Supramolecular assembly of the *Escherichia coli* LdcI upon acid stress

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Pathogenic and commensal bacteria often have to resist the harsh acidity of the host stomach. The inducible lysine decarboxylase LdcI buffers the cytosol and the local extracellular environment to ensure enterobacterial survival at low pH. Here, we investigate the acid stress-response regulation of *Escherichia coli* LdcI by combining biochemical and biophysical characterization with negative stain and cryoelectron microscopy (cryo-EM) and wide-field and superresolution fluorescence imaging. Due to deleterious effects of fluorescent protein fusions on native LdcI decamers, we opt for three-dimensional localization of nanobody-labeled endogenous wild-type LdcI in acid-stressed *E. coli* cells and show that it organizes into distinct patches at the cell periphery. Consistent with recent hypotheses that in vivo clustering of metabolic enzymes often reflects their polymerization as a means of stimulus-induced regulation, we show that LdcI assembles into filaments in vitro at physiologically relevant low pH. We solve the structures of these filaments and of the LdcI decamer formed at neutral pH by cryo-EM and reveal the molecular determinants of LdcI polymerization, confirmed by mutational analysis. Finally, we propose a model for LdcI function inside the enterobacterial cell, providing a structural and mechanistic basis for further investigation of the role of its supramolecular organization in the acid stress response.

cryo-EM | SMLM | lysine decarboxylase | LdcI

Cell survival requires the adaptation of metabolism to changing environmental demands. Biochemical regulation of metabolic enzymes by cellular metabolites has been intensely studied for many decades. In addition, a growing body of recent light microscopy observations highlights the spatial regulation of enzymes by stimulus-induced phase separation into distinct loci—liquid droplets, amyloid-like aggregates, or ordered polymers (1, 2). In eukaryotes, the observed condensation of fluorescently labeled metabolic enzymes is triggered by stress, including medium acidification, hypoxia, and nutrient limitation. Enterobacteria such as *Escherichia coli*, *Salmonella*, and *Vibrio* encounter these types of conditions in the host gastrointestinal tract (3). One of the key enterobacterial proteins expressed during the acid stress response, upon oxygen limitation and regulated by the nutrient stress alarmone guanosine tetraphosphate (ppGpp), is the acid stress-inducible lysine decarboxylase LdcI (4–7). LdcI has been scrutinized since the early 1940s because of direct links between the efficiency of the acid stress response and pathogenicity (8–10). This enzyme transforms lysine into cadaverine while consuming protons and producing CO2, thereby contributing to buffering of the bacterial cytosol and the extracellular medium under acid stress conditions to promote bacterial survival. While both the structure and the function of LdcI have been thoroughly studied (7, 11), nothing is known about its localization inside the bacterial cell, because its fluorescent protein fusions were previously described to form inclusion bodies (12).

Whereas the overwhelming majority of superresolution fluorescence imaging is focused on eukaryotes, optical studies of bacterial systems are nearly exclusively centered on large macromolecular complexes with obvious superstructure such as cytoskeletal, cell division, chromosome partitioning, RNA degradation, and secretion machineries (13–15). As for bacterial metabolic enzymes, a handful of examples of their regulation by phase separation through stimulus-triggered polymerization concern well-conserved oligomeric proteins involved in nucleotide and amino acid metabolism, such as CTP synthase (16, 17) and glutamine synthetase (18). Interestingly, these enzymes are also able to assemble into filaments in vitro, and their in vivo condensates, detected both in bacteria and in eukaryotes, have been suggested to correspond to the polymerized state of the enzymes. Other examples of bacterial metabolic enzymes purified as polymers from bacterial extracts or forming polymers in vitro are the aldehyde-alcohol dehydrogenase AdhE (19) and the hydrogen-dependent CO2 reductase HDCR (20).

Specific to bacteria, LdcI is a decamer composed of five dimers tightly arranged into pentameric double rings (Protein Data Bank [PDB] ID: 3N75) (7). Interestingly, in vitro these double rings were observed by negative stain electron microscopy (ns-EM) to stack on top of one another into filament-like structures (7, 9) but only at nonphysiological pH below 5, under which LdcI should not be significantly expressed. Here, to address the spatial localization of the *E. coli* LdcI, we start by critically evaluating the effects of

**Significance**

Bacteria possess a sophisticated arsenal of defense mechanisms that allow them to survive in adverse conditions. Adaptation to acid stress and hypoxia is crucial for the enterobacterial transmission in the gastrointestinal tract of their human host. When subjected to low pH, *Escherichia coli* and many other enterobacteria activate a proton-consuming resistance system based on the acid stress-inducible lysine decarboxylase LdcI. Here we develop generally applicable tools to uncover the spatial localization of LdcI inside the cell by superresolution fluorescence microscopy and investigate the in vitro supramolecular organization of this enzyme by cryo-EM. We build on these results to propose a mechanistic model for LdcI function and offer tools for further in vivo investigations.

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is known to interact precisely with the C-terminal β-characterized binding partner of LdcI, the AAA+ ATPase RavA, N-terminal labeling with an FP would be likely to interrupt the outside. Therefore, on the one side, one could assume that an point to the ring periphery and are readily accessible from the toward the central pore of the double ring, while the C termini LdcI decamer structure, the N termini are oriented inwards and high monomericity, and high fluorescence quantum yields. In the high monomer behavior of mGeosM in gel filtration (25). This is also in line with the known propensity of mEos2, from which mGeosM is derived, to form tetramers at high concentration (33). To conclude, both N-terminal fusion constructs were unable to interact with RavA as shown by biolayer interferometry (BLI)-binding studies (Methods and SI Appendix, Fig. S1). This means that the functionality of these fusions cannot be considered as entirely retained. Thus, none of the four fusions were directly suitable for an in-depth imaging analysis under conditions of native LdcI expression upon acidic stress that we planned to undertake. Altogether, we demonstrated that structural integrity and unaltered interaction with known partners are useful readouts for functional preservation. Based on this result we propose that, when feasible, purification and structure-function analysis of FP fusions should be performed prior to interpretation of the protein localization inferred from observation of FP fusions by optical methods. This workflow can be used for example in cases where chromosomal manipulation for assessment of intact function is difficult and/or the phenotype is condition dependent or unclear, whereas basic in vitro biochemical and ns-EM characterization can be efficiently set up.

