ratio of 3:1, with the latter being generated by a translational frameshift adding a fibronectin type III (FN3) domain to the C-terminus of the protein. However, virions only containing gp17.1 are still viable and infectious, indicating that the additional C-terminal FN3 domain is dispensable for phage assembly and infection15. gp17.1 monomers are unstable in solution and spontaneously self-polymerize into long tubes in vitro which are indistinguishable from native tubes6,16. Previously, we presented the proton-detected solid-state NMR (ssNMR) assignment of deuterated, 100% back-exchanged gp17.1 tubes and deduced secondary structure information from the assigned chemical shifts, which confirmed and extended an existing homology model of a polymerized gp17.1 subunit6,16. Additionally, we introduced new concepts based on the use of specifically labeled isoleucine-methyl groups to simplify ssNMR experiments and 4D proton-detected ssNMR experiments at 40 kHz magic-angle spinning (MAS) and 900 MHz proton Larmor frequency were used to probe long-range distance restraints between the following: 1) amide groups (Fig. 1c, left panel); 2) methyl groups; 3) methyl and amide groups globally (Fig. 1c, right panel); 4) methyl and amide groups at protein–protein interfaces. All of these experiments generated highly unambiguous restraints due to their high-dimensionality (4D) or spectral simplicity (amino-acid specific methyl labeling)—as visualized in Fig. 1d where a set of consistent restraints (amide-amide and methyl-amide contacts) defines the inner β-barrel motif of the tail tube formed by the β-strands β2, β3, β6.1, and β5.2. In mixed labeled samples (as detailed in Table S1) magnetization transfer between methyl and amide groups is solely possible at protein–protein interfaces because half of the subunits are 15N labeled and the other half are methyl labeled. Hence, these samples deliver a set of restraints that defines the relative organization of gp17.1 subunits within the tube. Figure 1e shows an exemplary protein interface between the N-terminus of a subunit, including the Ile18 labeled methyl group, and the C-terminus of another subunit.

For cryo-EM experiments, we purified ΔN–3 gp17.1 which is indistinguishable from wt gp17.1 as judged by solid-state NMR (see Figs. S1 and S2). Curvy tubes were observed in the micrographs (Fig. S3). For image processing those tubes that

Results

Hybrid structure calculation. To determine the structure of the tail-tube of SPP1, we performed a hybrid structure calculation using the inferential structure determination (ISD) approach37 integrating data from solid-state NMR and cryo-EM simultaneously. During structure calculation only the structure of a single monomer was represented and refined. The structures of the other subunits were generated by applying symmetry operators to the subunit structure. The use of an exact symmetry is justified by the NMR data that are only consistent with a highly symmetric sample. We represented two stacked rings in the structure calculation. Each ring was composed of six subunits such that interactions between twelve subunits were considered in the structure calculation.

For the collection of solid-state NMR long-range distance restraints, we overexpressed a set of differently labeled TTP gp17.1 in E. coli, purified them, and let them self-polymerize into native-like tail tubes as detailed in the “Methods” section. Torsion angles were predicted based on assigned backbone chemical shifts (Fig. 1a). Specific precursor molecules were supplemented during protein expression in deuterated media, introducing NMR visible methyl groups within certain amino acids of gp17.1 (Fig. 1b). We produced samples that were homogeneously methyl and 15N labeled, as well as samples that were heterogeneous mixtures of 50% methyl-labeled and 50% 15N labeled subunits. The latter samples were used to detect intermolecular interfaces, as previously described by us19. All eleven investigated methyl-labeled and/or deuterated samples and their precursors are listed in Table S1. 4D and 3D proton-detected ssNMR experiments at 40 kHz magic-angle spinning (MAS) and 900 MHz proton Larmor frequency were used to probe long-range distance restraints between the following: 1) amide groups (Fig. 1c, left panel); 2) methyl groups; 3) methyl and amide groups globally (Fig. 1c, right panel); 4) methyl and amide groups at protein–protein interfaces. All of these experiments generated highly unambiguous restraints due to their high-dimensionality (4D) or spectral simplicity (amino-acid specific methyl labeling)—as visualized in Fig. 1d where a set of consistent restraints (amide-amide and methyl-amide contacts) defines the inner β-barrel motif of the tail tube formed by the β-strands β2, β3, β6.1, and β5.2. In mixed labeled samples (as detailed in Table S1) magnetization transfer between methyl and amide groups is solely possible at protein–protein interfaces because half of the subunits are 15N labeled and the other half are methyl labeled. Hence, these samples deliver a set of restraints that defines the relative organization of gp17.1 subunits within the tube. Figure 1e shows an exemplary protein interface between the N-terminus of a subunit, including the Ile18 labeled methyl group, and the C-terminus of another subunit.

