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Validated near-atomic resolution structure of bacteriophage epsilon15 derived from cryo-EM and modeling

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High-resolution structures of viruses have made important contributions to modern structural biology. Bacteriophages, the most diverse and abundant organisms on earth, replicate and infect all bacteria and archaea, making them excellent potential alternatives to antibiotics and therapies for multidrug-resistant bacteria. Here, we improved upon our previous electron cryomicroscopy structure of Salmonella bacteriophage epsilon15, achieving a resolution sufficient to determine the tertiary structures of both gp7 and gp10 protein subunits that form the T = 7 icosahedral lattice. This study utilizes recently established best practice for near-atomic to high-resolution (3–5 Å) electron cryomicroscopy data evaluation. The resolution and reliability of the density map were cross-validated by multiple reconstructions from truly independent data sets, whereas the models of the individual protein subunits were validated adopting the best practices from X-ray crystallography. Some sidechain densities are clearly resolved and show the subunit–subunit interactions within and across the capsomeres that are required to stabilize the virus. The presence of the canonical phage and jellyroll viral protein folds, gp7 and gp10, respectively, in the same virus suggests that epsilon15 may have emerged more recently relative to other bacteriophages.

Results and Discussion

Cryo-EM Data and Map. In the current investigation, we acquired a unique cryo-EM dataset of ε15 (Fig. L4) using a different electron microscope than the one used in our previous study (4). A substantially larger fraction of high-quality images was recorded with the current electron microscope at liquid nitrogen specimen temperature over the previous instrument, operated at liquid helium temperature. The power spectrum of the boxed-out particles generally shows signals out to 4–5 Å resolution (Fig. S1). The reconstruction was based on a recently implemented protocol (19) (Materials and Methods). We adopted the gold standard definition for the resolution estimate (20), whereby we split the particle images into two subsets at the onset of image processing, reconstructed the datasets individually, and combined after determining the resolution estimate (Fig. 1B). Independent initial models were built de novo and used for the subsequent particle refinements in each of the two subsets of particle images (Materials and Methods) (Fig. S2). The Fourier Shell Correlation (FSC) between the two independently determined reconstructions was computed and indicated a resolution 4.5 Å using the 0.143 threshold (21) for the combined dataset (Fig. 1C). This map resolution measure differs from the commonly used approach in that, whereas two maps were computed with particle images split after all of the particles’ orientations were determined using a single initial model and common models for all subsequent refinements. This conventional procedure has now been recognized to be a substandard measure of the map resolution (20, 22, 23). Additionally, better image data, improving both image-processing and model-building protocols, more rigorously measuring map resolution and adopting best practice in model refinement as established in X-ray crystallography. The final ε15 full-atom model clearly demonstrates an intricate set of interactions between the proteins, both within and across capsomeres, contributing to overall viral stability, as well as illustrating new potential interactions during its assembly and maturation.

Supporting Information


The authors declare no conflict of interest.

Data deposition: The atomic coordinates and density maps have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 3J40) and Electron Microscopy Data Bank, www.emdbdata.org (EMDB ID code 5678).

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a model for the entire virus was transformed into a 3.5 Å resolution density map, which was compared with the raw cryo-EM density map. Using the 0.5 FSC criterion, a more appropriate measure in map-to-model comparisons (21), estimates their structure match up to ~5 Å on average (Fig. 1D).

In line with the new resolution measure and the overall structural features, we computed a FSC between our current map [Electron Microscopy Data Bank (EMDB) ID code 5678] and the previously published map (EMDB ID code 5003). (Fig. 1C). At FSC = 0.333 (21), these two maps are structurally similar up to ~4.8 Å on average. This measure suggests the resolution of the previous map (claimed to be 4.5 Å based on 0.5 criterion for FSC computed from maps with split data after they were refined with a conventional common model approach) (4) was slightly overestimated. Nevertheless, the resolvable features in many places within the two density maps are quite comparable (Fig. S3). The resolvability of structural features like the β-strand separation and loop connectivity in the current map is a clear improvement over the previous map, particularly in gp10 (Fig. S3 C and D). Furthermore, as shown in Figs. 2 and 3, the new density map is more uniformly resolved throughout, improving the interpretability of the structural features and the accuracy of the model building.

