Enterotoxigenic *Escherichia coli* (ETEC) is a bacterial pathogen that causes diarrhea in children and travelers in developing countries. ETEC adheres to host epithelial cells in the small intestine via a variety of different pili. The CS1 pilus is a prototype for a family of related pili, including the CFA/I pili, present on ETEC and other Gram-negative bacterial pathogens. These pili are assembled by an outer membrane usher protein that catalyzes subunit polymerization via donor strand complementation, in which the N terminus of each incoming pilin subunit fits into a hydrophobic groove in the terminal subunit, completing a β-sheet. Here we determined a crystal structure of the CS1 major pilin subunit, CooA, to a 1.6-Å resolution. CooA is a globular protein with an Ig fold and is similar in structure to the CFA/I major pilin CfaB. We determined three distinct negative-stain electron microscopic reconstructions of the CS1 pilus and generated pseudoatomic-resolution pilus structures using the CooA crystal structure. CS1 pili adopt multiple structural states with differences in subunit orientations and packing. We propose that the structural perturbations are accommodated by flexibility in the N-terminal donor strand of CooA and by plasticity in interactions between exposed flexible loops on adjacent subunits. Our results suggest that CS1 and other pili of this class are extensible filaments that can be stretched in response to mechanical stress encountered during colonization.
Structure of the ETEC CS1 Pilus

TABLE 1  Bacterial strains, plasmids, and primers

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Description</th>
<th>Source/reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacterial strains</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. coli MC4100/pEU494</td>
<td>F− araD139 Δ(argF-lac)169 leuA14− thyB501 Δ(fruK-yeiR)725 (fusA25) relA1 rpsL150 (Stb') thyB501 deoC1 ptsF25 rbs22 Δ(fimB-fimE)632::(IS1) pEU494</td>
<td>24, 51</td>
</tr>
<tr>
<td>ETEC LMC10/pEU2030</td>
<td>ETEC lacking the plasmid encoding the toxins and the positive regulator of coo</td>
<td>35</td>
</tr>
<tr>
<td>E. coli RosettaBlue</td>
<td></td>
<td>Novagen</td>
</tr>
<tr>
<td>Plasmids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pET-15b</td>
<td>T7 promoter, Amp'</td>
<td>Novagen</td>
</tr>
<tr>
<td>pEU494</td>
<td>pUC18 encoding ns under its own promoter, Amp'</td>
<td>52</td>
</tr>
<tr>
<td>pEU2030</td>
<td></td>
<td>This study</td>
</tr>
<tr>
<td>pET-15b-CooBA</td>
<td>pET-15b, cooB-cooA gene segment in NdeI/BamHI sites</td>
<td>This study</td>
</tr>
<tr>
<td>pETDuet-CooB1</td>
<td>pETDuet, cooB inserted into BamHI and Sall sites</td>
<td>This study</td>
</tr>
<tr>
<td>pETDuet-CooB1-Ntd11CooA2</td>
<td>pETDuet-CooB1 with cooA gene encoding residues 12−148 (cooAΔN11) inserted into NdeI and Xhol sites</td>
<td>This study</td>
</tr>
<tr>
<td>pET-28b-Ntd11CooA-dsc13</td>
<td>pET-28b with Ntd11CooA-dsc13 inserted into NdeI and Xhol sites</td>
<td>This study</td>
</tr>
<tr>
<td>Primers</td>
<td></td>
<td></td>
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<tr>
<td>ETEC-CooBA-IPCR</td>
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<td>ETEC-CooBA-rPCR</td>
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<tr>
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<td>CooA-R</td>
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<td>This study</td>
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<tr>
<td>Ntd11CooA-dsc13-R</td>
<td>GCCTCGAGGATATGCGAAGAATTATTTTGAAGTTATGACTATTTC</td>
<td>This study</td>
</tr>
</tbody>
</table>

Base of the pilus filament, which grows out through the usher pore into the extracellular space.

Electron microscopy and three-dimensional (3D) image processing of P pilus and type I pilus from UPEC and CFA/I pilus from ETEC, combined with X-ray crystallography of the pilin subunits and subunit complexes, reveal a conserved filament architecture, where subunits are associated in a head-to-tail manner, with the Nte on one subunit, n + 1, inserted into the hydrophobic groove of the adjacent subunit, n (7, 18–20). Subunits align with their cylindrical axes almost perpendicular to the filament axis and wrap around the filament with a rotation of ~110° and a rise of ~8 Å per pilin monomer for a right-handed 1-start helix. The filament diameter is ~8 nm, and an ~25 Å channel runs through the filament. These data together indicate that although the class 1 and 5 pili are dissimilar in amino acid sequence, their subunit structures, filament architectures, and assembly mechanisms are very similar. CFA/I filaments often possess a thin fibrillar structure at their tips, which appears to represent extended CFA/I filaments. The crystal structure of recombinantly expressed ETEC CooA together with a 3D reconstruction of the CS1 pilus filament. CooA, like CfaB and the class 1 pilins, undergoes donor strand complementation. The CS1 filaments exhibit structural polymorphism that likely relates to pilus extensibility.