Immunofluorescence of LdcI in E. coli under Acid Stress Reveals Its Supramolecular Organization. Imaging techniques relying on genetically encoded FPs fused to the protein of interest are unmatched in terms of labeling specificity and efficiency. In addition, they do not require delivery of fluorescent molecules across the cell wall and do enable live cell imaging and single molecule tracking. In principle, because none of the FP fusions possessed the properties of native LdcI, to minimize the deleterious effect of the N-terminal or C-terminal FP fusions on the LdcI function, one could have opted for coexpression of a small amount of the FP-labeled LdcI with the wild-type (WT) LdcI. However, knowing the intricate regulation of the native LdcI promoter (see below), we decided to rather turn to immunofluorescence superresolution imaging of the endogenous enzyme. One of the drawbacks of this technique is the requirement for exogenous labeling and therefore the fixation and permeabilization of cells. These procedures are

Results

Fluorescent Protein Fusions Affect LdcI Structure without Modifying Localization of the Overexpressed Fusion Constructs. Because of the small size of the E. coli cell, we opted for imaging by single molecule localization microscopy (SMLM) (21–24). We therefore started by creating fusions of LdcI with genetically encoded phototransformable FPs. Considering that LdcI is an acid stress response enzyme, we chose either mGeosM (25) or Dendra2T69A (26) as FP markers because of their relatively low pKa values, high monomericity, and high fluorescence quantum yields. In the LdcI decamer structure, the N termini are oriented inwards and toward the central pore of the double ring, while the C termini point to the ring periphery and are readily accessible from the outside. Therefore, on the one side, one could assume that an N-terminal labeling with an FP would be likely to interrupt the LdcI tertiary structure. On the other side, the only well-characterized binding partner of LdcI, the AAA+ ATPase RavA, is known to interact precisely with the C-terminal β-sheet of LdcI (27). Assembly of two copies of double-pentameric rings of LdcI and five copies of hexameric RavA spirals results in a huge 3.3-MDa macromolecular cage of intricate architecture and largely unknown function (27–29). The LdcI–RavA cage is proposed to assist assembly of specific respiratory complexes in E. coli and to counteract acid stress under starvation conditions by alleviating ppGpp-dependent inhibition of LdcI (30–32). Thus, although these functions still require further investigation, preservation of the RavA-binding propensity should be one of the criteria for assessing the suitability of an LdcI–FP fusion. Therefore, both N- and C-terminal fusion constructs with either mGeosM or Dendra2T69A attached to LdcI via an appropriate linker were cloned into dedicated plasmids and overexpressed at conditions optimized for LdcI overproduction (Methods). Expression of fusion proteins was immediately detected by wide-field fluorescence imaging that showed a similar distribution for the four fusions (Fig. 1A). Each construct was then purified in order to assess its structural integrity and RavA-binding capacity in vitro, with a goal of defining the most suitable construct for the subsequent creation of a corresponding chromosomal fusion.

The N-terminal Dendra2T69A–LdcI fusion exclusively formed dimers at pHs between 5.7 and 7.5, confirming the structure-based prediction that the fluorescent tag at this position would perturb the dimer–dimer interaction (Fig. 1B and SI Appendix, Fig. S1). Admittedly, native LdcI was shown to dissociate into dimers in vitro at pH above ~7.5, but in this pH range LdcI is not supposed to be expressed in the cell and therefore this dissociation may be irrelevant. Surprisingly, in contrast to Dendra2T69A-LdcI, mGeosM–LdcI assembled into regular symmetric nonnative higher-order oligomers, with a dramatically altered quaternary structure (Fig. 1B and SI Appendix, Fig. S1). These oligomers were built of three LdcI tetramers, bridged together by additional densities. Noteworthy, mEos2, the fluorescent protein from which mGeosM was derived, crystallizes as a tetramer (PDB ID: 3505) that can be straightforwardly fitted into the LdcI-bridging densities (Fig. 1B). This illustrates that despite the fact that mGeosM was designed as a monomeric FP (with the first “m” explicitly standing for monomeric), some residual oligomeric tendency is still maintained. This propensity of mGeosM to oligomerize when bound to LdcI may be driven by avidity effects—as LdcI dimers begin to assemble into a decamer, the local concentration of mGeosM increases to such a point that oligomerization becomes energetically favorable, despite the apparent monomer behavior of mGeosM in gel filtration (25). This is also in line with the known propensity of mEos2, from which mGeosM is derived, to form tetramers at high concentration (33). To conclude, both N-terminal fusion constructs were considered as entirely retained. Thus, none of the four fusions were directly suitable for an in-depth imaging analysis under conditions of native LdcI expression upon acid stress that we planned to undertake. Altogether, we demonstrated that structural integrity and unaltered interaction with known partners are useful readouts for functional preservation. Based on this result we propose that, when feasible, purification and structure-function analysis of FP fusions should be performed prior to interpretation of the protein localization inferred from observation of FP fusions by optical methods. This workflow can be used for example in cases where chromosomal manipulation for assessment of intact function is difficult and/or the phenotype is condition dependent or unclear, whereas basic in vitro biochemical and ns-EM characterization can be efficiently set up.

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known to potentially affect cell morphology and/or ultrastructure, and therefore, when possible, the use of live cell imaging with FP fusions, or observation of unfixed samples under cryogenic conditions would be preferred. In addition, fixation, permeabilization, and specific exogenous labeling in bacteria present unique challenges because of the complex, often multilayered cell wall (34, 35). A significant advantage of immunolabeling, however, is the possibility of direct imaging of the WT protein, which should circumvent the dangers associated with FP fusions described above. The three prerequisites for imaging of WT systems by immuno-based labeling in general and of LdcI in particular are 1) availability of an antibody or nanobody (antibody fragment derived from heavy-chain-only camelid antibody) coupled to an organic fluorescent dye and directed toward the native protein, 2) precise knowledge of endogenous expression conditions, and 3) validation of an efficient permeabilization and immunolabeling technique that enables the antibody/nanobody to enter the cell and specifically target the protein of interest without altering its native organization.

As a first step toward immunofluorescence SMLM imaging of endogenous LdcI, we probed an anti-LdcI nanobody, hereafter called anti-LdcI-Nb. The complex between purified E. coli LdcI and the nanobody was purified by size exclusion chromatography (SEC) (Methods), and imaged by ns-EM. The resulting three-dimensional (3D) map of the LdcI/anti-LdcI-Nb complex demonstrates binding of anti-LdcI-Nb to each LdcI monomer in the decamer and reveals the location of the interaction site which is clearly distinct from the RavA-binding site (Fig. 2A and SI Appendix, Fig. S2). This nanobody was thus identified as a suitable labeling agent for LdcI imaging in E. coli cells and labeled with the fluorescent dye Alexa Fluor 647 (AF647) or Alexa Fluor 488 (AF488) (Methods).

Consistent with published data, LdcI expression could be induced by a pH shift experiment (i.e., transfer of bacteria from pH 7.0 into a pH 4.6 growth medium) in the presence of lysine under oxygen-limiting conditions (SI Appendix, Fig. S3A). While the WT strain grew well under these conditions and efficiently buffered the extracellular medium up to pH 6.2 in ~1.5 to 2 h (SI Appendix, Fig. S3A) concomitantly with the increase in the level of LdcI expression (SI Appendix, Fig. S3 B and C), the growth and the acid stress response of the ΔldcI mutant strain were severely impaired (SI Appendix, Fig. S3A). Since the peak of LdcI expression by the wild-type cells was achieved between 1 and 2 h after exposure to acid stress (SI Appendix, Fig. S3 B and C), a time point of 90 min was chosen for the subsequent labeling and imaging experiments. The specificity and performance of anti-LdcI-Nb in immunofluorescence labeling of permeabilized E. coli cells were characterized by flow cytometry and wide-field fluorescence imaging (SI Appendix, Fig. S3 D and E). Both techniques demonstrated that in the absence of LdcI expression no specific fluorescence is seen, confirming the suitability of anti-LdcI-Nb for immunofluorescence studies. Thus, the above-mentioned prerequisites for immuno-based imaging of cellular LdcI have been fulfilled. Noteworthy, the LdcI expression profile highlighted a considerable asset of the usage of immunolabeling instead of FP fusions for visualization of the endogenously expressed LdcI: indeed, the transient nature of the expression and the necessity of work under oxygen-limiting conditions may have created difficulties due to the longer maturation time of the FPs under these conditions.