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Fig. 1 Solid-state NMR data for hybrid structure calculation defines local structure. a Backbone dihedral angles define the secondary structure (arrows represent β-sheets, barrels α-helices). The loop (40–59) and the C-arm (143–176) are highlighted in purple and pink, respectively. b Isoleucine Cβ1-methyl, alanine Cβ-methyl, leucine-valine Cβ/Cγ1-methyl, threonine Cγ2-methyl, and methionine Cε-methyl labeling renders those moieties NMR-visible and yields highly resolved NMR spectra (green). c Long-range restraints between amide protons are extracted from a 4D HNNHN spectrum (orange), whereas long-range restraints between amide and methyl groups are extracted from a series of 3D HNNH spectra (blue). 2D planes from both types of spectra are superimposed on a 2D hHN fingerprint spectrum (gray). d Representative long-range distance restraints (dashed lines) defining the inner β-barrel of the tail tube. The colored β-strands belong to one monomer, whereas the gray β-strands are from the neighboring subunits. Amide-amide contacts are highlighted in violet, amide-methyl contacts in cyan. e Schematic representation of long-range distance restraints between the Cβ1 methyl group of Ile18 and amide groups, defining the interface between the N-terminus of one monomer i (including Ile18; in cyan) and the C-arm (143–176) of another monomer j (in pink) within the tail tube. Protons are colored in red.

appeared most straight were selected. 3D reconstruction (see "Methods" section) yielded a density map with an average resolution of 4.3 Å (Figs. 2a and S4). The local resolution varies significantly (Fig. 2b–d), from about 3.5 Å at the inner ring where the β-strands are well resolved (Fig. 2e, f), to worse than 5 Å at the periphery.

Overall structure of the SPP1 tail tube. Figure 3 shows the structure of the tail-tube of gp17.1 as determined by hybrid structure calculation (see also Movie S1 and Fig. S5) with a heavy atom RMSD of 1.8 ± 0.9 Å (Backbone RMSD of 1.1 ± 0.5 Å) over the entire protein sequence. The monomer of gp17.1 forms a central β-sandwich-type fold consisting of eight β-strands. This fold is flanked by an α-helix (74–86) and loop regions, of which the long C-terminal arm (C-arm, 143–176) stands out (Fig. 3a). Six gp17.1 subunits assemble into a ring with the inner 24 β-strands forming a β-barrel that defines the inner lumen of the tube. This inner lumen exhibits a negative electrostatic potential (see Fig. S6) which facilitates sliding of the viral DNA through the tube by repelling it from the surface. Additionally, inner ring contacts are mediated by the large loop region 40–59. The α-helices are arranged almost parallel to the tail tube axis (Fig. 3b). These rings stack on each other with a rotation of 21.9° forming a right-handed helical, hollow tube (Fig. S3c). The surface of one gp17.1 subunit features a hydrophobic patch that is shaped by sidechains of various hydrophobic amino acids (Fig. 3d). In the context of the tail tube complex the C-arm of the superjacent subunit folds onto the outer β-sheet of the β-sandwich fold by anchoring the sidechain of Gln162 into a pocket (Fig. 3e). This interaction obscures the lipophilic area whereupon the complex is stabilized, because the number of unfavorable hydrophobic contacts with the solvent is reduced (Fig. 3f). This explains why a previously reported C-terminally truncated mutant of gp17.1 remains monomeric. Additional ring-to-ring contacts are mediated by the loop region 40–59 which interacts with five neighboring subunits—mostly by establishing electrostatic contacts (Fig. 3g).

