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Discovery of a Novel Inner Membrane-Associated Bacterial Structure Related to the Flagellar Type III Secretion System

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ABSTRACT The bacterial flagellar type III secretion system (fT3SS) is a suite of membraneembedded and cytoplasmic proteins responsible for building the flagellar motility machinery. Homologous nonflagellar (NF-T3SS) proteins form the injectisome machinery that bacteria use to deliver effector proteins into eukaryotic cells, and other family members were recently reported to be involved in the formation of membrane nanotubes. Here, we describe a novel, evolutionarily widespread, hat-shaped structure embedded in the inner membranes of bacteria, of yet-unidentified function, that is present in species containing fT3SS. Mutant analysis suggests a relationship between this novel structure and the fT3SS, but not the NF-T3SS. While the function of this novel structure remains unknown, we hypothesize that either some of the fT3SS proteins assemble within the hat-like structure, perhaps including the fT3SS core complex, or that fT3SS components regulate other proteins that form part of this novel structure.

IMPORTANCE The type III secretion system (T3SS) is a fascinating suite of proteins involved in building diverse macromolecular systems, including the bacterial flagellar motility machine, the injectisome machinery that bacteria use to inject effector proteins into host cells, and probably membrane nanotubes which connect bacterial cells. Here, we accidentally discovered a novel inner membrane-associated complex related to the flagellar T3SS. Examining our lab database, which is comprised of more than 40,000 cryo-tomograms of dozens of species, we discovered that this novel structure is both ubiquitous and ancient, being present in highly divergent classes of bacteria. Discovering a novel, widespread structure related to what are among the best-studied molecular machines in bacteria will open new venues for research aiming at understanding the function and evolution of T3SS proteins.

KEYWORDS bacteria, cryo-ET, flagella, secretion systems

Cryogenic electron tomography (cryo-ET) images intact cells in 3D in a native, frozenhydrated state, occasionally revealing unexpected structures (1–4). Examining a database of tens of thousands of electron cryo-tomograms of phylogenetically diverse bacterial species **Editor** Michael Y. Galperin, NCBI, NLM, National Institutes of Health

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FIG 1 Identification of a novel hat-like structure in *H. pylori*. A slice through an electron cryo-tomogram of an *H. pylori* cell, showing two hat-like structures (white circles in the enlargement). Scale bars represent 100 nm in the main panel and 25 nm in the enlargement.

that our lab has imaged over the past 15 years (5, 6), we found an inner membrane (IM)-associated, hat-shaped structure in diverse Gram-negative and Gram-positive bacteria (Fig. 1 and S1). In many cases, we observed multiple such structures, up to 10, distributed around the cell (see Movie S1 for an example from an *Escherichia coli* cell that was partially lysed, enhancing visibility of periplasmic structures). Subtomogram averages of the structure from different species revealed conserved characteristics: a hat-shaped part in the periplasmic (of diderms) or extracellular space (of monoderm *Bacillus subtilis*) and two cytoplasmic densities beneath, located ~10 nm below the IM (Fig. 2). In general, the periplasmic portion had a diameter of ~25 nm at its widest point at the outer leaflet of the IM, a diameter of ~10 nm at its top, and a height of ~12 nm.

The cytoplasmic densities were absent in the averages from three species: *Pseudoalter-omonas luteoviolacea, Hylemonella gracilis,* and *Bacillus subtilis.* In *P. luteoviolacea* and *B. subtilis,* this could be due to the fact that the tomograms from these species, which were collected for other projects, were of lysed, not intact, cells. While lysing cells decreases their thickness, thereby increasing the quality of cryo-tomograms, it can also affect the integrity of macromolecular complexes, e.g., the flagellar motor (7). However, it is unclear why the cytoplasmic densities were also missing in *H. gracilis,* since these cryo-tomograms were of intact cells. In the averages from several other species, the cytoplasmic density did not resolve into two distinct sections (Fig. 2).