Initial characterization of the cellular distribution of LdcI 90 min after exposure of E. coli cells to acid stress was carried out by wide-field fluorescence imaging. Based on these images, it appeared that natively expressed LdcI did not display a homogeneous cytoplasmic distribution but rather showed a patchy localization pattern (Fig. 2B). Three-dimensional stochastic optical reconstruction microscopy (STORM) imaging subsequently provided a more detailed view of this patchy distribution.
As shown in Fig. 3 and in SI Appendix, Fig. S4 and Movies S1–S5, the labeling density was lower in the center of the bacterium. This indicates a propensity for LdcI to cluster near the cell periphery, toward the inner membrane and the cell poles, rather than being distributed homogeneously through the volume. Recent investigations argue that, in most cases, in vivo clustering of metabolic enzymes corresponds to their polymerized states and represents an efficient means of regulation of enzymatic activity and metabolic homeostasis in response to a stimulus (1, 2). Thus, structure determination of these polymers is a crucial step toward the understanding of regulation mechanisms. We previously documented that in vitro at pH below 5 and high concentration, LdcI decamers indeed tend to stack (7). However, at pH 5.7, optimal for LdcI expression and enzymatic activity and likely corresponding to the internal pH upon acid stress, the oligomeric state of LdcI has not yet been addressed by EM. Supposing that the high local concentration of LdcI, clustered in patches in the E. coli cell under acid stress conditions, is likely to induce enzyme polymerization via stack formation, we next examined the LdcI assembly state at pH 5.7 by cryo-EM and discovered that it does indeed form straight polymers (Fig. 4).

Structural Determinants of LdcI Stack Formation Revealed by Cryo-EM. In order to understand the molecular determinants of LdcI polymerization at low pH and to provide a framework for the analysis of LdcI function under acid stress conditions, we solved the 3D structures of the LdcI decamer at pH 7.0 and of LdcI stacks at pH 5.7 by cryo-EM (Methods, Fig. 4, and SI Appendix, Figs. S5 and S6 and Table S1). The 2.8-Å resolution structure of the LdcI decamer at pH 7.0 (Fig. 4A) is extremely similar (SI Appendix, Tables S2 and S3) to the LdcI crystal structure solved at pH 8.5 in an inhibited ppGpp-bound state (7). However, contrary to pH 7.0 and even pH 6.2 where LdcI is still predominantly decameric (11), LdcI forms straight rigid filaments on the cryo-EM grid at pH 5.7, which corresponds to the pH of maximum LdcI enzymatic activity (Fig. 4B). The structure of a three-decamer stack was solved to a resolution of 3.3 Å, revealing the structural details of acid stress-induced LdcI polymerization (Fig. 4C). LdcI decamers stack tightly on top of one other, with negligible rotation between decamers along the stack. Each dimer fits snugly in the interdimer groove of decamers above and below.

A comparison of LdcI decamer structures taken from the LdcI stack cryo-EM map (at pH 5.7), the LdcI decamer cryo-EM map (at pH 7.0), and the crystal structure of decameric LdcI crystallized with bound ppGpp at pH 8.5 (PDB ID: 3N75) reveals some remarkable differences between the stack structure and the two decamer structures (Fig. 4D and E and SI Appendix, Tables S2 and S3). While the three structures do not show any major differences at the monomer level, a structural alignment of an LdcI dimer extracted from the LdcI decamer structures with a dimer extracted from the LdcI stack structure uncovers a rigid body-like rotation between monomers around a hinge region located at the monomer-to-monomer interface (SI Appendix, Tables S2 and S3 and Fig. 4D). This rotation results in a 5° tilt when comparing the N-terminal wing domains in LdcI dimers, and an overall slightly decreased diameter of the central cavity inside the stacked LdcI rings (Fig. 4D and E), which may
contribute to the tight packing of each monomer into the grooves of an opposing decamer in the stack.

A careful examination of the LdcI stack structure shows that two major interdecamer interfaces situated at a two-fold symmetry axis perpendicular to the stack direction contribute to stack formation (Fig. 5A). In particular, the first interface (Fig. 5B) is formed between residues R468, D470, E482, and Y485 situated in the ppGpp-binding domain (amino acids 418 to 564), and residues N314, D316, and G352 from the pyridoxal phosphate (PLP)-binding domain (amino acids 184 to 417). R468 sits between D316 and E344′ from a neighboring decamer, and makes electrostatic interactions through its η1 and η2 nitrogen atoms with D316. In addition, D470 interacts with the backbone of G352′, and E482 forms hydrogen bonds with N314′ and Y485. The second interface (Fig. 5B) is formed between residue N94 of the wing domain (amino acids 1 to 130) of one set of opposing dimers and a stretch of four residues in the ppGpp-binding domain—T444, E445, S446, and D447—at the end of helix α16 from a second set of opposing dimers. The wing domain residue N94 makes hydrogen bonds with E445′ of an opposing dimer. A second charged residue, D447, interacts with the backbone of T444′ and is held in place by R97 from the wing domain of a neighboring dimer in the same decamer.

Considering that the LdcI polymerization is induced by acid stress, we wondered which residues in the interface would be sensitive to pH changes. Surprisingly, most of the side chains involved in the interdecamer interface are charged arginine (pKa of 13), aspartate, and glutamate residues (pKa of 4 and 3, respectively), which do not change protonation state in the pH window relevant for LdcI activity (pH 5 to 7) (36). Nonetheless, other residues, situated outside the interaction interfaces, may drive stack formation through pH-dependent interactions that would in turn lead to the observed intermonomer rotation, and the associated constriction of the LdcI central cavity, coupled to the alignment of complementary contacts at interfaces one and two (Fig. 4 D and E). We note for example that H694 should be protonated in the LdcI stack structure at pH 5.7 but deprotonated in the two ring structures at pH 7.0 and pH 8.5, and that an electrostatic interaction between H694 and D192 situated in the linker region is present in the stack structure only (Fig. 5E). The formation of this salt bridge between H694 and D192 at pH 5.7, but not at pH 7.0 or higher, may thus directly contribute to the reorientation of monomers in the LdcI dimers, tightening of LdcI decamers, and alignment of interaction interfaces leading to subsequent stack formation at low pH.