Structural alignments of gp17.1 and existing TTP structures of other systems show high similarity as expected (Fig. S7)—all featuring hexameric, helically stacked rings with subunits consisting of a β-sandwich-type fold and one parallel α-helix. Loop 40–59 is present at the interface between subunits in all described Siphoviridae phages, Siphoviridae-like systems, and T4 phage, suggesting that it is a conserved structural element across both Siphoviridae and Myoviridae families as it was also proposed to play a regulatory role during tail polymerization. The mentioned C-armed is only present in SPP1 and 80a (even if not completely resolved). It might be a critical element regulating the tail structure in a subgroup of Siphoviridae. YSD1 phage features a similar intermolecular contact—an inserted domain after the α-helix that similar to the C-arm folds onto the outer β-sheet of the β-sandwich of a neighboring subunit. However, this contact is established within the ring and not between the rings. Phages, like Siphoviridae, YSD1 carry an additional N-terminal loop that promotes intermolecular contacts. The organization of the tail tube of T5 phage is different since it uncommonly exhibits a trimeric ring resulting from the fusion of every two subunits within the hexamerization domain. The Siphoviridae-like gene
transfer agent tail tube does not bear any of these elements. The Myoviridae T4 phage TTP gp19 features two additional linkers that mediate intermolecular interactions—which facilitates contact to ten different subunits over an area of 6706 Å² within the tail tube. gp17.1, however, only interconnects with six different subunits over 4850 Å². This dramatically reduced contact—in addition to not being bundled in a sheath—is expected to enable flexibility of this Siphoviridae tail tube as proposed previously for phages T5 and λ.

**Dynamic regions mediate tail bending.** To determine the driving forces contributing to the flexibility of the tail tube of SPP1, we created a model of a bent tail tube based on the structure of polymerized gp17.1 subunits does not only allow to deduce symmetry restraints but also limits the position of all atoms within the density. Rings with a thickness of 38.5 Å (as represented by the straight arrows) stack onto each other with a rotation of 21.9° (as represented by the bent arrow). b-d Local resolution of the cryo-EM map increases going from outer to the inner surface of the tail tube (as represented by the color gradient). e The inner region of the map reveals a highly resolved β-barrel and allows for the positioning of bulky sidechains as exemplified for Tyr67 and Tyr68. The direction of the tail structure is baseplate upwards.

Fig. 2 Cryo-EM data for hybrid structure calculation provides global and local information. a 3.5-6 Å cryo-EM map of the tail tube of SPP1 consisting of polymerized gp17.1 subunits does not only allow to deduce symmetry restraints but also limits the position of all atoms within the density. Rings with a thickness of 38.5 Å (as represented by the straight arrows) stack onto each other with a rotation of 21.9° (as represented by the bent arrow). b-d Local resolution of the cryo-EM map increases going from outer to the inner surface of the tail tube (as represented by the color gradient). e The inner region of the map reveals a highly resolved β-barrel and allows for the positioning of bulky sidechains as exemplified for Tyr67 and Tyr68. The direction of the tail structure is baseplate upwards.

shows representative 15N relaxation dispersion curves; flat profiles indicate the absence of motion, whereas decaying profiles indicate the presence of dynamics. Most residues are involved in slow motions (Fig. S12). The residues belonging to the β-barrel can be fitted in a combined approach to a two-state model assuming in a simplified way the existence of two distinct conformations of the β-barrel (Fig. 4f). Calculated chemical shift differences between both states are higher for residues in proximity to the hinge regions. Thus, we propose that these slow motions represent tube bending. The global, collective nature of this motion does not impose heterogeneity onto the cryo-EM map since only straight tubes are considered for structure calculation.

Overall, our hybrid data support a model where the C-arm (143–176) and the loop (40–59) act as bellows contributing to tail tube bending by stretching. Our dynamic structure of the tail tube is reminiscent of a molecular spinal column. The hexameric rings forming the inner β-barrel would be in this picture the vertebrae, while the flexible parts (C-arm and loop) correspond to the intervertebral disks. The flexibility of the system might facilitate the screening of the bacterial membrane to find the receptor for infection initiation. We expect that the combination of sophisticated ssNMR experiments and cryo-EM will help to characterize structures of other dynamic and/or flexible supramolecular assemblies, in particular those systems where the conformational flexibility leads to a lack of resolution in cryo-EM reconstructions—while depending on the timescale of these dynamics the quality of NMR spectra may not be affected.