**Model Building.** In X-ray crystallography, model building in 4–5 Å resolution density maps is considered to be particularly challenging (24). However, in our previous study of ε15, we developed a manual modeling approach to trace the Ca chain of gp7 (4, 25). Briefly, the presence of five distinct helices in gp7 was sufficient to anchor a sequence to structure correspondence, from which we were able to trace the Ca backbone aided by a density skeleton connectivity (26, 27). However, in the case of gp10, model building was not possible due to limited resolvability of the structural features. Since our initial success in building the gp7 model, our de novo modeling protocol has been enhanced with improved algorithms and automation capability (28–30).

The enhanced resolvability along with improved modeling methods allowed us to generate complete all-atom models for gp7 and gp10 from our new density map (Figs. 2 and 3). As illustrated in Fig. S4, we used an iterative model refinement process using ROSETTA (31), COOT (32), and PHENIX (33) to refine mainchain and sidechain fit to density, while simultaneously improving model geometry, bond distances, and inter/intrasubunit interfaces (SI Materials and Methods). Best practice procedures from X-ray crystallography were adopted to improve and validate our models (Table 1) based on recent recommendations by the EMDB model validation taskforce (34). Although fully refined and validated, it should be noted that density at this resolution does not provide “perfect” restraints and thus potential errors (i.e., imperfect H-bonding, registry errors, etc.) may be present (24).

**Coat Protein, gp7.** For constructing and validating the gp7 model, we began with our original models of the seven gp7 subunits [Protein Data Bank (PDB) ID code 3CSB] constructed from the previous ε15 density map (4). We first fit these models to our new density map (Fig. 1B), segmented the subunits in the asymmetric unit, and computed an average gp7 density map (equivalent to Non-Crystallographic Symmetry (NCS) averaging in crystallography) with the six hexon subunits (Fig. 2A). We used our Pathwalker protocol (28) to determine various possible topologies of gp7 constrained by our averaged gp7 subunit density. Our results showed two different populations of models. One model was equivalent to our previous Ca trace, whereas the other model had a strand swap in the A-domain and a slight reorientation of the C terminus. Comparing both models to our latest density map, the new gp7 model, with the strand swap, agreed more favorably with the density (Fig. S3 A and B). Seven independent all-atom models were subsequently constructed for the corresponding subunits in the asymmetric unit and refined using this initial Ca trace.

![Image](image-url)
Fig. 3. Structure of the staple protein, gp10. (A) An averaged gp10 density map is shown in two views with the corresponding model (Chain C in rainbow representation). Individual sections of the model are highlighted (insets 1 and 2). (B) Correlation values were computed between the map and model on a per amino acid basis, then averaged for all seven subunits in the asymmetric unit. Regions shown in blue had a strong correlation between the map and model, as opposed to the red regions, which had a weak correlation between the map and model. (C) VP2 of infectious bursal disease virus (PDB ID code 2DF7) shows a similar jellyroll fold with four strands on the top and bottom. A full view of the infections bursal disease virus is shown (Inset, Upper Left). (D) The gp10 model is colored based on Cα rmsd between gp10 and the corresponding domain of 2DF7. The largest differences are shown in red. Residues highlighted in green were not used in computing the rmsd. (Fig. S4, Fig. S5). In comparison with the previous model of gp7, the overall model structures are similar with a Cα rmsd of 2.33 Å between the two models over 54% (182 aa) of the protein’s structure. The new models have considerably improved fit to the density and are geometrically favorable (Table 1, Fig. S5A).

Several distinct structural features are clearly observed in the model of gp7 and are listed as follows (from N to C termini): N-arm, F-loop, E-loop, P-domain, the long α-helix, A-domain, and the C terminus (Fig. 2B, Fig. S5B). Superimposing the seven subunit models revealed nearly identical structures except for variations at the E-loop and N-arm (Fig. S5B). Furthermore, the penton subunit shows additional variation at the penton opening, with a loop in the A-domain folding inward, narrowing the opening at the penton.

To examine the agreement between each subunit model and the corresponding density, we visualized sidechain placement (Fig. 2A and 2C) and calculated a map–model correlation per amino acid as exemplified in one subunit (Fig. 2C, Fig. S5C). In all but three small regions, the average correlation with the density was above 0.70, although it should be noted that the main source of the low correlation was not in the overall mainchain trace but the absence of well-resolved sidechain density. The poorly correlated regions were primarily located in the A-domain and composed of one strand (155–159) and two loops (246–256 and 267–276) on either end of one helix (Fig. S5C). These regions are surface exposed and likely to be flexible.