To validate the observed interactions at the interdecamer interface, and to assess the individual importance of key residues involved in LdcI stack formation, we constructed four LdcI point mutants (R468E, R97E, H694A, and H694N), two double mutants (R97E/R468E and E445A/D447A), and one triple mutant (E445A/D447A/R468E). The mutants were purified following the protocol for wild-type LdcI (Methods), diluted into a buffer at pH 5.7 and observed by ns-EM (SI Appendix, Figs. S7 and S8). Although the grid preparation procedure for ns-EM yields stacks that are shorter and more curved and distorted when compared to the cryo-EM data (Fig. 4 B and SI Appendix, Figs. S7 and S8), our previous observations of the five paralogous E. coli amino acid decarboxylases justify the validity of this approach for a qualitative comparative analysis (7, 36). Ns-EM images make immediately apparent that the E445A/D447A double mutant does not have a significantly altered capability of stack formation at pH 5.7 when compared to WT LdcI, whereas the single R468E mutation is sufficient to completely abolish stack formation. The R97E mutant has a moderate destabilizing effect and displays fewer and smaller stacks then the WT LdcI. Consistent with the major role of R468 in the LdcI stack formation, the R97E/R468E double mutant and E445A/D447A/R468E triple mutant exclusively occur as decamers at low pH. Altogether, these results reveal that R468 is one of the key determinants of LdcI stack formation. Finally, similarly to the R97E mutant, a modest destabilization of stack formation is observed for the two histidine mutants, H694A and H694N, favoring our hypothesis that H694 may have an influence on the propensity of LdcI polymerization at low pH (SI Appendix, Figs. S7 and S8).

Discussion

Our synergistic approach, combining several in vitro techniques including biochemical characterization of purified fluorescent
protein fusions, ns-EM observation of mutants, low-resolution ns-EM reconstruction, and high-resolution cryo-EM analysis, with in vivo flow cytometry, wide-field, and 3D-STORM imaging, provides insights into the supramolecular LdcI assembly upon acid stress. This work adds to the very few known examples of regular polymerization as means of regulation of enzymes involved in amino acid metabolism in bacteria. The cryo-EM structure of the LdcI stacks presented here offers a molecular framework for future investigation of the role of LdcI polymerization in the acid stress response.

The high-resolution cryo-EM structures of the stack at pH 5.7 and of the decamer at pH 7.0 complement the previously solved crystal structure of the ppGpp-bound LdcI decamer obtained at pH 8.5. Previous serendipitous cocrystallization of LdcI with the nutrient stress response alarmone ppGpp led to assessment of the effect of ppGpp on LdcI activity and to a proposal that ppGpp would act as an LdcI inhibitor that prevents excessive lysine consumption upon nutrient limitation during acid stress (7). In addition, a ppGpp-dependent disassembly of the LdcI stacks had already been observed but could not be structurally explained since the ppGpp-binding site is situated between two neighboring dimers inside the LdcI ring (7). Our cryo-EM structures show that one of the residues involved both in ppGpp binding and in the stack formation is arginine 97 (R97). In the crystal structure of ppGpp-bound LdcI, R97 makes a stacking interaction with the guanosine imidazole ring of ppGpp, while in LdcI stacks R97 is involved in a key interaction at the second interface (Fig. 5D), where it locks D447 in a conformation allowing interactions between helices α16 from opposing LdcI decamers. Binding of ppGpp to LdcI interferes with the R97–D447 interaction, thereby

![Fig. 4. Cryo-EM analysis of the Ldc decamer at neutral pH and LdcI stacks formed in acid stress conditions. (A) Cryo-EM reconstruction of the Ldc decamer at pH 7.0 from side (Left) and top (Right) views. (Scale bar, 50 Å.) (B) Cryo-EM micrograph of LdcI stacks at pH 5.7. (Scale bar, 100 nm.) (Inset) The 2D class averages displaying clear secondary structural features. (Scale bar, 100 Å.) (C) Top (above) and side (below) views of the cryo-EM reconstruction of a three-decamer Ldc stack. Four dimers are colored either blue/gold or green/coral, corresponding to the coloring of the atomic model presented in Fig. 5. (Scale bar, 50 Å.) (D) Overlay of LdcI dimers at pH 8.5 (PDB ID: 3N75, shown in lilac), 7.0 (shown in cyan), and 5.7 (shown in gold). Alignment was carried out on a single monomer in the dimer pair. There is a 5° shift in the angle between the wing domains of LdcI at pH 5.7 and pH 7.0/8.5. The Inset shows an LdcI dimer (colored bluelyellow), clipped from the map shown in C, and positioned in the same orientation as the dimers shown above. WD, wing domain. (E) Comparison of the central decamer pore diameter between LdcI at pH 8.5, 7.0, and 5.7, showing a 5 Å decrease in the pore size upon stack formation at low pH.](https://doi.org/10.1073/pnas.2014383118)
most likely prohibiting correct positioning of D447 at the tip of helix α16, and resulting in a disruption of the second stack interface, leading to a moderate yet notable stack destabilization (SI Appendix, Figs. S7 and S8). Furthermore, our cryo-EM structure of ppGpp-free LdcI decamers at neutral pH enables us to rule out the effect of ppGpp on the differences observed between the LdcI stack structure at pH 5.7 and the ppGpp–LdcI crystal structure at pH 8.5. Indeed, despite the absence of ppGpp, R97 is still oriented toward the ppGpp-binding site and away from the interdecamer interface in the pH 7.0 decamer cryo-EM map (Fig. 5D). This suggests that the conformational changes in LdcI driving stack formation are mostly driven by low pH and not by the absence of ppGpp, although the D192 and H694 hinge residues are similarly far apart in the pH 8.5 and pH 7.0 structures (Fig. 5E). Our current work provides a solid experimental and structural basis for a future closer evaluation of the hypothesized role of ppGpp in LdcI regulation in vitro and in vivo (7).

Although the optical imaging experiments performed in this study do not allow quantitative evaluation of the nature of the clusters of endogenous LdcI at the bacterial periphery, we suppose, based on the parallel with observations in other systems (1, 2), that they may correspond to LdcI assembled into stacks. Possible reasons for such assembly may be to provide an efficient way to locally increase the LdcI concentration and to enhance its activity. But why would LdcI, an apparently highly soluble protein, be driven toward the inner membrane? What would be the advantage for acid-stressed E. coli cells to increase the concentration of LdcI via stack formation in these particular peripheral locations? Localization of proteins to specific sites in the bacterial membrane was shown to be generally driven by chemical factors such as the phospholipid composition of the lipid microdomains and by physical factors such as the degree of local curvature or the electric potential of the membrane (37). An attractive hypothesis would be that as an acid stress response protein performing a proton-consuming enzymatic reaction, LdcI may be attracted to proton sinks formed by anionic phospholipids which compartmentalize oxidative phosphorylation (OXPHOS) complexes for efficient functioning in bacterial respiration and adaptation to environmental changes. Indeed, OXPHOS complexes were often described to be unevenly distributed in the membrane in the form of mobile patches (38–42), providing evidence for highly dynamic and spatially organized bioenergetic membranes in E. coli cells (42). In addition, certain bacterial flottilins, which are essential scaffold proteins of the functional membrane microdomains, equivalent to the lipid rafts of eukaryotic cells, also show a patchy distribution and were