MUTANT ANALYSIS

We identified the same novel structure in an *H. pylori* strain which contains a naturally occurring point mutation that disrupts the function of FliP (8), the platform on which other flagellar type III secretion system (fT3SS) proteins assemble (Fig. 3). We imaged this strain (henceforth referred to as *H. pylori fliP**) in the course of a study of flagellar assembly (9). Curiously, the diameter of the hat-like density in *H. pylori fliP** was reduced to only ~20 nm at its widest part, and the two cytoplasmic densities were missing (Fig. 4A and C). These changes reminded us of the changes that we observed in flagellar intermediates in *H. pylori fliP**. The flagellar membrane/supramembrane-ring complex (MS-complex) of *H. pylori fliP** similarly lacks its two cytoplasmic densities, and the MS-ring (FliF) adopts a smaller diameter (~20 nm) compared to that of motile *H. pylori* (~25 nm) (Fig. 4B and D). In the MS-complex, the two cytoplasmic densities, which also appear ~10 nm below the IM, are known to be formed by the C-terminal domain of the fT3SS protein FlhA (FlhA_c), which forms a nonameric ring (Fig. 3 and 4B) (10). The similar dimensions of the hat-like structure and the MS-complex as well as the comparable changes that occur in *H. pylori fliP** suggested to us that the hat-like structure is related to the fT3SS and that its cytoplasmic densities could also be formed by FlhA_c.

The fT3SS consists of a cytoplasmic part containing an ATPase and an IM-embedded part known as the core complex (fT3SScc). The fT3SScc is comprised of five proteins (FliP, FliQ,

Journal of Bacteriology



FIG 2 The hat-like complex is a widespread bacterial structure. A gallery of the hat-like structure in various bacterial species (*Xanthomonas citri, Vibrio harveyi, V. fischeri, V. cholerae, Salmonella enterica, Pseudoalteromonas luteoviolacea, Proteus mirabilis, Pseudomonas flexibilis, P. aeruginosa, Legionella pneumophila, Escherichia coli, Hyphomonas neptunium, Agrobacterium tumefaciens, Azospirillum brasilense, Brucella abortus, Helicobacter hepaticus, H. pylori, Campylobacter jejuni, Hylemonella gracilis, and Bacillus subtilis). Subtomogram averages are shown for all species except <i>C. jejuni* and *H. hepaticus*, for which insufficient data were available for averaging. For these two species, single tomogram average. Color coding indicates taxonomic class: green, Gammaproteobacteria; pink, Alphaproteobacteria; light orange, Epsilonproteobacteria; purple, Betaproteobacteria; and black, Bacilli. Scale bars represent 20 nm.

FliR, FlhB, and FlhA), with another protein, FliO, being required for assembly but not forming part of the complex (11, 12). Initially, FliP forms a pentameric platform on which FliQ, FliR, and FlhB assemble to create a $FliP_5FliQ_4FliR_1FlhB_1$ subcomplex upon which an FlhA ring is built (13) (Fig. 3). To explore our hypotheses that the hat-like structure is related to the fT3SS and that its cytoplasmic densities are formed by $FlhA_c$, we further mined our database for mutants lacking other fT3SScc proteins in species in which we had identified the hat-like structure.

First, we examined *Campylobacter jejuni* mutants lacking the C-terminal domains of FlhA ($\Delta flhAc$) and FlhB ($\Delta flhBc$) (10). In the $\Delta flhAc$ cells, compared to wild-type cells, the periplasmic



FIG 3 Schematic representation of the fT3SS assembly. The process starts with assembly of the fT3SScc proteins in the inner membrane (IM) and continues with the addition of the other components of the flagellum.

hat-like part was again smaller in diameter, and the two cytoplasmic densities again disappeared, supporting our hypothesis that they are formed by $FlhA_c$ (Fig. 5A). Of course, it is possible that $FlhA_c$ does not directly constitute the cytoplasmic densities but rather regulates the localization (or expression) of another protein or group of proteins that does. Such a regulatory role has been identified for another fT3SScc protein, FliO, which is responsible for the optimal expression of other flagellar genes (14). The $\Delta flhAc$ mutant arrests flagellar assembly at the C-complex (constituting the MS-ring, the fT3SS, the cytoplasmic [C-] ring, and other associated periplasmic components [9]), and we again observed that this intermediate lacked the cytoplasmic densities of FlhA_c (Fig. 5B). In contrast, in $\Delta flhBc$ cells, the hat-like structure was indistinguishable from the wild-type complex, both in diameter and in the presence of the associated cytoplasmic densities (Fig. 5C), and the cytoplasmic FlhA_c densities were







FIG 4 Comparison of the hat-like structure and MS-complexes in *H. pylori* strains. Central slices through subtomogram averages of the hat-like structures (A, C) and MS-complexes (B, D) of motile wild-type *H. pylori* (A, B) and nonmotile *H. pylori* fliP* (C, D). The diameters of the widest parts of the hat-like structures and the MS-rings are indicated. Numbers in parentheses indicate how many particles were used to produce the subtomogram average. (B, D) Adapted from (9). Scale bars represent 20 nm.