**Methods**

Preparation of deuterated protein samples for proton-detected solid-state NMR measurements. Protein samples and their preparation are summarized in Table S1. gp17.1 protein was expressed, purified and polymerized as described in the following. E. coli BL21 were transformed with a pETM13 vector containing the gp17.1 sequences including a C-terminal His-tag (Fig. S1). In a three step protocol, the bacterial cultures were adapted to D2O conditions: In the first step, 12.5 mL LB medium was mixed with 12.5 mL fully deuterated M9-medium with 13CD6-glucose and 15ND2Cl as the sole carbon and nitrogen sources. By lyophilizing and redissolving in D2O twice, exchangeable protons of the M9 medium components
Fig. 3 Structure of polymerized gp17.1 forming the tail tube of the bacteriophage SPPI. a Final ten lowest-energy structures of a gp17.1 subunit which consist of a central β-sandwich-type fold (turquoise) that is flanked by an α-helix (pink), a large loop and an extended C-terminal arm (C-arm). b Six gp17.1 monomers form a hexameric ring. The inner β-sheets of the β-sandwiches organize in a β-barrel motif that forms the lumen of the tube. c These hexameric rings stack onto each other in a helical fashion creating a hollow tube. Ring-to-ring contacts are mediated by the two loop regions (highlighted in red)—especially by the C-arm that folds onto the subjacent ring. d The molecular lipophilicity potential of gp17.1 reveals a hydrophobic patch on the surface of one subunit i. The color gradient represents the lipophilicity potential. e, f This unpolar area is obscured by the C-arm (pink) of the superjacent subunit j within the complex of the tail-tube—by anchoring the sidechain of Gin162 into a pocket. g The loop of subunit i (turquoise) features mostly electrostatic interactions with five neighboring subunits (gray, purple, green, beige, and orange) within the complex. Charged amino acids are colored in red (negative) and blue (positive). The direction of the tail structure is baseplate upwards.

Preparation of fully protonated protein samples for carbon-detected solid-state NMR and cryo-EM measurements. gp17.1 and ΔN–3 gp17.1 were expressed, purified, and polymerized as described above, only that D₂O was exchanged to H₂O. ΔN–3 gp17.1 for cryo-EM was expressed in LB medium. The protein pellets and a few DSS crystals for spectral referencing and temperature control were filled into 3.2 mm rotors. For cryo-EM, 20 mM sodium phosphate in the final buffer was replaced by 20 mM Tris-HCl (pH 7.4).

Solid-state NMR spectroscopy. Solid-state NMR spectroscopy of the methyl-labeled and/or deuterated protein samples was conducted with a 1.9 mm, four-channel (1H, 13C, 15N, and 2H) probe at 40 kHz magic-angle spinning (MAS) frequency and an external magnetic field strength according to 900 MHz 1H Larmor frequency. The temperature was calibrated to around +18 °C by means of internally added DSS. 2D hCH and 3D HνH spectra were recorded as described previously19, a 2D kNH spectrum was recorded as detailed before16. Pulse program, acquisition, processing and reconstruction parameters for the 2D hCH, 3D HνH, 3D HCH, and 4D HνHνH spectra are summarized in Supporting Information Tables S2–S6.

Solid-state NMR spectroscopy of the fully-protonated samples was conducted with a 3.2 mm, triple-channel (1H, 13C, and 15N) probe at 11 kHz MAS frequency and an external magnetic field strength according to 900 MHz 2H Larmor frequency. The temperature was calibrated to around +10 °C by means of internally added DSS. 2D 13C–13C correlation spectra with 50 ms proton-driven spin diffusion (PDSD) mixing were recorded as fingerprints to compare wild-type gp17.1 with the ΔN–3 gp17.1 mutant.

Long-range distance restraints were extracted from the recorded spectra by peak picking in CcpNmr41. Methyl groups were assigned based on our previous work19, on the basis of the assignment precursor (Table S1) or by correlations to sequential amide groups. In the alanine-methyl sample (Table S1) protons scrambled into the Hy2 position of isoleucines.