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Fig. 5. Structural insights into molecular determinants of the LdcI polymerization under acid stress conditions. (A) Atomic model of a two-decamer LdcI stack at pH 5.7. Dimers are colored as shown in the cryo-EM map in Fig. 4. Dashed boxes annotated with “B,” “C,” “D,” and “E” on the two-decamer stack indicate the locations of the zooms shown in B–E. (B) Closeup of the first decamer–decamer interface, which includes the key stack-forming residue R468. (C) Closeup of the second decamer–decamer interface. (D) Overlay of the LdcI decamer structures at pH 8.5 and 7.0 with the LdcI stack structure at pH 5.7, focused on R97. R97 in the LdcI stack (green) adopts a different conformation compared to the one in the pH 8.5 crystal structure (purple, with ppGpp bound) and the pH 7.0 cryo-EM map (cyan, without ppGpp bound). Despite the absence of ppGpp in the pH 7.0 sample, R97 is still oriented toward the ppGpp-binding site. (E) Comparison between the H694-D192′ distance in the LdcI stack at pH 5.7 (colored gold and blue), the LdcI decamer at pH 8.5 (Left, colored purple), and pH 7.0 (Right, colored cyan). Key residues are labeled for all panels, and interactions are shown with dotted lines.

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shown to interact with specific OXPHOS complexes (42, 43). In this regard, two different lines of evidence would be interesting to note. First, LdcI was described to copurify with a partially assembled complex I (44), whereas the LdcI-binding partner RavA, as well as ViaA, the second protein expressed from the rav/avia operon and which also interacts with RavA, were shown to interact with both complex I and fumarate reductase (31, 32). Second, the other E. coli PLP-dependent lysine decarboxylase LdcC, exercising the role of cadaverine biosynthesis irrespectively of acid stress (45), neither binds RavA nor forms stacks (29, 36), despite its 69% identity with LdcI. It is interesting to note that in LdcC the key stack-forming residue R468 has irrespectively of acid stress (45), neither binds RavA nor forms stacks (29, 36), despite its 69% identity with LdcI. It is interesting to note that in LdcC the key stack-forming residue R468 has.

Our STORM images suggest that individual patches have a peripheral distribution with a long-range stripy or pseudohelical organization (Fig. 3, SI Appendix, Fig. S4, and Movies S1–S6). While similar distributions have been documented for bacterial cytoskeletal, cell division, chromosome partitioning, RNA degradation, and secretion machineries, the impact on these for bacterial distributions ofFPs and other tags used for labeling, demonstrated specifically for YFP-MreB (46), warrants caution in their interpretation (14, 47). In contrast, here we observed endogenous wild-type LdcI labeled with anti-LdcI-Nb, which means that the ensuing pattern is likely real. Admittedly, we cannot completely exclude a potential perturbation by the employed fixation and permeabilization procedures. Nevertheless, examination of published images of OXPHOS patches (for example ref. 40) also hints to a similar organization. Excitingly, anionic phospholipid-specific dyes and fluorescently labeled antibiotics specific for nascent peptidoglycan synthesis upon cell elongation were also proposed to be distributed on helical or stripe patterns (37, 48–51). It is therefore tempting to imagine that inside the bacterial cell, LdcI has a tendency to follow a general path upon polymerization governed by the underlying chiral patterns in the cell envelope (52).

Finally, from the methodological view, our work convincingly illustrates that different FP fusion constructs can share the same cellular distribution despite a completely different structure, necessitating caution when inferring intact function from the preservation of the protein localization inside the cell. Our findings emphasize the importance of characterizing FP fusions using both biochemical and structural techniques, such as ns-EM, to ensure that the FP tag disrupts neither structure nor function of the target protein.

**Methods**

**Expression Constructs.** For fluorescence studies, several FP fusion constructs were generated starting from an available placid containing the coding sequence of LdcI (Uniprot entry P0A9H3), cloned in the pET22b(+) vector with a C-terminally 6His-tag (7). All constructs were generated using the Gibson cloning strategy and verified by sequencing analysis. The Gibson assembly was performed using 0.4 U T5 exonuclease, 2.5 U Phuson polymerase, and 400 U Taq ligase (New England Biolabs) in 1× IS buffer consisting of 100 mM Tris-HCl pH 7.5, 10 mM MgCl₂, 0.8 mM DNTP mix, 10 mM dithiothreitol (DTT), 50 mg polyethylene glycol (PEG)-8000, 1 mM nicotinamide adenine dinucleotide (NAD). A total of 7.5 μL of the Gibson Master Mix was mixed with 2.5 μL of D3 containing 100 ng of vector. The mix was incubated for 60 min at 50 °C. Transformations were performed in Top10 competent bacteria (One Shot TOP10 chemically competent E. coli, Invitrogen) and selected using 100 μg/mL ampicillin or 50 μg/mL kanamycin sulfate (Euromedex). Agarose gel purification and DNA plasmid extraction kits were purchased from Macherey-Nagel.

Dendra2tag-LdcI and mGeosM-LdcI were both cloned in the pET-TEV vector containing an N-terminal 6His-tag and a TEV cleavage site between Dendra2tag or mGeosM and the LdcI gene. LdcI-Dendra2tag and LdcI-mGeosM were both cloned in the pET22b(+)-vector with LdcI followed by either Dendra2tag or mGeosM containing an uncleavable C-terminal 6His-tag.

The anti-LdcI-Nb was obtained from the nanobody generation platform of the Architecture et Fonction des Macromolécules Biologiques (AFMB) laboratory (Marseille, France). A llama (Llama glama) was immunized with the purified wild-type LdcI. Lymphocytes were isolated from blood samples, a nanobody phage display library was generated, and enrichment of antigen-specific clones was performed using standard procedures (53, 54). Sequences of three positive clones were subcloned in the pHEN6 vector containing the pelB leader sequence from Erwinia carotovora for secretion into the periplasm, and a C-terminal 6His-tag (kindly provided by Aline Desmyter, AFMB, Marseille, France). One of the clones was used for the present study to yield anti-LdcI-Nb.

Plasmids, primers, and cloning strategy are summarized in SI Appendix, Table S4.

**Protein Purification.** LdcI–FP fusions and LdcI mutants were expressed in BL21 (DE3) cells grown in Luria-Bertani (LB) medium supplemented with 100 μg/mL ampicillin or 50 μg/mL kanamycin sulfate. Protein expression was induced using 40 μM isopropyl-β-D-1-thiogalactopyranoside (IPTG, Euromedex) and carried out overnight at 18 °C. The LdcI–FP fusions and LdcI mutants were purified as previously described for wild-type LdcI (27, 29, 36), in a final buffer containing 25 mM Hepes (pH 7.0), 0.3 M NaCl, 5% glycerol, 1 mM DTT, and 0.1 mM PLP.