The overall structure of gp7 resembles the canonical bacteriophage coat protein fold found in tailed dsDNA bacteriophages (Fig. 2D, Fig. S6) (35, 36). When superimposed on HK97 gp5, the prototypical example of such a fold, the overall structures are remarkably similar despite ε15 gp7 being 52 aa longer. Over the first ~145 aa, the two structures are nearly identical, at which point a domain swap in ε15 gp7 can be seen, where the β-strands in the P-domain occur before the two helices in the A-domain. In the A-domain, the five strands in the central β-sheet are oriented identically to the corresponding domain of the HK97 coat protein, although their ordering is different due to the aforementioned strand swap occurring just before the A-domain. Smaller differences can also be seen in some of the gp7 loops to accommodate the extra residues. Results from the CLICK webserver (37) revealed that 55% of the structure (153 aa) is aligned within 2.19 Å rmsd and has a topology score of 1.0 (i.e., identical topology; topology score of 0 corresponds to irrelevant structures). This high structural similarity with the HK97 bacteriophage coat protein fold provides an additional validation for our gp7 model.

Staple Protein, gp10. As mentioned, ε15 contains a second capsid protein, gp10, which binds to the outer surface of the virus capsid (Fig. 1B). Gp10s from neighboring capsomeres come together to form a dimer at the local twofold axes; visually each gp7 hexon is surrounded by 6 dimers of gp10. With respect to gp7, the gp10s reside adjacent to the A-domain and on top of both the E-loop and long helix from gp7 of neighboring asymmetric units. Due to this arrangement, we previously hypothesized that this protein acted as a molecular staple, performing similar stabilizing functions as the chainmail crosslinks in HK97. However, the exact structural mechanisms necessary for gp10 attachment and its contribution to capsid stability remained unclear.

In our new density map, gp10 appears to contain two sheets consisting of four β-strands each (Fig. 3). Although this map clearly shows β-strand separation, loops connecting the strands at one end of the two sheets were less resolved. A consensus fold model could not be directly established visually. To resolve this, multiple initial models for gp10 were generated using Pathwalker (28). From this pool of possible models, we selected, evaluated, and validated 16 possible topologies based on the density match. Each of these backbone models was converted to an all-atom model using REMO (38) and refined with ROSETTA (39) to evaluate a number of criteria, including density occupancy, model geometry, clashes, interactions, and structural similarities to known folds. Only six of the 16 accounted for all of the gp10 density. Three of these models were eliminated as they displayed clashes and poor protein–protein interfaces when combined with the gp7 model and fit back into the density map. Of the remaining three candidate folds, the top Pathwalker scoring model had the best fit to density. As such, this model was further refined to the density and chosen.

For this model, densities corresponding to aromatic and charged sidechains were generally visible. As with gp7, individual gp10 models in the asymmetric unit were iteratively refined and evaluated with standard X-ray crystallographic modeling tools (40) (Fig. 3, Fig. S7). The final refined model of gp10 clearly showed an eight-stranded jellyroll fold, a commonicosahedral virus structural fold found in viruses infecting organisms from all three domains of life (41–44). Using DALI (45), relevant structural similarity to other viruses containing the jellyroll fold were identified, with the top score belonging to the projecting (P) jellyroll domain of the

Table 1. Model statistics for 3,122 residues in 14 chains (one asymmetric unit) in e15

<table>
<thead>
<tr>
<th>Statistical measure</th>
<th>Score</th>
<th>MolProbity summary*</th>
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<td>60%</td>
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<tr>
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<td>21st percentile</td>
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<tr>
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<td>0%</td>
</tr>
<tr>
<td>Bad backbone angles</td>
<td>11</td>
<td>0.07%</td>
</tr>
</tbody>
</table>

*Percentiles based on structures at similar resolution in PDB.
infectious bursal disease virus VP2 subviral particle (PDB ID code 2DF7, Z-score = 6.3) (Fig. 3 C and D) (43, 46). Other high-scoring jellyroll folds from the DALI search included the coat protein from Satellite Panicum Mosaic Virus (PDB ID code 1STM), the B-domain of the large capsid protein in Cowpea mosaic virus (PDB ID code 2BFU), human rhinovirus 3 coat protein (PDB ID code 1W8X), and the projection domain of the infectious pancreatic necrosis virus VP2 protein (PDB ID code 3IDE). A list of relevant structural similarities is shown in Table S1; a diagram of the gp10 fold with labeled (B–G) strands is shown in comparison with selected jellyroll fold viral proteins (Fig. S8).