Relaxation measurements by solid-state NMR. 15N R1 and 15N R1ρ relaxation rates were measured by a series of pseudo-3D experiments (1H, 15N, delay/spinlock strength). The delay times for the R1ρ experiments were 0.5, 1, 1.5, 2, 3, 4, 8, 16, and 32 s. For the R1ρ experiments pseudo-3D experiments spinlock strengths of 9, 7, 5.5, 5, 4.5, 4, 3.5, 3, 2.5, and 2 kHz and spinlock durations of 5, 10, 20, 40, 80, 100, 140, and 200 ms were used. The peak heights from the resulting 2D hνH correlation spectra were extracted with CcpNmr41 and fitted as a function of relaxation time to a monoexponential function. This results in one global R1 relaxation rate and 15NR 1 and 15NR 1
**Fig. 4 Bending of the tail tube is mediated by flexible hinge regions.** a Model of a bent SPP1 tail tube with a curvature radius of 655 Å (as indicated by the arrow). The model is based on the structure of straight tubes and 2D class averages of bent tubes. Most structural changes are found on the outside of the tube (red) which implies that the bending process is mediated by stretching. b Regions that act as hinges during bending of the tube are colored in pink. c Variances in the cryo-EM map (pink) match the hinge regions. d Also, hinge regions are associated with highest $^{15}$N R1 (color key) and R2 ($\phi$ thickness of wire) relaxation rates. White coloring represents missing values. e Decaying relaxation dispersion profiles indicate the presence of slow motions. f Relaxation dispersion profiles of the inner $\beta$-barrel can be fitted in a correlated manner to a two-state exchange process. Smallest chemical shift changes correlate with middle regions of the $\beta$-barrel which are furthest away from the hinge regions (color key). The direction of the tail structure is baseplate upwards. Source data are provided as a Source Data file.

The spinlock strength-dependent R2 relaxation rates, R1 relaxation rates and the angle between the spinlock offset frequency ($\omega_2$) and the chemical shift offset from that (0) in a residue specific manner:

$$
R_{2,\text{obs}} = R_{2,\text{p},0} \cos^2 \theta,
$$

$$
\theta = \tan^{-1} \frac{\omega_2}{\Omega},
$$

The errors were estimated by Monte Carlo simulations. The fits were repeated 250 times with the R2 errors being multiplied by a random number between 0 and 1. The R2 errors were calculated in a similar way by performing the fits 1000 times using average noise from the spectra as input.

**Cryo-EM image processing and helical reconstruction.** MotionCor2 was used for movie correction and CTFFIND parameters were fitted with Gctf. All image processing was done using RELION 2.1. Fibrils were manually picked, and segments were extracted with an interbox distance of 10% of the box sizes, which yielded 69,282 segments. Box sizes were chosen as 200 pixels. Segments from micrographs for which Gctf estimated a resolution worse than 5 Å were discarded, which left 64,760 segments. 2D classification was used to select those classes that show sufficient detail in the class averages (Fig. S13). The initial model for 3D reconstruction was built with the Relion relion_helical_toolbox program with the simulate helix option, which places spheres along a helix, using a rise of 40 Å and a twist of 21°. The initial model was refined using 3D classification ($K=3$). The class yielding the highest resolution was used as initial model for further 3D classification with all particles using $K=5$ classes and a $T$ value of 4. The best resolved class contained 10,682 segments which were used for further refinement. A soft mask enclosing three rings was used for further refinements. The number of filaments and segments used for the final reconstruction was 1866 and 5965, respectively. The final optimized helical symmetry was C6, with a helical rise of 38.46 Å and a twist of 21.89°. Gold-standard refinements were performed by selecting entire fibrils and splitting the data set accordingly into an even and odd set. The Fourier shell correlation was computed between two half maps. According to the 0.143 criterion the obtained resolution is 4.3 Å (Fig. S14). To obtain a robust resolution estimate, the FSC curve was fitted using $1/[e^{(A-A^{\infty})/B}]+1)^2$, yielding $A=0.122$, $B=0.015$, and $C=0.228$. The 0.143 criterion then yields a resolution of 4.0 Å (Fig. S14). Image processing and reconstruction details can be found in Table S8. The final map was sharpened with the EMAN2 tool e2picro3d.py with a B-factor of $-150$ Å$^2$, locally normalized and filtered to 3.5 Å.