The anti-LdcI-Nb was expressed in E. coli WK6 cells, following the protocol described in ref. 53, and purified with immobilized metal affinity chromatography (IMAC, using a Ni-NTA column) followed by SEC using a superdex 75 increase 10/300GL column (GE-Healthcare) equilibrated with a buffer containing 25 mM Tris pH 7.4 and 0.3 M NaCl.

In order to characterize the LdcI complex with anti-LdcI-Nb, the two proteins were mixed at a 1:5 molar ratio and submitted to size exclusion chromatography as carried out for LdcI alone but without DTT in the buffer. The top of the peak was taken for subsequent ns-EM analysis.

**Ns-EM on LdcI–FP Fusions, LdcI/anti-LdcI-Nb, and LdcI Mutants.** FP fusion samples after gel filtration were diluted to a concentration of ~0.025 mg/mL. A total of 3 μL was applied to the clean side of carbon on a carbon–mica interface and stained with 2% uranyl acetate (Dendra2T69A, LdcI-mGeosM, LdcI-Dendra2tag, LdcI/anti-LdcI-Nb, LdcI mutants) or 2% sodium silicotungstate (Dendra2tag-LdcI). Images were collected on a 120-kV Tecnai T12 electron microscope with an Orius 1000 camera (Gatan) or on a 200-kV Tecnai F20 electron microscope with either a OneView camera (Gatan) or a Ceta camera (Thermo Scientific). All images were collected with a defocus range of approximately −1.0 μm to −2.0 μm and with pixel sizes between 2.29 Å/pixel and 3.42 Å/pixel.

**Image Processing of LdcI–FP and LdcI/anti-LdcI-Nb.** Thirty-six micrographs of Dendra2tag-LdcI with a pixel size of 2.73 Å/pixel, 368 micrographs of mGeosM-LdcI with a pixel size of 2.29 Å/pixel, 23 micrographs of LdcI-Dendra2tag, 92 micrographs of LdcI-mGeosM with a pixel size of 3.42 Å/pixel, and 124 micrographs of LdcI/anti-LdcI-Nb with a pixel size of 2.82 Å/pixel were used for image analysis.

**Contrast Transfer Function (CTF) Estimation.** CTF estimation was performed with CTFIND3 (55). Semiautomatic particle selection was carried out with BOXER (56), with box sizes of 98 × 98 pixels for Dendra2tag-LdcI, 180 × 180 pixels for mGeosM-LdcI, 128 × 128 pixels for LdcI-Dendra2tag, and 112 × 112 pixels for LdcI/anti-LdcI-Nb, respectively. Particle extraction followed by several rounds of cleaning by two-dimensional (2D) classification in RELION 1.4 (57), resulted in the following number of particles for each dataset: Dendra2tag-LdcI = 7,140, mGeosM-LdcI = 5,514, LdcI-Dendra2tag = 832, LdcI-mGeosM = 12,211, and LdcI/anti-LdcI-Nb = 14,075.

For Dendra2tag-LdcI, initial model generation was carried out in RELION 2.1 (58) without any symmetry applied. For mGeosM-LdcI, initial model generation was carried out in RELION 2.1 with either C1, C3, or D3 symmetry applied. The results of all three calculations being very similar, the model with applied D3 symmetry was selected. For LdcI-Dendra2tag, LdcI-mGeosM, and LdcI/anti-LdcI-Nb, the previously determined LdcI decamer structure (PDB ID: 3N7S) was filtered to 60 Å and used as an initial model for 3D refinement.

The 3D refinement was carried out for each dataset with applied C2 symmetry for Dendra2tag-LdcI, L3 symmetry for mGeosM-LdcI, and D5 symmetry for LdcI-Dendra2tag, LdcI-mGeosM, and LdcI/anti-LdcI-Nb. Rigid body fitting of LdcI (PDB ID: 3N7S), mEos2 (PDB ID: 3S05), and Dendra2 (PDB ID: 2VZK) crystal structures was then carried out in Chimera (59) for the five datasets.
Cryo-EM on LdcI Stacks (pH 5.7). Wild-type LdcI was purified as previously described (11) from an E. coli strain impaired in the production of ppGpp (Novy serpentinidus ppGpp binding). Purified LdcI was diluted to a final concentration of ~0.25 mg/mL in a buffer containing 25 mM 2-(N-morpholino)ethanesulfonic acid (MES, pH 5.7), 0.3 M NaCl, 5% glycerol, 1 mM DTT, and 0.1 mM PLP. A total of 3 μL of the sample was applied to a glow-discharged R 2/1 300 mesh holey carbon copper grid (Quantifoil Micro Tools GmbH) and plunge frozen in liquid ethane using a Vitrobot Mark IV (FEI) operated at 100% humidity. Data sets were recorded at the European Synchrotron Radiation Facility in Grenoble, France (60), on a Titan Krios microscope (Thermo Scientific) equipped with a BioQuantum LS/967 energy filter (Gatan) and a K2 summit direct electron detector (Gatan) operated in counting mode. A total of 2,256 movies of 30 frames were collected with a total exposure of 6 s, total dose of 29.3 e-/Å², and a slit width of 20 eV for the energy filter. All movies were collected at a magnification of 130,000 × and corresponding to a pixel size of 1.652 Å/pixel at the specimen level. A summary of cryo-EM data collection parameters can be found in SI Appendix, Table S1.

Image Processing of LdcI Stacks (pH 5.7). Motion correction and dose weighting of the recorded movies were performed using MotionCor2 (61). CTF parameters were determined on the aligned and dose-weighted sums using CTFFIND4 (62). After manual inspection of the dose-weighted sums, the best micrographs were selected for further processing. LdcI stacks were manually picked using e2helixboxer in EMAN2 (63). A total of 15,165 LdcI-stack particles were extracted in RELION 3.0 (64) with an extract size of 320 pixels, resulting in boxes containing three LdcI decamers, and with the –helix option with an outer diameter of 160 pixels and a helical rise of 77 pixels. After particle extraction, per-particle CTF correction was performed using patch motion correction in CryoSPARC, discarding the first two frames. Initial 3D refinement with imposed D5 symmetry was carried out in RELION 3.0, resulting in a cleaned dataset containing 15,157 particles. Initial 3D refinement with imposed D5 symmetry was carried out in RELION 3.0, using an initial model generated by manually stacking three LdcI decamers (PDB ID: 3N75) in Chimera (59) and low-pass filtering the resulting LdcI stack to 40 Å. The resulting 4.3 Å resolution 3D reconstruction, along with the cleaned particle stack, was subsequently imported into CryoSPARC (66). A final real space refinement, using the best 558 (21.7%) micrographs were subjected to 2D classification in RELION 3.0, resulting in a map with a resolution containing 15,157 particles. Initial D5 refinement with imposed D5 symmetry was carried out in RELION 3.0, using an initial model generated by manually stacking three LdcI decamers (PDB ID: 3N75) in Chimera (59) and low-pass filtering the resulting LdcI stack to 40 Å. The resulting 4.3 Å resolution 3D reconstruction, along with the cleaned particle stack, was subsequently imported into CryoSPARC (66). A final real space refinement, using the best 558 (21.7%) micrographs were subjected to 2D classification in RELION 3.0, resulting in a map with a resolution of 3.2 Å based on the 0.143 gold-standard Fourier shell correlation (FSC) criterion (67). The final map was sharpened using a B-factor of ~96 Å². A local resolution estimation of the final 3D reconstruction was calculated in RELION 3.0. A summary of cryo-EM data collection parameters and imaging processing steps for LdcI stacks can be found in SI Appendix, Table S1 and Fig. S5, with local resolution and FSC curves shown in SI Appendix, Fig. S6.