Although the infectious bursal disease virus VP2 subviral particle is not a bacteriophage, its crystal structure (PDB ID code 2DF7) (Fig. 3C) (46) offered significant insight into gp10. Visually, gp10 and the 2DF7 domain structure share a remarkable similarity in the overall fold despite a difference in length (141 aa in VP2 versus 111 aa in gp10) and no significant sequence homology; the topology of the two structures is identical (Click Score of 1.0). Over 87 residues in common (81.31%), the two structures have a Co rmsd of 2.44 Å with the majority of the structural difference occurring in the loops joining the β-strands (Fig. 3D). It is interesting to note that the observed jellyroll fold is found in capsids of various triangulation

Subunit–Subunit Interactions. To maintain capsid stability, ε15 forms an extensive network of interactions between gp7 and gp10, in both their own asymmetric units and across neighboring asymmetric units (Fig. 4A and B). Based on our models, 14 possible salt bridges were identified using PDBePISA (47), a web-based utility to interactively detect macromolecular interfaces (Table S2). Despite our limited map resolution, corresponding densities were present for some of these modeled salt bridges.

Although ε15 does not have the same covalent chainmail network as HK97 (5), these salt bridges likely play an equivalent role as evidenced of their location when drawing parallels to other bacteriophage capsids, such as P22 and HK97 (9, 12). Of particular interest is a series of three salt bridges found in the gp7 E-loop that tether the long extended loop within and between capsomeres (Table S2). As such, these salt bridges may stabilize the capsid by linking neighboring gp7 subunits in the mature virion, and possibly during capsid maturation.

For gp10, the two subunits, located at the twofold (Fig. 4C), interact along three β-strands (N63–T69, D73–E81, and R31–E37) and are dominated by hydrophobic residues. A number of additional subunit interactions are also seen at the gp7–gp10 interfaces. Like the capsid proteins in HK97 and PSSP-7, the N-arm of gp7 extends parallel to the P-domain; in other bacteriophages like P22 and T4, the N-arm is bent and runs underneath the long helix (Fig. S6). For ε15 gp7, the first ~20 aa of the N-arm takes an abrupt turn toward the capsid surface at Asn21 and forms a small helix and a short strand (blue in Fig. 4C). The first 10 residues of the N-arm from a gp7 subunit in an adjacent capsomere buttress one of the gp10s in the dimer along its exterior surface. Specifically, residues 3–6 of gp7 appear to form a strand parallel to strand C (residues 21–24) of the exposed B-strands in gp10; strands C, H, E, and F are surface exposed whereas strands B, I, D, and G are buried against the gp7 capsid layer (Fig. S8).

Based on the model, additional salt bridges between gp7 and gp10 are predicted with PISA. At the F-loop in gp7, Lys51 appears to form a salt bridge with Asp107 in the C terminus of gp10. Likewise, Lys2 in gp10 appears to coordinate with Ser10 of gp7 in an adjacent capsomere to anchor the N terminus of gp10. Additionally, Asp135 in gp7, located after the long α-helix, interacts with Arg41 of the gp10 model. Based on the location and potential interactions, it is conceivable that gp10 plays a key role in contributing to the capsid stability by joining neighboring capsomeres. This is further

![Fig. 4. ε15 subunit interactions. (A) An asymmetric unit for ε15 consists of seven gp7 (cyan) and seven gp10 (red) subunits. For illustration purposes, additional gp10 subunits are highlighted to illustrate the dimeric relationship. (B) One gp10 (bold red subunit) and one gp7 (bold cyan subunit) are shown with potential salt bridges shown as spheres (Table S2). (C) The gp10 dimer is highlighted in red and spans the twofold axis (Left). Four gp7 subunits are directly below a single gp10. The N-arm regions of gp7 are colored in dark blue and flank the outer strands of gp7. The color annotations for gp7 are the same as defined in Fig. 2B. Directly below the gp10 dimers are the gp7 E-loops (cyan). Rotating 90° shows a slice through the capsid with both gp10 termini positioned at the twofold interface, protruding down into the gp7 subunits (Center). A zoomed-in view of this region highlights the positioning of the N-arm of gp7 and the termini of gp10 (Right).](www.pnas.org/cgi/doi/10.1073/pnas.1309947110)
reinforced by the fact that even in 5 M guanidinium chloride (GuHCl), the gp7 and gp10 capsid shell remains intact (4).