MATERIALS AND METHODS

Bacterial strains and plasmids. CS1-producing E. coli strains were a gift from June Scott, Emory University. E. coli MC4100/pEU494 (24) was used to PCR amplify the cooA gene for recombinant expression. ETEC strain LMC10/pEU2030 was used to express CS1 pilin for electron microscopy (EM) analysis. All bacterial strains, plasmids, and oligonucleotides are listed in Table 1. Antibiotic concentrations were as follows: kanamycin (Kan), 45 μg/ml; ampicillin (Amp), 100 μg/ml.

Cloning and expression of CooAΔN. The cooB-cooA genes were PCR amplified from pEU494 (24) using primers ETEC-CooBA-IPCR and ETEC-CooBA-rPCR (Table 1). The PCR product was ligated into the NdeI and BamHI restriction sites of the pET-15b vector (Novagen) to produce pET-15b-CooBA. The cooA gene fragment encoding residues 12 to 148 of the mature CooA protein (cooAΔN11) was subcloned from this vector using primers Ntd11CooA-dsc13-F and CooA-R and ligated into the pETDuet-CooB1 vector at the NdeI and Xhol sites, which had the cooB gene inserted into the BamHI and Sall sites, to produce pETDuet-CooB1-Ntd11CooA2. Primers Ntd11CooA-dsc13-F and Ntd11CooA-dsc13-R were used to PCR amplify cooAΔN11 from pETDuet-CooB1-Ntd11CooA2 and add a gene segment to its 3' end encoding a flexible linker, DNKQ, followed by the 13 N-terminal amino acids of mature CooA. This new gene fragment, Ntd11CooA-dsc13, was inserted into the NdeI and Xhol restriction sites of pET-28b to produce pET-28b-Ntd11CooA-dsc13 for expression of a soluble CooA protein lacking its N-terminal 11 residues, with its 13 N-terminal residues fused to its C terminus via a DNKQ linker (donor-strand-complemented CooA [CooAΔN]). CooAΔN was expressed with an N-terminal hexahistidine tag for the purposes of protein purification.
E. coli RosettaBlue (Novagen) was transformed with pET-28b-Ntd11CooA-dsc13. Cells were grown overnight in Luria broth (LB)–Kan, and 100 ml of the overnight cultures was used to inoculate 1 liter LB-Kan. Cells were grown at 37°C to an optical density of 0.75 at a 600-nm wavelength (OD<sub>600</sub>) at which time isopropyl-β-D-thiogalactopyranoside (IPTG) was added to a final concentration of 1 mM to induce CooA<sub>dsc</sub> expression. Cells were grown for an additional 3.5 h at 30°C with vigorous shaking and harvested by centrifugation at 5,000 × g for 30 min at 4°C. Cells were lysed by incubation in lysis buffer (500 mM NaCl, 50 mM phosphate buffer [pH 7.4], and 40 mM imidazole with 0.2 mg/ml lysozyme) followed by sonication 3 times (2 min each) at 45% amplitude. The lysed cells were centrifuged at 25,000 × g for 40 min at 4°C to remove cell debris, and CooA<sub>dsc</sub> was isolated from the supernatant using a nickel-nitrilotriacetic acid (NTA) column (GE Healthcare). CooA<sub>dsc</sub> was eluted using an imidazol gradient. Fractions containing CooA<sub>dsc</sub> were pooled and concentrated on an Amicon stirred-cell concentrator (Millipore) and exchanged into CooA buffer (100 mM NaCl and 20 mM Tris-HCl, pH 7.5). CooA<sub>dsc</sub> was further purified on a Sephacryl S-100 HR size exclusion column (GE Healthcare) and concentrated to 15 mg/ml for crystallization.