Fitting of Structures and Refinement. For fitting of atomic models in the 3D reconstructions of LdcI at pH 5.7 (stack) or pH 7.0 (decamer), two (for the stack) or one copy (for the decamer) of the LdcI X-ray crystal structure (PDB ID: 3N75) were first rigid-body fitted in the corresponding 3D reconstructions using Chimera. Refinement was performed using the Phenix software (68) and was identical for both 3D reconstructions. A first round of real space refinement was carried out with enabled rigid-body, global minimization, local grid search and atomic displacement parameter (ADP) refinement parameters, and imposing rotamer, Ramachandran, non-crystallographic symmetry (NCS), and reference model (PDB ID: 3N75) restraints. After several rounds of manual correction in Coot (69), a final round of real space refinement was then performed using the same settings, but without rigid body refinement and without applying reference restraints, and setting the “nonbonded_weight” parameter to 500. A summary of refinement and model validation statistics can be found in SI Appendix, Table S1.

pH Shift Experiment. Stationary phase cultures, which were grown overnight from single colonies in LB medium, were diluted to OD600 ~0.01 and regrown at 37 °C to OD600 ~1–1.5. After 1 h, LB medium was replenished to a final OD600 of 0.1. From this culture, 14 mL was transferred to 15-mL Falcon tubes and pelleted by centrifugation at room temperature for 5 min. The supernatant was decanted, whereby systematically around 200 μL LB remained in the Falcon tube. The pellet of bacteria was resuspended and afterward, LB-4.6 containing 30 mM l-lysine was added to the cells up to 14 mL. To prepare LB-4.6 medium, LB powder (Sigma-Aldrich) was completely dissolved in distilled water by stirring for 30 min. The pH of 4.6 was then adjusted using HCI. After autoclaving, sterile filtered l-lysine was added to LB-4.6. For this, l-lysine was dissolved in an aliquot of LB-4.6, sterile filtered, and mixed with the remaining LB-4.6. A total of 30 mM l-lysine was used. In order to grow the culture under oxygen-limiting conditions, the lid of the Falcon tube was closed and the tubes were placed at 37 °C on a shaker (150 rpm). After the OD600 had reached 0.3, the cultures were taken for western blotting, using Odyssey infrared imaging system (LI-COR), followed by immunofluorescence staining. For each time point, a new tube was opened and not reused further.

Western Blotting. SDS-PAGE was performed with a Bio-Rad electrophoresis chamber using standard 12% reducing SDS-PAGE gels. Proteins were transferred to a nitrocellulose membrane (Bio-Rad) using a Trans-Blot Turbo Transfer System (Bio-Rad). The membrane was blocked for 1 h using 5% bovine serum albumin (BSA) in Tris-buffered saline (TBS) buffer supplemented with 0.1% Tween (TBS-T). Afterward, the membrane was incubated for 1 h with an anti-LdcI antibody (Qalam-Antibodies) in BSA/TBS-T (1:5,000). The membrane was subsequently washed 3 times in TBS-T and incubated for 1 h with horseradish peroxidase (HRP)-coupled anti-rabbit antibody (Sigma-Aldrich) in BSA/TBS-T. Finally, the membrane was washed 3 times in TBS-T. The membrane was rinsed once with TBS, prior to detection of antibody-labeled proteins using ECL reagent (GE Healthcare). Incubation and detection of antibody were performed at 25 °C.

Nanobody Labeling. For the labeling reaction, 50 μL nanobody (i.e., 200 μg) was pipetted into a 1.5 mL tube, placed on ice, and supplemented with 5.5 μL of 1 M bicarbonate buffer at pH 8.3. A total of 100 μg of Alexa-647 or Alexa-488 N-hydroxysuccinimide (NHS) ester dye (Life Technologies, A37573 and A20000) were dissolved in 10 μL dimethyl sulfoxide (DMSO) to final concentration of 10 mg/mL. A total of 5 μL (i.e., 40 nmol) dye in DMSO was added to the protein and incubated for 1 h at room temperature in a shaking block, covered with an aluminum foil to protect the dye from the light. Excess dye was removed by iterative buffer exchange using a 3K spin column (Amicon-Ultra-4 Centrifugal Filters Ultracell 3K, Millipore UFC800396). Finally, the sample was centrifuged at 10,000 rpm in a TBS-T. The membrane was rinsed once with TBS, prior to detection of antibody-labeled proteins using ECL reagent (GE Healthcare). Incubation and detection of antibody were performed at 25 °C.

Cell Preparation for Immunofluorescence Staining of E. coli Cells with Nanobodies. For immunofluorescence staining, cells were subjected to a pH shift in order to induce LdcI expression, as described in the pH Shift Experiment above. The OD600, measuring the OD, was measured, and the volume of cells corresponding to OD 4, with OD = 1 corresponding to about 8 × 10^10 cells per mL, were collected by centrifugation. After removal of LB by pipetting, cells were resuspended in 2 mL of...
4% formaldehyde (FA) in PBS (made from 16% formaldehyde solution, methanol-free from Thermo Scientific). Falcons were placed on a rotor for centrifugation at 600 rpm at room temperature. After fixation, cells were collected by centrifugation and the solution was removed by pipetting. Cells were then resuspended in 14 mL PBS (Gibco, Thermo Scientific) to remove and dilute the fixative. Cells were permeabilized for 10 min with 2 mL 0.1% Triton X-100 in PBS and subsequently washed three times with 10 mL PBS. Finally, cells were transferred to 1.5-mL tubes, centrifuged, and resuspended in 200 μL 1% BSA/PBS (BSA/PBS solution was dissolved for 30 min and sterile filtered to avoid clumps). After 30 min of incubation, 0.5 μg anti-LdcI-Nb, labeled with the dedicated dye, was added to the 200 μL bacteria BSA/PBS suspension. Cells were incubated with the labeled anti-LdcI-Nb for 16 h at 4 °C. The next day, cells were washed three times with 1 mL PBS, centrifuged to remove antibody solution, and resuspended in 250 μL PBS. When needed, Hoechst 33342 (Sigma-Aldrich) was added to a final concentration of about 100 ng/mL.