August 2022 Volume 204 Issue 8



FIG 5 Comparison of the hat-like structure and flagellar subcomplexes in fT3SScc mutants and lysed cells. (A–D) Central slices through subtomogram averages of the hat-like structure (A, C) and flagellar C-complex (B, D) in *C. jejuni* $\Delta flhA_c$ (A, B) and $\Delta flhB_c$ (C, D) cells. (E, F) Central slices through subtomogram averages of the hat-like structure in *P. aeruginosa* wild-type (E) and $flhA^*$ (F) cells. (G, H) Central slices through subtomogram averages of the hat-like structure (G) and a flagellar subcomplex constituting the MS-ring and rod (H) from lysed *B. subtilis* cells. (H) Adapted from (7). Numbers in parentheses indicate how many particles were used to produce the subtomogram average. Scale bars represent 20 nm.

also present in the flagellar C-complex (Fig. 4D). This lack of a visible difference was not surprising, since, unlike the large pentameric FliP ring or the nonameric FlhA ring, FlhB is a small protein present in a monomeric form in the fT3SScc. To confirm the generality of the relationship between the fT3SScc and the hat-like complex, we imaged an *flhA* mutant of *P. aeruginosa (flhA**, obtained from a transposon insertion mutant library). Here, also, the hat-like structure appeared smaller in size (18 nm versus 20 nm) and lacked clear cytoplasmic densities compared to that of the wild-type cells (Fig. 5E and F), although the number of identified structures in the *flhA** mutant was fewer than that of wild-type cells and produced a poorer-quality subtomogram average.

Another similarity between the hat-like structure and the flagellar system was the absence of the cytoplasmic densities in both structures in the cryo-tomograms of lysed cells. Recently, we showed that cell lysis can lead to the loss of the C-ring and FlhA_c densities in flagellar motors (7). In cryo-tomograms of lysed *Bacillus subtilis*, in which the hat-like structure lacked its two cytoplasmic densities, we identified a flagellar subcomplex consisting of the MS-ring and rod but lacking the cytoplasmic FlhA_c densities (Fig. 5G and H). Moreover, the MS-ring in this flagellar subcomplex looked morphologically similar (at the resolution of our subtomogram averages) to the hat-like structure (Fig. 5G and H).

These observations strengthened our hypothesis that the hat-like structure is related to the fT3SS, and we speculated that the periplasmic hat may be formed by the flagellar MS-ring protein, FliF, which assembles on the fT3SScc during flagellar assembly (Fig. 3) in a different, more closed conformation than that observed in the flagellar motor. We therefore generated and imaged a $\Delta fliF$ mutant in the *H. pylori fliP** background. To our surprise, while we could not identify any MS-complexes in this mutant, the hat-like complex was still present, though again with a smaller diameter and lacking clear cytoplasmic densities (Fig. 6A). The periplasmic hat, therefore, cannot be FliF. Subsequently, we speculated that it could be FliO, which is believed to have a large periplasmic domain in *H. pylori* (14) and is required for fT3SScc formation. We produced and imaged a $\Delta fliO fliP^*$ *H. pylori* mutant but still

H. pylori Δ fliF fliP* (146)H. pylori Δ fliO fliP* (29)H. pylori Δ fliG fliP* (60)H. pylori Δ fliG fliP* (27) A_{20} nm
 10^{40} B_{20} nm
 10^{40} C_{20} nm
 10^{40} D_{20} nm
 10^{40} A_{20} nm
 10^{40} B_{20} nm
 10^{40} C_{20} nm
 10^{40} D_{20} nm
 10^{40} A_{20} nm
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 10^{40} B_{20} nm
 10^{40} D_{20} nm
 10^{40} D_{20} nm
 10^{40} A_{20} nm
 10^{40} B_{20} nm
 10^{40} $H. pylori \Delta$ fliG fliP* (42) $H. pylori \Delta$ fliM fliP* (50) $H. pylori \Delta$ fliQ fliP* (42) F_{20} nm
 10^{40} F_{20} nm
 10^{40}

FIG 6 The effect of various flagellar-related mutations on the hat-like structure in *H. pylori flip**. (A–F) Central slices through subtomogram averages of the hat-like structures in the indicated mutants of *H. pylori*. Numbers in parentheses indicate how many particles were used to produce the subtomogram average. Scale bars represent 20 nm.

observed hat-like structures (again lacking the cytoplasmic densities and with a reduced width of \sim 20 nm at the base), ruling out FliO, as well (Fig. 6B).