CooA<sub>dsc</sub> crystallization, data collection, and structure determination. CooA<sub>dsc</sub> was crystallized by the hanging drop vapor diffusion method in 1.0 M sodium citrate and 0.1 M imidazole, pH 8.0, using 2 μl protein solution and 2 μl reservoir buffer. Crystals were flash-frozen and stored in liquid nitrogen in reservoir buffer with 25% glycerol as a cryoprotectant. X-ray diffraction data were collected at the Stanford Synchrotron Radiation Lightsource (SSRL) Beamline 11-1. The raw data were processed for Lorentz and polarization corrections and integrated by using the XDS software program, followed by scaling with the program XSCALE (25). The crystal belonged to the triclinic space group P1, with two molecules in the unit cell. The solvent content was 35% as determined from the Matthews coefficient (26). The structure was solved by molecular replacement using the CfaB crystal structure as a search model (PDB code 3F83) (7). Nonidentical residues in CfaB were changed to Ser, Ala, or Gly in the software program COOT. An initial search by using the program Phaser (27) yielded two unambiguous solutions with fairly high Z scores (100 and 14.9). Several iterative cycles of rigid-body refinement with tight NCS (noncrystallographic symmetry) restraint and density modification procedures were performed using the software programs REFMAC5 (28) and DM (29), respectively. The electron density map was interpretable and clearly showed density for many missing side chains, which were built using COOT (30). Many iterative cycles of model building and restrained refinement by the software program ARP/wARP (31) improved the model and revealed density for most of the side chains as well as the missing portions of the main chain. Water oxygens were located using ARP/wARP, and their positions were manually checked in COOT with difference map output by REFMAC5 and the composite annealed omit map calculated using the software program CNS (32). Further refinement using TLS (thermal liberation screw-motion) gave an R<sub>free</sub> value of 0.222 and an R<sub>free</sub> value of 0.251. No electron density was observed for the His tag or linker. The model was validated by the software programs PROCHECK (33) and MOLPROBITY (34). Data collection and refinement statistics are shown in Table 2.

### Purification of CS1 filaments
CS1 pili were purified from ETEC LMC10 cells carrying the pEU2030 plasmid (35). ETEC LMC10 has lost the plasmid carrying rns, which encodes the positive transcriptional regulator of coo operon expression. pEU2030 encodes rns under its own promoter. Cells were grown for 16 h in LB-Amp with shaking at 37°C. Cells were harvested by centrifugation at 8,000 × g for 30 min at 4°C. The cell pellet was resuspended in phosphate-buffered saline (PBS) (pH 7.4; 137 mM NaCl, 2.7 mM KCl, 10 mM Na<sub>2</sub>HPO<sub>4</sub>, and 2 mM KH<sub>2</sub>PO<sub>4</sub>) with 10 mM EDTA. CS1 pili were sheared from the cells by passing the mixture through a 25-gauge needle four times followed by vigorous vortexing for 1 min. Cell debris was removed by centrifugation at 8,000 × g for 30 min at 4°C. Ammonium sulfate (20% [wt/vol]) was added to the supernatant and incubated at 4°C for 2 h to precipitate CS1 filaments, which were isolated by centrifugation. The CS1 pellet was resuspended in PBS-EDTA and dialyzed exhaustively in the same buffer to remove residual ammonium sulfate. Purity was assessed by SDS-PAGE analysis (see Fig. S1 in the supplemental material), and the major 15-kDa CooA band was excised from the gel and analyzed by tandem mass spectrometry at the University of Victoria (UVic) Genome BC Proteomics Centre to confirm its identity.

### Electron microscopy and image processing
CS1 pili in PBS buffer were diluted 20× with distilled water. Five microliters of sample was applied to glow-discharged continuous carbon-coated grids, negatively stained with 2% uranyl acetate, and imaged on an FEI Tecnai T12 microscope at 80 keV with a nominal magnification of ×30,000. Images were recorded on Kodak SO163 film and digitized using a Nikon Coolpix 8000 scanner at a raster of 4.16 Å per pixel. The EMAN software package (36) was used to extract filament images from micrographs, and the SPIDER software package (37) was used for most image processing.