Beyond stabilizing the mature ε15 virion, gp10 may play a larger role during capsid maturation. Previous examples of phage maturation in P22 and HK97 show a fairly substantial change in the N-arm conformation (9, 12, 48). In P22, it is postulated that the N-arm binds the C terminus of the scaffolding protein and that the conformational change in the N-arm contributes to scaffolding protein release. If gp10 is present during procapsid assembly, it raises the question if and how the N-arm of gp7 would simultaneously interact with the scaffolding protein (49) and gp10. Alternatively, gp10 may be incorporated in the mature phage, like in lambda and phage L (15, 50). Regardless of when gp10 is introduced into the capsid, the large conformational change during maturation would require dynamic, and possibly transitory, interactions to maintain capsid stability.

Interestingly, two protruding densities corresponding to the cytotoxic residues (C65) are prominent at the gp10-gp10 interface. Their proximity suggests a possible disulfide bond. However, neither connecting densities nor biochemical evidence exists to support such interaction in the mature form of ε15. Nevertheless, it is possible that if gp10 is present in the procapsid, this disulfide bond may serve as a temporary mechanism to stabilize the capsid during maturation until a full set of inter- and intracapsomeric interactions can be established in the mature virus. Such temporary disulfide bonds are not without precedence as early transient disulfide-dependent steps have been shown in influenza virus nucleocapsid (51) and Simian Virus 40 maturation (52). Additionally, transient interchain disulfide bonds in the P22 tailspike adhesion proteins have been seen (53).

**Evolutionary Lineage of ε15 Phage.** Tailed dsDNA bacteriophages have demonstrated a highly conserved structural fold as first defined in gp5 of HK97 (35) (Fig. S6). This fold has also been found to exist in the major capsid protein of Herpesvirus (36), suggesting that this fold, despite host, sequence, and structural variations, is a basic template that contributes to the ability of the major capsid proteins to form the primary proteinaceous shell that envelopes the genome (Fig. 2A, Fig. S6). The presence of the swapped strands in gp7 further supports the notion that this fold is a template that is conducive for virus formation. As such, the canonical fold found in the major capsid proteins of tailed dsDNA bacteriophages is essentially a scaffolding that can tolerate varied interactions, insertion domains, and decorating proteins, as well as sequence and structural permutations in forming a stable yet flexible capsid that can efficiently accommodate the viral genome.

Likewise, the jellyroll fold is seen in a wide variety of nonrelated viruses including animal viruses, plant viruses, and bacteriophages (Fig. S6) (41, 42, 44), as well as nonviral cellular proteins. Until our structure of ε15, no virus structure has been shown to contain both of these common viral folds. Because Salmonella, a bacterial pathogen in higher vertebrates, is the host of ε15 phage, it is conceivable that the ε15 evolved in the vertebrate host, which is a reservoir for both bacteriophage and other viruses. Genetic transfer and recombination within the microbial environment may have resulted in the emergence of ε15 phage.

**Materials and Methods**

**Cryo-EM.** Infectious bacteriophage ε15 was purified as described previously (3). The purified infectious phage particles were plunge frozen at liquid N2 temperature using a FEI Vitrobot device. The images were taken at −17 e Å2 dose in a JEM3200FS electron cryomicroscope operated at 300 kV and at liquid N2 specimen temperature. The imaging condition was set with spot size = 2, condenser aperture = 50 μm, and objective aperture = 120 μm. The in-column energy filter of this instrument was activated to remove electrons with larger than 25 eV energy loss. The images were recorded on Kodak SO163 films at 50,000x nominal magnification. A total of 1,309 films were digitized using Nikon Super CoolScan 9000 ED scanner at 6.35 μm/pixel and resulted in calibrated sampling of 1.19 A/pixel for the scanned images.

**Image Processing.** Individual particles (720 x 720 pixels) were first automatically selected using the ethan method (54) followed by manual screening using EMAN boxer program (53). A total of 54,161 particles were selected for initial processing. The selected particles within a micrograph were incoherently averaged to generate 2D power spectrum for contrast transfer function (CTF) parameter determination. CTF parameters were first automatically estimated (56) and then visually verified using the EMAN1 ctf program (55). The defocuses range from 0.5 to 2.5 μm. The dataset was divided into two data subsets for the following reconstruction steps.