Wide-Field Imaging and Flow Cytometry. For epifluorescence imaging, 2 μL of cells was placed between a glass slide and a coverslip, which had been carefully pressed together, and observed using an inverted IX81 microscope, with a UPLFLN 100x oil immersion objective (numerical aperture [N.A.] 1.3) (Olympus), using the appropriate specific excitation and emission filters for Alexa Fluor 488 (A1-488-35315B set, Semrock) and DAPI (D1-50606B set, Semrock). Acquisitions were performed with Velocity software (Quorum Technologies) with a sCMOS 2048 × 2048 camera (Hamamatsu ORCA Flash 4, 16 bit/pixel) achieving a final magnification of 64 nm per pixel. For flow cytometry, 50 μL of cell suspensions were injected in a MACSQuant VYB flow cytometer (Miltenyi Biotech) using the 488-nm excitation and 525/50-nm emission channel (B1). AF488 positive populations were estimated after forward scatter and side scatter gating on the cells. Data were further processed with MACSQuantify software (Miltenyi Biotech).

STORM Imaging. For single molecule localization microscopy, cells were transferred to a glucose buffer containing 50 mM NaCl, 150 mM Tris (pH 8.0), 10% glucose, 100 mM MEA (mercaptoethylamine) and 1 μM ADP for 10 min. Streptavidin-coated biosensors (FortéBio) were incubated with 1 mM ADP for 10 min. Streptavidin-coated biosensors (FortéBio) were functionalized with biotinylated Rava-AviTag, then quenched with 10 μg/mL biocytin. Experiments with the wild-type LdcI are adapted from ref. 28. For Ct-term labeled FP fusions, pins were dipped in wells containing a range of LdcI-FP concentrations from 0 to 1,000 nM, with no binding signal recorded at any concentration of LdcI-FP.

Data Availability. Cryo-EM maps, along with the corresponding fitted atomic structures, have been submitted to the EM Data Bank (EMDB) and Protein Data Bank (PDB) with accession codes EMDB 10850 and PDB 6YNL for LdcI stacks (pH 5.7), and EMDB 10849 and PDB 6YN5 for LdcI decamers (pH 7.0).

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36. U. Kanjee, I. Gutsche, S. Ramachandran, W. A. Houry, The enzymatic activities of the

31. K. S. Wong et al., Structural insights into ATP hydrolysis by the MoxAR Pase Rava and the

24. S. T. Hess, T. P. K. Girirajan, M. D. Mason, Ultra-high resolution imaging by fluores-

42. A. Magalon, F. Alberge, Distribution and dynamics of OXPHOS complexes in the

43. M. Bramkamp, D. Lopez, Exploring the existence of lipid rafts in bacteria. Microbiol.

40. I. Fishov, C. L. Woldringh, Visualization of membrane domains in Escherichia coli. Micro-

45. Y. Yamamoto, Y. Miwa, K. Miyoshi, J. Furuyama, H. Ohmori, The Escherichia coli ldcC
gene encodes another lysine decarboxylase, probably a constitutive enzyme. Genes Genet.

44. H. Erhardt et al., Disruption of individual nuc-genes leads to the formation of par-
tially assembled NADH:ubiquinone oxidoreductase (complex I) in Escherichia coli. Biochim.

47. H. Chang et al., A unique series of reversibly switchable fluorescent proteins with

21. J. T. Dervan, V. J. Hruby, E. L. Roche, Nat. Methods 7, 785–793 (2010).

26. R. Berardozzi, V. Adam, A. Martins, D. Bourgeois, Arginine 66 controls dark-state

23. M. Ingerson-Mahar, A. Briegel, J. N. Werner, G. J. Jensen, Z. Gitai, The metabolic

20. T. T. Hess, D. A. Stone, P. A. Sajdak, Nat. Methods 9, 703–707 (2012).

18. T. G. Frey, D. Eisenberg, F. A. Eiserling, Glutamine synthetase forms three- and seven-

17. M. Ansari, J. P. Deisboeck, S. M. Deisboeck, Nat. Struct. Mol. Biol. 8, 697–701 (2001).

16. M. Ingersoll-Mahar, A. Briegel, J. N. Werner, G. J. Jensen, Z. Gitai, The metabolic

19. J. S. Wang, L. Furchtgott, K. C. Huang, J. W. Shaevitz, Helical insertion of peptidoglycan

22. C. J. DeMoss, J. S. Lu, J. M. Wang, Nat. Methods 5, 778–782 (2008).

28. A. Rohou, N. Grigorieff, SCrying: Accurate determination of local defocus and specimen tilt

27. H. Malet et al., Single-molecule imaging of bacterial protein complexes labeled with fluorescent

13. A. K. Munshi, D. A. Spaeth, Nat. Methods 5, 255–259 (2008).

14. R. B. Davis et al., Spatial resolution of fluorescent protein labels within live Escherichia coli

15. S. T. Hess, T. P. K. Girirajan, M. D. Mason, Ultra-high resolution imaging by fluores-

12. V. DeCinque, J. W. Shaevitz, Nat. Methods 4, 909–911 (2007).

11. M. Ptak, R. A. Daniel, J. Errington, Curr. Opin. Microbiol. 10, 121–129 (2007).

10. A. J. Mindell, N. Grigorieff, Accurate determination of local defocus and specimen tilt in electron microscopy. J. Struct. Biol. 142, 338–347 (2003).

9. S. T. Hess, T. P. K. Girirajan, M. D. Mason, Nat. Methods 4, 909–911 (2007).

8. L. E. Hinkle, E. L. Roche, Nat. Struct. Mol. Biol. 9, 263–268 (2002).

7. E. F. Pettersen et al., UCSF Chimera—a visualization system for exploratory research and analysis. J. Comput. Chem. 25, 1605–1612 (2004).

6. K. Zhou, P. J. O’Toole, N. Pavlenkov, Lipid spirals in Bacillus subtilis and their role in cell division. Mol. Microbiol. 68, 1315–1327 (2008).

5. M. T. Swulius, G. J. Jensen, The helical MreB cytoskeleton in Escherichia coli MC1000/p

4. Y. Yamamoto, Y. Miwa, K. Miyoshi, J. Furuyama, H. Ohmori, The Escherichia coli ldcC gene encodes another lysine decarboxylase, probably a constitutive enzyme. Genes Genet. Syst. 72, 167–192 (1997).

3. G. K. Kim et al., Aldehyde-alcohol dehydrogenase forms a high-order spirochrome archi-

1. G. G. Abney, M. J. Hoffmann, K. M. Smith, J. Bacteriol. 187, 4989–4997 (2005).

2. A. Magalon, F. Alberge, Distribution and dynamics of OXPHOS complexes in the

Supramolecular assembly of the Escherichia coli LdcI upon acid stress https://doi.org/10.1073/pnas.2014383118

Biochim. Biophys. Acta 1817, 863–871 (2012).

Biochim. Biophys. Acta 1817, 863–871 (2012).

Biochim. Biophys. Acta 1817, 863–871 (2012).

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https://doi.org/10.1073/pnas.2014383118

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