Segments (100 × 100 pixels each; n = 10,443) of the CS1 filaments

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### Table 2 Crystallographic data collection and refinement statistics

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Name or value(s) for CooA&lt;sub&gt;dsc&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
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<td>Data collection</td>
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<tr>
<td>Beamline</td>
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<td>Wavelength (Å)</td>
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</tr>
<tr>
<td>Space group</td>
<td>P1</td>
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<td>Cell a, b, c (Å)</td>
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</tr>
<tr>
<td>Cell α, β, γ (°)</td>
<td>83.72, 89.95, 74.58</td>
</tr>
<tr>
<td>Resolution (Å)</td>
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</tr>
<tr>
<td>Completeness (%)&lt;sup&gt;a&lt;/sup&gt;</td>
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</tr>
<tr>
<td>No. of observed reflections</td>
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</tr>
<tr>
<td>No. of unique reflections</td>
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</tr>
<tr>
<td>R&lt;sub&gt;free&lt;/sub&gt; (%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.025 (0.117)</td>
</tr>
<tr>
<td>I/σ(I)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>34.1 (10.8)</td>
</tr>
<tr>
<td>Redundancy</td>
<td>3.9 (4.0)</td>
</tr>
<tr>
<td>Wilson B value (Å&lt;sup&gt;2&lt;/sup&gt;)</td>
<td>24.5</td>
</tr>
<tr>
<td>Mosaicity (°)</td>
<td>0.2</td>
</tr>
</tbody>
</table>

**Refinement statistics**

- Resolution limits (Å): 19.7–1.6
- Molecules/unit cell: 2
- No. of reflections used: 31,826
- R<sub>cryst</sub> (%)<sup>a</sup>: 22.2
- R<sub>free</sub> (%)<sup>b</sup>: 25.1
- No. of atoms: 884
- Protein: 2,295
- Ligands: 6
- Water: 156
- Avg B factor (Å<sup>2</sup>): 21.6
- Chain A: 22.0
- Chain B: 22.0
- Ligands: 29.7
- Imidazole: 29.7
- Sodium ion: 21.1
- Water oxygen: 29.0
- RMSD bond angles (°): 1.37
- RMSD bond lengths (Å): 0.008
- Ramachandron plot
  - Favored (%): 99.0
  - Allowed (%): 1.0

<sup>a</sup> Overall/final shell.
<sup>b</sup> R<sub>free</sub> is the unweighted R value on I between symmetry mates.
<sup>c</sup> I/σ(I) is the cross-validation R factor for 5% of reflections against which the model was not refined.

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RESULTS AND DISCUSSION

X-ray crystal structure of CooAdsc. The CooA pilin subunit was expressed as a soluble recombinant protein in which the N-terminal 11 residues were removed and the Nte (residues 1 to 13) was appended to the CooA C terminus (Fig. 1A) with an intervening 4-residue Asp-Asn-Lys-Gln linker sequence (the DNKQ linker) (Fig. 1B). This strategy has been used to produce soluble subunits and complexes for CfaB and pilins in the chaperone-usher pilus family to prevent misfolding and spontaneous subunit polymerization (7, 8, 40, 41). The crystal structure of recombinant donor-strand-complemented CooA, CooAdsc, containing an N-terminal hexahistidine tag was solved to a 1.6 Å resolution by molecular replacement using CfaBdsc (3F84 [7]) as a model (Table 2).