From our study of flagellar assembly, we had cryo-tomograms of several flagellar-related mutants of *H. pylori*: $\Delta flas fliP^*$, $\Delta fliM fliP^*$, $\Delta fliG fliP^*$, and $\Delta fliQ fliP^*$. These mutants remove an additional fT3SScc protein (Δ *fliQ*), the C-ring proteins (Δ *fliM* and Δ *fliG*), or the tyrosine kinase responsible for the expression of the class II flagellar genes ($\Delta flgs$) (15), and they were all constructed in the *fliP** background to capture early flagellar intermediates (9). In all of these mutants, we observed hat-like structures, again with reduced diameter (\sim 20 nm at the widest part), with missing or less well-resolved cytoplasmic densities (Fig. 6C-F). These differences were not due to decreased resolution, since in each case, more particles were averaged than from wild-type cells (see numbers above, subtomogram averages). We also investigated whether any of the H. pylori mutants affected the abundance of the hat-like structure but found no obvious correlation between the strain and the number of hat-like structures per cell (Fig. S2). While H. pylori flip* cells had more structures per cell than did motile cells, other mutants, namely, $\Delta fliM$ fliP*, $\Delta fliO$ fliP*, and $\Delta fliQ$ fliP*, had fewer. Interestingly, the hat-like structure was still present in two mutants that did not contain visible flagellar intermediates, $\Delta fliG fliP^*$ and $\Delta fliF fliP^*$ (Fig. S2). Taken together, our observations suggest that while the cytoplasmic densities of the novel complex are likely FlhA_c, the periplasmic density is not formed by FliF or other fT3SScc proteins.

CORRELATION WITH FT3SS But Not With NF-T3SS

Next, we examined whether the hat-like structure is also related to the nonflagellar (NF-) T3SS. Bioinformatics analysis has shown that the NF-T3SS evolved from the flagellar system through a series of gene deletions, initially producing a machine thought to be involved in secreting proteins across the IM, an extant example of which is present in Myxococcales (16). Subsequently, the acquisition of an outer membrane-associated secretin resulted in the contact-dependent NF-T3SS (the injectisome) (16). First, we examined *Myxococcus xanthus*, which has NF-T3SS genes but no gene encoding a secretin, but found no hat-like structures in 95 cryo-tomograms (Fig. S3). Next, we examined *Prosthecobacter vanneervenii*, which has genes for the NF-T3SS and a secretin, but found no hat-like structures in 91 cryo-tomograms

TABLE 1 Summary of the species investigated in this study, indicating the presence (+) or absence (-) of fT3SS genes, NF-T3SS genes, and hat-like structures in each

| Species name | fT3SS genes | NF-T3SS genes | Hat-like structures |
|---------------------------------|-------------|---------------|---------------------|
| Helicobacter pylori | + | _ | Yes |
| Xanthomonas citri | + | _ | Yes |
| Vibrio harveyi | + | - | Yes |
| Vibrio fischeri | + | - | Yes |
| Vibrio cholerae | + | _ | Yes |
| Pseudoalteromonas luteoviolacea | + | - | Yes |
| Pseudomonas flexibilis | + | - | Yes |
| Legionella pneumophila | + | _ | Yes |
| Escherichia coli | + | - | Yes |
| Hyphomonas neptunium | + | - | Yes |
| Agrobacterium tumefaciens | + | _ | Yes |
| Azospirillum brasilense | + | - | Yes |
| Brucella abortus | + | - | Yes |
| Helicobacter hepaticus | + | - | Yes |
| Campylobacter jejuni | + | - | Yes |
| Hylemonella gracilis | + | - | Yes |
| Salmonella enterica | + | + | Yes |
| Proteus mirabilis | + | + | Yes |
| Pseudomonas aeruginosa | + | + | Yes |
| Bacillus subtilis | + | + | Yes |
| Prosthecobacter vanneervenii | _ | + | No |
| EPEC ^a | _ | + | No |
| Myxococcus xanthus | _ | + | No |
| Flavobacterium johnsoniae | _ | - | No |
| Coxiella burnetii | _ | - | No |
| Amoebophilus asiaticus | _ | _ | No |

^aThis EPEC mutant lacks all the NF-T3SS genes except the core complex ones and has all the flagellar genes except the fT3SScc ones.