The local resolution as shown in Fig. 2b–d, was estimated by comparing a density map computed from the atomic model with the reconstructed density map by FSC. Both density maps were first interpolated on finer grid (pixel size of 70 e$^-$/Å$^2$. Applied underfocus values ranged between 0.2 and 1.7 µm. The pixel size was calibrated to 0.935 Å as calibrated using gold diffraction rings within the powerspectra of a cross grating grid (EMS, Hatfield). Details of data acquisition are summarized in Table S8.
0.468 Å. The (local) FSC calculation was done with the EMAN2 program e2fsc.py, using a FSC cutoff of 0.5 for defining the resolution.

Structure calculation. The hybrid structure calculation aims to combine the experimental information from NMR and cryo-EM. Distance restraints were derived from NMR peak lists and incorporated using a logistic restraint potential similar as described before. The density map from cryo-EM was incorporated as a real-space map restraint. Both data sets pose their own challenges: The NMR distance restraints are highly ambiguous due to the helical symmetry of the tail tube. NMR peaks stemming from the homogeneously mixed samples can result from a contact within the monomer or from contacts between different subunits (i.e., between the monomer and one of its virtual copies generated by the symmetry operators). However, NMR peaks stemming from the heterogeneously mixed samples can clearly be assigned as intermolecular long-range restraints. We considered all possible interactions between all members of both hexameric rings. Therefore, in total 12 possible contacts were combined as an ambiguous distance restraint. Another challenge is posed by the generous upper bound of 7 Å for the distance restraints. Also, the cryo-EM map itself has a substantial resolution inhomogeneity and is not sufficient for an unambiguous tracing of the backbone in the outer β-strands and the ε-terminus.

A particular challenge was posed by the estimation of the registers of adjacent β-strands. Due to the insufficient resolution of the density map in the flexible regions (e.g., residues 40–49) and the large distance upper bounds, many relative registers between adjacent strands seem possible in principle. To infer the register that is most consistent with the NMR and cryo-EM data, we developed a new probabilistic restraint that probes all possible registers between adjacent strands. The estimated registers were then imposed as additional hydrogen bonding restraints to increase the regularity of the gp17.1 tail tube structure. Only the combination of the local information provided by the NMR restraints with the global shape information encoded in the cryo-EM map allowed us to compute a model that is closest (in local structure and subunit contacts) to the straight tail tube, as we do not have high-resolution information on the curved tail tube. Therefore, a network of harmonic distance restraints (random atom pairs between 3 and 15 Å) was defined with target distances from the straight tail tube. In addition the β-sheets at the inside of the tubes were position-restrained to keep curvature and relative ring positions. DireX was used to optimize the model under these distance and position restraints (without density map restraints). The curved model (Fig. 4a) represents a model that is closest (in local structure and subunit contacts) to the straight tail tube, while adopting the imposed curvature and ring distances. The amount of fulfilled restraints after bending reveals regions of the protein that are exposed to environment changes (red color coding in Fig. 4a).

ChimeraX morph command was used to create a trajectory between the straight and the bent tail tube (standard settings). During that procedure hinge and core regions are identified by a reimplementation of the morph server.

Figure creation. Figures were created with ChimeraX.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

Solid-state NMR chemical shift assignments are deposited at the Biological Magnetic Resonance Data Bank under the accession code 27469. The cryo EM electron density map is deposited at the Electron Microscopy Data Bank under the accession code EMD-10792. Protein structures are deposited at the Protein Data Bank under the accession codes 6EGY (hybrid structure of the SPPI tail tube by solid-state NMR and cryo EM; final EM refinement) and 6FQZ (hybrid structure of the SPPI tail tube by solid-state NMR and cryo EM; NMR ensemble). The authors declare that any other data supporting the findings of this study are available within the article and in its Supplementary Information, or from the authors upon request. Source data are provided with this paper.

Code availability

Codes for curve fitting, error analysis, and structure calculation are available from the corresponding authors upon request.

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