CooAdsc is a single-domain protein that forms a 7-stranded β-sandwich with an Ig fold (Fig. 1C to E). CooAdsc crystallized as a dimer with the two monomers packed together by hydrogen bonding between the second sheet of the β-sandwich in each subunit. The monomers cross over each other at an ~45° angle and are essentially identical in structure (see Fig. S2A in the supplemental material), but the N- and C-terminal residues Asp12 and Pro165, respectively, are resolved only for chain A. The first β-sheet of the β-sandwich is antiparallel, comprising strands A1, A2, C, and D (Fig. 1C). The second β-sheet comprises strands A2, G, F, and C. Strands G, F, and C are antiparallel, whereas A2, which lies at the edge of the second β-sheet, runs parallel to G. Several of the β-strands span both β-sheets and are interrupted by loops that protrude from the surface of the protein. The N- and C-terminal halves of strand A, A1 and A2, are part of β-sheets 1 and 2, respectively, and are connected by an 8-residue loop. A similar arrangement is observed for strand D on the opposite side of the protein, with D1 being part of β-sheet 1 and D2 being part of β-sheet 2, connected by a 9-residue loop. D2 is followed by a single helical turn and then D3, which rejoins the first β-sheet. Strand C also has a short loop interruption, with its N- and C-terminal halves being part of β-sheet 2 but C2 also forming canonical backbone hydrogen bonds with strand D1 of β-sheet 1. The connectivity and orientation of the β-strands are typical of a c-type Ig fold (42), but the protein has an overall cylindrical shape and resembles a β-barrel due to the curvature of the β-sheets and their continuity at strand C2. The loops within the β-strands, as well as those at the ends of the β-sheets, dominate the surface of the protein. These loops are considerably more flexible than the β-strands, as shown by their higher B factors (see Fig. S2A), although they are constrained by extensive interactions with adjacent molecules in the closely packed crystal lattice, which has a solvent content of only 35%.
FIG 1 Crystal structure of ETEC CooA<sub>ΔCoo</sub>. (A) Amino acid sequence alignment of CooA and CfaB. Secondary structure elements of CooA are indicated above its sequence. Red background, identical residues; yellow background, similar residues. (B) Schematic representation of wild-type CooA (top) and the CooA<sub>ΔCoo</sub> construct (bottom). The sequence shown with orange background represents the Nte, which has been removed from the N terminus and attached to the C terminus. (C) Schematic of the secondary structure of CooA<sub>ΔCoo</sub>. Green β-strands belong to the first β-sheet and blue and orange strands belong to the second β-sheet in the Ig fold. The orange G strand is the Nte, which has been fused to the C terminus. (D) Ribbon representation of the 1.6-Å-resolution CooA<sub>ΔCoo</sub> crystal structure. (E) Space-filling representation of CooA with the Nte shown in stick representation. Carbon atoms are colored orange in the Nte, yellow in the DNKQ linker, and gray in the remainder of the protein. Oxygen atoms are red, and nitrogens are blue. Residues in the Nte are numbered as in native mature CooA, where the Nte of subunit <i>ni</i> (not shown) is donated to subunit <i>n</i> (shown). The Nte fits into the hydrophobic groove with hydrophobic side chains at positions P2 to P5, indicated in parentheses, filling corresponding pockets in the groove. (F) CS1 assembly scheme showing donor strand complementation between adjacent CooA subunits. Each subunit has a different color to illustrate donor strand complementation with an adjacent subunit in the filament.
The conformation of the loop between strand A2 and B differs slightly between the two molecules in the crystallographic dimer, with the Cα atoms of Ala36 at the tip of this loop being ~2.2 Å apart. The CooA structure is very similar to that of the CFA/I major pilin, CfaB, as expected from the high sequence identity (53%) (Fig. 1A; see also Fig. S2B in the supplemental material). The only significant differences in backbone structure occur at the ends of the proteins: the D3-E1 loop, which is a -hairpin in CooA and an irregular loop in CfaB, and divergent conformations for the E2-F loop.

Strand G of the second -sheet (residues 153 to 165) is the Nte in native mature CooA that has been fused to the C terminus in CooAdsc (Fig. 1C to E). Strand G fills a hydrophobic groove between strands A2 and F to complete the second -sheet. Strand G forms canonical backbone hydrogen bonds with strands A2 and F and interacts with A1 in sheet 1 via its side chains. Although strand G is the C-terminal strand of CooAdsc, its residues are numbered in Fig. 1E according to their position at the N terminus of the protein, since this strand would originate with an adjacent subunit (n+1) in the pilus filament via donor strand complementation (Fig. 1F). As is seen for pilins in the chaperone-usher family, four alternating hydrophobic side chains at positions P2 to P5 fit into their respective P2 to P5 pockets in the hydrophobic groove. In addition, Lys3 forms a salt bridge with Asp16 in strand A1. Clear electron density is seen for residues 1 to 12 of the Nte (residues 153 to 164 in CooAdsc) in chain A, but only the backbone atoms are apparent for Pro13 (Pro165). Pro13 is conserved among the class 5 fimbrial family and was proposed to form a flexible hinge between subunits in CFA/I due to cis-trans isomerization (7). Pro13 is also present at the N terminus in our recombinant CooAdsc; however, neither the electron density at the N terminus nor that at the C terminus is sufficient to distinguish between the cis and trans proline conformations. The next residue in native CooA, Thr14, is the first residue in β-strand A1 in the body of the pilin subunit. Since residues 1 to 11 of the Nte for subunit n+1 form the donor G strand for subunit n in the pilus filament and residues 14 to 19 form the first β-strand of subunit n+1, the linker between the subunits is comprised of only two residues, Asp12 and Pro13. Thus, the donor strand complementation interaction requires the ends of the CooA subunits to be very close together in a head-to-tail interaction in the CS1 filament (Fig. 1F), as was seen in the crystal structure of the ternary complex between two CfaB subunits and the CfaE chaperone (7).