(Fig. S3). Enteropathogenic *Escherichia coli* (EPEC) has both the NF-T3SS injectisome and the fT3SS (17). In our database, we have cryo-tomograms of an EPEC strain that has the core complex proteins of the NF-T3SS (*escR*, *escS*, *escT*, *escU*, and *escV*) but lacks the other components of the injectisome and the fT3SScc (see Materials and Methods). We identified no hat-like structures in 200 cryo-tomograms of this strain (Fig. S3). Finally, as a negative-control, we investigated bacterial species that do not have any T3SS-related genes (neither fT3SS nor NF-T3SS). We examined *Flavobacterium johnsoniae* (206 cryo-tomograms), *Coxiella burnetii* (149 cryo-tomograms), and *Amoebophilus asiaticus* (9 cryo-tomograms) but did not identify any hat-like structures in any of these species (Fig. S3). Together, these results suggest that the hat-like structure is specifically related to the fT3SS (Table 1).

DISCUSSION

What could the function of the hat-like structure be? We do not know; we can only speculate. One possibility, for example, is that it plays a role in the assembly of early flagellar components. The fT3SScc is the first to assemble during flagellar biogenesis, followed by the MS- and C-rings (11, 12). Then, the process proceeds in a cooperative way in which the absence of a certain component prevents subsequent parts from assembling (18–20). However, we recently found that in an *H. pylori fliP** strain, the MS-ring, which depends on Sec for secretion through the IM (21), can still assemble in the absence of the fT3SScc (9). One hypothesis for the function of the novel structure described here is that it is somehow involved in this process of early flagellar biogenesis. A physical association with the fT3SScc would explain the disappearance of the cytoplasmic densities and the smaller diameter of the periplasmic portion in the *fliP** and *flhA* mutants. While bioinformatics analysis in the future might help to identify candidates for this structure, the fact that flagellar genes do not always cluster in the same way in all species (22) makes this challenging.

Another possibility is that this apparently ancient structure may have diverged to serve a function independent of flagellar assembly. Supporting an independent role, the hat-like

TABLE 2 Total electron dose used for each tilt-series in each species

| Species name | Class | Cumulative electron |
|---------------------------------|-----------------------|---------------------|
| Xanthomonas citri | Gammaproteobacteria | 120 |
| Vibrio harvevi | Gammaproteobacteria | 160 |
| Vibrio fischeri | Gammaproteobacteria | 150 |
| Vibrio cholerae | Gammaproteobacteria | 160 |
| Salmonella enterica | Gammaproteobacteria | 200 |
| Pseudoalteromonas luteoviolacea | Gammaproteobacteria | 180 |
| Proteus mirabilis | Gammaproteobacteria | 160 |
| Pseudomonas flexibilis | Gammaproteobacteria | 100 |
| Pseudomonas aeruginosa | Gammaproteobacteria | 170 |
| Legionella pneumophila | Gammaproteobacteria | 100 |
| Escherichia coli | Gammaproteobacteria | 130 |
| Hyphomonas neptunium | Alphaproteobacteria | 180 |
| Agrobacterium tumefaciens | Alphaproteobacteria | 200 |
| Azospirillum brasilense | Alphaproteobacteria | 200 |
| Brucella abortus | Alphaproteobacteria | 160 |
| Helicobacter hepaticus | Epsilonproteobacteria | 200 |
| Helicobacter pylori | Epsilonproteobacteria | 120 to 130 |
| Campylobacter jejuni | Epsilonproteobacteria | 200 |
| Hylemonella gracilis | Betaproteobacteria | 75 |
| Bacillus subtilis | Bacilli | 160 |
| Myxococcus xanthus | Deltaproteobacteria | 200 |
| Prosthecobacter vanneervenii | Verrucomicrobiae | 180 |
| Flavobacterium johnsoniae | Flavobacteria | 100 |
| Coxiella burnetii | Gammaproteobacteria | 130 |
| Amoebophilus asiaticus | Cytophagales | 200 |
| EPEC | Gammaproteobacteria | 120 |

structures did not show a preferred spatial distribution in the cell, even in species with polar flagella. It is also possible that the structure serves different functions in different species. This could explain the structural variations we observed, such as the absence of the cytoplasmic densities in *H. gracilis*. Whatever the function of the hat-like complex, it joins the already rich repertoire of the (f)T3SS, which has roles in flagellar motility and protein translocation and was recently implicated in membrane nanotube formation (23).