The particle images were first binned 4x for initial model building and initial determination of orientation and center parameters. The initial model was built de novo by iterative refinement of a subset of 300 particles randomly selected from the half dataset with randomly assigned initial orientations. The initial orientations of all particles in each of the half datasets were determined using the EMAN1 projection matching program classbymat with an angular projection step size of 3 degrees (55). The orientations were then refined to higher accuracy using the program jalign, which is based on simplex optimization of matching between the particle image and model projections (19). The particle orientation parameters were then transferred to particles binned at 2x and ultimately to particles without binning for further refinements. In the last stage of refinement, magnification, astigmatism, and defocus parameters were also included (19). 3D maps with isosahedral symmetry enforcement were reconstructed using a newly developed program e2shell developed program j2d using EMAN2 library (57) and parallelized with message passing interface (MPI) to speed up the reconstruction process. These steps were then iterated until the refinement converged.

The map for each data subset was reconstructed from ~7,000 particles by removing particles with poor alignment scores and unstable alignment parameters. The resolution of the map was evaluated using the FSC. Only the isosahedral shell region was included in this FSC analysis by masking out the inner and outer background. As expected, the internal DNA densities using the icosahedral shell masks with a half width of 6 Å. The final map of the entire dataset was then built from ~14,000 particles by combining these two subsets of particles.

**Model Building.** For model building, each of the seven individual gp10 and gp7 subunits from one asymmetric subunit was cropped out of the full map using CHIMERA, from the University of California, San Francisco (UCSF) (58). For gp7, the previous structure (4) was used as the template to be developed program j2d using EMAN2 library (57) for each of the seven positions in the asymmetric unit.

For gp10, initial pseudoatom placement for pathwaying was done using e2segment3d.py, with minimum and maximum spacings of 2.1 and 4 Å, respectively, resulting in placement of 111 C atoms within the gp10 density map. Pseudos for gp7 (52) were obtained by taking the initial Cα backbone from the initial averaged density map with UCSF’s CHIMERA. For both gp7 and gp10, initial pseudoatom placement was varied (n = 1) and 100 models were calculated in e2pathwalker.py. A threshold was also set, corresponding to the smallest isosurface value at which the density for a single subunit remained connected. Pathwaying was performed with the default distance constraints using the LKH TSP solver (20). The corresponding sequences were then threaded onto the corresponding models in both the forward and reverse directions.

For gp7, a representative model for each of the two possible fold classes was fit into the ε15 density map at each of the seven positions in the asymmetric unit and evaluated for fit to density. The best model was visually selected and converted to an all-atom model using REMO, a webserver that transforms Cα backbone models into all-atom models (38).

For gp10, we identified 16 potential model types from the gallery of Pathwalker models. Each was loaded into GORGON (29) and was interactively adjusted to correct any bond distance and clash problems. These models were then converted to all-atom models using REMO and density refined with ROSETTA (39). Initial evaluation of the models considered (i) geometry, (ii) fit to density, (iii) density occupancy, and (iv) interface between neighboring gp10s and gp7s. Additionally, the models were submitted to the DALI to identify possible structural homologs (45). A final model was selected based on fit to density, model geometry, and structural similarity to known homologs.

Using these all-atom models, slightly more than an asymmetric unit (12 gp10s and 7 gp7s) was assembled in the context of the entire capsid
structure. Fitting of these subunits was done manually and refined with the "Fit in Map" option in UCSF's CHIMERA (58). Individual subunits were iteratively refined with ROSETTA (39) and COOT (32) to optimize sidechain positions, improve density agreement, and improve the dihedral angles, as well as eliminate subunit clashes. To optimally refine an r15:1 entire asymmetric unit using best practices established in X-ray crystallography, PHENIX (33) and COOT (32) were used iteratively.

PHENIX refinement was implemented using phenix.refine (60). To use our density map data, a python script was written, converting the phases of the map to HKL coefficients, which could then be used during the refinement process. Refinement was done by applying secondary structure restraints, NCS restraints, and mapstereochemistry weight restraints. Initial rounds of refinement used Ramachandran restraints and a strong geometry weight. To minimize potential overfitting, geometric and density statistics (Table 1) were monitored during the various rounds of refinements. Later rounds of refinement were done maintaining the Ramachandran restraints, relaxed geometry restraints, and NCS restraints. A final simulated annealing COOT was used to manually remove clashes between refinement iterations. With one asymmetric unit refined, neighboring subunits to the asymmetric unit were added and thus a large complex was refined ensuring that asymmetric unit interfaces could be properly interpreted. An entire virus model was then constructed by applying isocahedral symmetry to the asymmetric unit model.

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