**3D reconstruction of the CS1 filament.** We used quick-freeze/deep-etch metal shadowing EM to visualize the surface features of the CS1 pilus (Fig. 2A). F-actin filaments were used as a control because they possess a left-handed 1-start helix that gives rise to a sharp layer line in the power spectrum at 1/(59 Å) (Fig. 2A, inset, left). An averaged power spectrum generated from 174 nonoverlapping segments of the CS1 filaments (Fig. 2A, inset, right) revealed a layer line at 1/(145 Å) arising from a left-handed helix. To further determine the helical symmetry of the CS1 pilus, we calculated power spectra for 10,443 overlapping segments from negatively stained CS1 filaments and added these together (Fig. 2B). This method is independent of any alignment needed to average together images and is thus unbiased. The resulting power spectrum shows four layer lines. The layer line at ~1/(27.6 Å) arose from a 1-start helix, based upon the diameter of the filaments (~75 Å) and the distance of the peak from the meridian (Fig. 2B, inset). The mass per unit length was independently determined to be 1.76 kDa/Å by STEM (see Fig. S3 in the supplemental material). Taking into account the molecular mass of CooA (~15.2 kDa), the average rise per subunit was determined to be ~8.6 Å. This gives a twist of ~3.2 subunits per turn of the 1-start helix. Index-
ing of the remaining layer lines in the power spectrum shown in Fig. 2B suggested that the reflection at 1/(145 Å) arose from the 3-start helix, while the layer line at 1/(23.2 Å) originated with the 2-start helix. Since metal shadowing unambiguously showed that the 3-start helix is left-handed, the 1-start helix must be right-handed. Thus, the average symmetry of the CS1 pili was determined to be a 113.5° right-handed rotation around the helical axis with an 8.7-Å rise per subunit. These helical parameters are close to those of CFA/I pili (113.6°/8.3 Å [19]).

We used IHRSR (38) to determine the CS1 pilus structure. IHRSR was run using 10,443 overlapping pilus segments with three starting symmetries distributed about the mean helical parameters determined from the power spectra and the STEM data (Fig. 2B; see also Fig. S3 in the supplemental material). The IHRSR run converged to the same solution of 113.1° and 8.7 Å from the two starting values of 113°/8.7 Å and 114°/7.7 Å but failed to converge to a solution from a starting symmetry of 112°/9.7 Å (Fig. 3A). In our experience, when two or more closely related starting symmetries do not converge to the same solution, it indicates structural heterogeneity for the filaments. Thus, a single-particle approach was used to sort CS1 pilus segments into three more homogeneous groups (see Materials and Methods). IHRSR was then run for each group using a starting symmetry of 113° and 8.7 Å. Each group converged to a stable solution: the symmetries determined for group 1, the largest group, which accounts for ~58% of all segments, and for group 2 (~20%) are very similar, 113.2°/8.7 Å and 113.1°/8.6 Å, respectively, whereas group 3 (~22%) has a symmetry of 111.9°/10.0 Å (Fig. 3B). Importantly, all three symmetries were randomly distributed among the CS1 filament particles extracted for image analysis, indicating that the heterogeneity exists within the filaments rather than between them.

**Pseudoatomic-resolution CS1 filament structures.** The three independent CS1 reconstructions generated using IHRSR are shown in Fig. 4A. Each reconstruction was estimated to be at ~20-Å resolution using an FSC of 0.5 for two completely independent reconstructions generated from different starting symmetries (see Materials and Methods). Since the best measure of resolution is the quality of the map itself, we filtered our high-resolution structure of the CooA_\text{disc} pilin to 20 Å, docked it into
FIG 4 Characterization of three CS1 structural groups. (A) 3D reconstructions of the three structural groups (gray transparent surfaces) with the corresponding pseudoatomic-resolution models (colored ribbons). (B) Pseudoatomic models filtered to 20-Å resolution (cyan surfaces) superimposed on the 3D reconstructions (gray transparent surfaces). (C) Reference-free 2D averages of each structural group are compared with the projections of the corresponding 3D reconstructions.

Each of the three 3D reconstructions, and generated pseudoatomic-resolution filament models by imposing the symmetry parameters onto the subunit (Fig. 4A). An important fitting constraint was the requirement for a head-to-tail interaction between each subunit such that donor strand complementation could occur. To test the validity of the CS1 filament models, each model was filtered to a 20-Å resolution and superimposed on the corresponding 3D reconstruction (Fig. 4B). The FSC for the models and reconstructions (see Fig. S4 in the supplemental material) confirm the resolution estimates and provide confidence in the 3D reconstructions of the three CS1 structural classes. Since each reconstruction was generated starting from a featureless solid cylinder in the IHRSR procedure, all the details in each reconstruction that emerge during the iterative process arose from the information contained in the pilus images. To exclude any bias from the projections used for the sorting of the CS1 segments into the three groups (see Materials and Methods), we calculated reference-free 2D averages for each group and compared those with the corresponding projections from the 3D reconstructions (Fig. 4C). An excellent match between the 2D averages and the projections of the actual reconstructions excluded the possibility that the 3D reconstructions are model biased.