MATERIALS AND METHODS

Strains and growth conditions. E. coli cells were grown as described (24). X. citri cells were grown in 2xTY medium for 14 h to stationary phase. V. cholerae, V. harveyi, and V. fischeri were grown as described (25). P. luteoviolacea were grown as described (4, 26). P. mirabilis were grown as described (27). P. aeruginosa were grown in LB medium at 37°C overnight. The P. aeruginosa flhA* mutant was obtained from a transposon library (mutant number 3296 from the nonredundant library http://pa14.mgh.harvard.edu/cgi-bin/pa14/ downloads.cgi) from Dianne Newman's lab at Caltech. L. pneumophila were grown as described (28). S. enterica were grown as described (29). P. flexibilis were grown in lactose growth medium. H. neptunium were grown to exponential phase in PYE medium. A. tumefaciens wild-type cells with a plasmid-borne VirC1-GFP translational fusion under control of the VirB promoter were grown in AB medium with 150 to 300 μ g/mL of kanamycin. A. brasilense and B. abortus were grown as described (3). H. hepaticus ATCC 51449 and H. gracilis were grown as described (24, 30). C. jejuni and its mutants were grown as described (10, 29, 31). B. subtilis protoplasts and lysate were prepared with lysozyme, following protocol (32). A motile revertant H. pylori 26695 isolate was selected by serial passage in Brucella broth supplemented with 10% heat-inactivated fetal bovine serum at 37°C, 5% CO₂ for 4 days until cultures reached an OD₆₀₀ of \sim 0.4. Nonmotile *H. pylori fliP** mutants were propagated on TSAII blood agar plates (BD Biosciences) at 37°C, 5% CO₂ for either 24 or 48 h prior to collection with a sterile cotton swab for grid preparation. H. pylori mutants ($\Delta fliM$ fliP*, $\Delta fliO$ fliP*, $\Delta flqS$ fliP*, $\Delta flqS$ fliP*, $\Delta fliG$ $fliP^*$, $\Delta fliQ$ fliP*, and $\Delta fliF$ fliP*) were grown directly from glycerol stocks on sheep blood agar at 37°C with 5% CO, for 48 h. Cells were either collected from the plate using a cotton swab, resuspended in PBS, then spun down and plunge-frozen directly, or passaged onto a new plate and allowed to grow for 24 h under the same conditions prior to collection and plunge-freezing. No difference could be discerned between the two samples by cryo-ET.

M. xanthus DK1622 cells were grown for 48 h in CTT medium at 30°C without antibiotics to OD_{600} =0.7. *F. johnsoniae* strain CJ2618 was grown as described (4). *P. vanneervenii* cells were grown as described (33). *A. asiaticus* strain 452471 was grown as described (34). *C. burnetii* Nine Mile phase II clone 4 strain was grown in the laboratory of Robert Heinzen at the National Institute of Allergy and Infectious Diseases. EPEC cells strain 8612 ($\Delta rorf1$ -escL $\Delta rorf3$ -mpc $\Delta escN$ -espF $\Delta flhBA \Delta fliOPQR \Delta minCD$) from a glycerol stock were grown in 2mL LB at 37°C in static conditions overnight. The next day, 1 mL of cells were spun down at 4,000 rpm for six minutes and concentrated 10 times. 3 μ L of cells were spotted onto glow-discharged grids placed on DMEM high-glucose agar plates (1.5% agar) for 3 h. Then, the grids were washed twice in PBS and transferred to starvation-medium (DMEM without phenol red, vitamins, and amino acids) agar plates and incubated for 1 h at 37°C. Subsequently, the samples were washed with PBS and plunge-frozen.

H. pylori mutagenesis. Flagellar mutants were generated in the nonmotile *H. pylori* 26695 background as previously described (9, 35).

Electron cryo-tomography sample preparation and imaging. Sample preparation for cryo-ET imaging was done as described (20, 25, 36). All data were collected by the Jensen Lab at Caltech. For the total cumulative electron dose used for each tilt-series in each species, see Table 2.

Image processing and subtomogram averaging. The three-dimensional reconstruction of the tiltseries was performed either automatically through the RAPTOR pipeline used in the Jensen Lab (5) or with the IMOD software package (37). Subtomogram averaging was done using the PEET program (38) with a 2-fold symmetrization applied along the particle *y*-axis. The numbers of particles included in each subtomogram average are indicated on the figures.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. SUPPLEMENTAL FILE 1, PDF file, 6.3 MB. SUPPLEMENTAL FILE 2, MOV file, 11.5 MB.

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