An important feature of the class 1 and class 5 pili from Gram-negative bacteria is the head-to-tail connectivity between pilin subunits $n+1$ and $n$ along the 1-start helix of the filament as a result of donor strand complementation (6). Bullitt and colleagues found that the head-to-tail interactions between the CfaB subunits of the 1-start helix in CFA/I were more extensive than those between subunits on consecutive turns or “layers” of that helix (19). Our structural analysis of the CS1 pilus shows that the extent of the head-to-tail interactions depends upon the structural state of the filament. To compare the connectivity along the 1-start helix, we contoured the EM reconstructions of the three structural groups at a higher threshold (Fig. 5A) than that required to accommodate the crystal structure of the CooA pilin (as shown in Fig. 4B). The group 1 reconstruction, which represents the majority of CS1 segments, possesses a prominent bridge of density between subunits along the 1-start helix (Fig. 5A, dashed oval). This connectivity is also apparent in the group 2 reconstruction, though the density is weaker. Only a narrow bridge of density connects subunits in the 1-start helix in the group 3 reconstruction. The extent of the end-to-end connectivity correlates with the distance between the C terminus of the Nte of CooA subunit $n$ (Pro13/Pro165, end of orange strand, Fig. 5) and the N terminus of subunit $n+1$ (Thr14, black sphere), which are joined by a peptide bond in native CooA. In the CS1 models 1 and 2, these distances are $\sim 8\AA$ and $\sim 5\AA$, respectively, which would allow the N terminus of subunit $n+1$ to complement the $\beta$-sheet of subunit $n$ without major conformational changes in the CooA structure. However, in filament model 3, these residues are $\sim 19\AA$ apart. Head-to-tail connectivity could be accommodated in this filament structure if a small conformational change occurs at the CooA N terminus such that $\beta$-strand A1 is released from hydrogen bonding with strand B and side chain interactions with strand G (Fig. 1C and D; see also Fig. S5 in the supplemental material), allowing it to extend far enough for the Nte to reach its neighboring subunit in the 1-start helix. Support for such a mechanism is provided by the crystal structure of EcpA, the major pilin from the E. coli common pilus (ECP) (43). EcpA is similar in structure to CooA and crystallized as a domain-swapped dimer, with the A1 $\beta$-strands crossing over to hydrogen bond with strand B of their partner monomer. The exposed loop connecting strands A1 and A2 is present in CfaB and other pilins belonging to the chaperone-usher family, and thus flexibility in the N terminus may be a common feature of these pili.

Despite the relatively low resolution of the reconstructions, the fitting of the CooA$_{adsc}$ structure into the EM maps was unambiguous. Surprisingly, small differences in the helical symmetries of the three reconstructions are accompanied by significant structural perturbations. Reconstructions 1 and 2 have similar helical symmetries but differ in orientations of the subunits, where CooA is rotated $\sim 25^\circ$ about its long axis in model 2 relative to its orientation in model 1 (Fig. 5B). Reconstruction 3, with its larger rise (10.0 Å), is more extended than reconstructions 1 and 2 (8.7 Å and 8.6 Å, respectively). CooA subunits are held together in the CS1 filament by head-to-tail donor strand complementation ($n$ to $n+1$) and by interactions between the subunit bodies, both head-
to-tail \((n \rightarrow n+1)\) and along the filament axis between each successive turn of the 1-start helix (i.e., between \(n+1\) and \(n-2\) in Fig. 5B). Only minor clashes occur between subunits in each of the three models, mainly between side chains in loops. This is not surprising for models derived from rigid body fits, and these clashes could be eliminated with minor adjustments in side- and main-chain torsion angles. In support of this idea, the A2-B loop at the “head” of CooA subunit \(n\) is closely associated with the tail of subunit \(n+1\) in models 1 and 2; this loop has relatively high temperature factors, and its conformation differs slightly between the A and B chains of the CooA crystallographic dimer (see Fig. S2A in the supplemental material), suggesting flexibility in this region. Interestingly, despite the fact that many of the surface loops in neighboring subunits contact each other in both the filament models and the crystallographic lattice, these contacts are not the same. This is not surprising given that there is no head-to-tail constraint for subunits in the crystal lattice.

The filament symmetries determined here for CS1, 112°–113° rotation and \(\sim 9\-Å\) rise, are comparable to those of CFA/I class 5 pili (113.6°/8.3 Å [19]) and those of UPEC P pili and type I pili from the class 1 family (109.6°/7.6 Å for P pili [18]; 106.7/7.2 Å for type I pili [20]). The orientation of the CooA subunits in filament models 1 and 2 is similar to that of CfaB in the CFA/I model (7), while model 3 differs significantly.

The origin of the filament polymorphism is not clear, but it is not introduced by negative staining for EM, which may actually limit the observed heterogeneity. In an effort to obtain a higher-resolution structure, we imaged and processed frozen hydrated CS1 pili in order to preserve the filaments in a near-native state. Surprisingly, the averaged power spectra for CS1 pili in vitrified ice showed a single layer line representing the left-handed 3-start helix, with other layer lines being weakened or absent altogether (see Fig. S6 in the supplemental material). The loss of higher-resolution data is indicative of structural heterogeneity in the frozen hydrated sample. It is possible that structural perturbations were introduced into the filaments during purification, where

![FIG 5](https://example.com/fig5.png)
they were mechanically sheared from *E. coli* cells. Yet *Vibrio cholerae* type IV pili are heterogeneous even without being subjected to mechanical shearing (44), and *Neisseria gonorrhoeae* type IV pili are not polymorphic although they were prepared by shearing (45). Interestingly, the *N. gonorrhoeae* type IV pili were shown to undergo a conformational change that exposed new epitopes when force was applied (46). Filament polymorphism implies structural plasticity, which would allow the pili to maintain adherence to the host epithelium when subjected to high flow forces. The ability of the filaments to alter their subunit orientations and contacts may be explained by having (i) most contacts mediated by flexible surface-exposed loops and (ii) an extensile N terminus.

In addition to the filament polymorphism, we observed a more dramatic structural perturbation of the CS1 pili that may provide further flexibility for the filaments. A fraction of the pili in the quick-freeze and deep-etch images appear to be partially unraveled at their ends, as well as within the filaments (Fig. 6). These thin fibrillar segments likely represent sections where the axial interactions between CooA subunits (i.e., between *n* + 1 and *n* − 2 in Fig. 5B) are disrupted but subunits remain associated via donor strand complementation. Pili unraveling has also been reported for CFA/I, P pili, and type I pili (18, 20, 47, 48) and may thus represent an important adaptation to stress for pili in the chaperone-usher family. Force measurements of CFA/I pili using optical tweezers showed that unraveling occurs at relatively low forces (49) compared to the forces required to unravel P pili and type I pili (50). These differences were explained by fewer layer-to-layer interactions between subunits in consecutive turns of the CFA/I filaments than was the case for those for P pili and type I pili (49). The close similarity between the CS1 and CFA/I structures suggests that CS1 pili may also unravel at low forces, consistent with our observations of structural polymorphism. Importantly, unraveling of CFA/I, P pili, and type I pili is reversible, implying that the pili can behave in a spring-like manner (18, 20, 47–49). Together, pilus extensibility and reversible unraveling would provide considerable pilus length variability while maintaining the structural integrity of the filament. These features are likely to be important to pathogenic *E. coli* in countering flow forces in the intestines and the urethra.

To our knowledge, this is the first report of structural heterogeneity for a pilus belonging to the chaperone-usher family. Most likely, structural polymorphism and filament extensibility exist in all members of this pilus family, where the filament structure is determined by donor strand complementation as well as flexible axial interactions between subunits. Such structural plasticity would represent an important pathogenic adaptation for these adhesive filaments and may also contribute to antigenic variability and immune evasion.

ACKNOWLEDGMENTS

We are grateful to June Scott at Emory University, who initially proposed structural studies for CS1 and provided the expression strains. We thank Joseph Wall at the BNL STEM facility. We also thank John Heuser and Robyn Roth at the Washington University School of Medicine for metal shadowing analysis, Darryl Hardie and Derek Smith at the Uvic Genome BC Proteomics Center for mass spectrometry analysis, and the staff at SSRL Beamline 11-1 for their assistance with remote collection of X-ray diffraction data. SSRL is a Directorate of SLAC National Accelerator Laboratory and an Office of Science User Facility operated for the U.S. Department of Energy Office of Science by Stanford University.

This work was supported by grants from the NIH (EB001567 to E.H.E.) and the Natural Sciences and Engineering Research Council (to L.C.). L.C. is supported by salary awards from the Canadian Institutes of Health Research (CIHR) and the Michael Smith Foundation for Health Research. J.L. is the recipient of a Frederick Banting and Charles Best Canada Graduate Scholarship from CIHR.

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