

1 **Title**

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3 **Supramolecular assembly of the *E. coli* Ldcl upon acid stress**

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14

15 **Abstract**

16 Pathogenic and commensal bacteria often have to resist the harsh acidity of the host stomach. The
17 inducible lysine decarboxylase Ldcl buffers the cytosol and the local extracellular environment to
18 ensure enterobacterial survival at low pH. Here, we investigate the acid-stress response regulation
19 of *E. coli* Ldcl by combining biochemical and biophysical characterisation with negative stain and
20 cryo-electron microscopy, and wide-field and super-resolution fluorescence imaging. Due to
21 deleterious effects of fluorescent protein fusions on native Ldcl decamers, we opt for three-
22 dimensional localisation of endogenous wild-type Ldcl in acid-stressed *E. coli* cells, and show that it
23 organises into patches following an apparent long-range pseudo-helical order. Consistent with
24 recent hypotheses that *in vivo* clustering of metabolic enzymes often reflects their polymerisation as
25 a means of stimulus-induced regulation, we show that Ldcl assembles into filaments *in vitro* at low
26 pH. We solve the structures of these filaments and of the Ldcl decamer formed at neutral pH by
27 cryo-electron microscopy, and reveal the molecular determinants of Ldcl polymerisation, confirmed
28 by mutational analysis. Finally, we propose a model for Ldcl function inside the enterobacterial cell,
29 providing a structural and mechanistic basis for further investigation of the role of its supramolecular
30 organisation in the acid stress response.

31

32 **Introduction**

33 Cell survival requires the adaptation of metabolism to changing environmental demands.
34 Biochemical regulation of metabolic enzymes by cellular metabolites has been intensely studied for
35 many decades. In addition, a growing body of recent light microscopy observations highlights the
36 spatial regulation of enzymes by stimulus-induced phase separation into distinct loci – liquid droplets,
37 amyloid-like aggregates or ordered polymers¹. In eukaryotes, and particularly in yeast, the observed
38 condensation of fluorescently-labelled metabolic enzymes is triggered by stress, including medium
39 acidification, hypoxia and nutrient limitation. Enterobacteria such as *Escherichia coli*, *Salmonella* and
40 *Vibrio* encounter these types of conditions in the host gastrointestinal tract². One of the key
41 enterobacterial proteins expressed during the acid stress response, upon oxygen limitation and
42 regulated by the nutrient stress alarmone guanosine tetraphosphate (ppGpp), is the acid stress-

43 inducible lysine decarboxylase LdcI³⁻⁶. LdcI has been scrutinised since the early 1940s because of
44 direct links between the efficiency of the acid stress response and pathogenicity⁷⁻⁹. This enzyme
45 transforms lysine into cadaverine while consuming protons and producing CO₂, thereby contributing
46 to buffering of the bacterial cytosol and the extracellular medium under acid stress conditions to
47 promote bacterial survival. While both the structure and the function of LdcI have been thoroughly
48 studied^{6,10}, nothing is known about its localisation inside the bacterial cell.

49 Whereas the overwhelming majority of super-resolution fluorescence imaging is focused on
50 eukaryotes, optical studies of bacterial systems are nearly exclusively centred on large
51 macromolecular complexes with obvious superstructure such as cytoskeletal, cell division,
52 chromosome partitioning, RNA degradation and secretion machineries¹¹⁻¹³. However, the relevance
53 of the documented patchy or long-range helical localisation of some of these assemblies is now
54 questioned and requires re-evaluation. Indeed, the vast majority of these studies were based on
55 labelling with either fluorescent proteins or epitope tags shown to be able to induce artefactual
56 associations and localisations¹²⁻¹⁵.

57 A handful of examples of regulation of bacterial metabolic enzymes by phase separation
58 through stimulus-triggered polymerisation concern well-conserved oligomeric proteins involved in
59 nucleotide and amino acid metabolism, such as CTP synthase^{16,17} and glutamine synthetase¹⁸.
60 Interestingly, these enzymes are also able to polymerise into filaments *in vitro*, and their *in vivo*
61 condensates, detected both in bacteria and in eukaryotes, have been suggested to correspond to
62 the polymerised state of the enzymes. Other examples of bacterial metabolic enzymes purified as
63 polymers from bacterial extracts or forming polymers *in vitro* are the aldehyde-alcohol
64 dehydrogenase AdhE¹⁹ and the hydrogen-dependent CO₂ reductase HDCR²⁰.

65 Specific to bacteria, LdcI is a decamer composed of five dimers tightly arranged into
66 pentameric double-rings (PDB ID: 3N75)⁶. Interestingly, *in vitro*, at pH below 6, these double-rings
67 were observed by negative stain electron microscopy (ns-EM) to stack on top of one another into
68 filament-like structures^{6,8}. Here, we address the spatial localisation of the *E. coli* LdcI. To critically
69 evaluate labelling artefacts and define the optimum constructs for subsequent chromosomal
70 manipulation, we start by overexpressing and purifying different fluorescent protein (FP) fusions of
71 LdcI, performing their structural and biochemical characterisation *in vitro*, and observing the
72 distribution of the overexpressed constructs *in vivo*. This methodological section enables us to
73 propose a workflow that brings together examinations of *in vivo* protein localisations with *in vitro*
74 biochemical and ns-EM characterisations of the purified FP fusions in order to ensure artefact-free
75 optical imaging investigations. This analysis is followed by unveiling a supramolecular organisation
76 of endogenous LdcI inside the *E. coli* cell upon acid stress, coupled to determination of the *in vitro*
77 structure of the LdcI polymers by cryo-electron microscopy (cryo-EM). For comparison, we also
78 determined the cryo-EM structure of the LdcI decamer at neutral pH. In addition, mutational analysis
79 of the LdcI stack-forming interfaces allowed identification of critical residues involved in stack
80 formation. Finally, we discuss the observed LdcI localisation pattern in the light of the wealth of
81 available functional and imaging data, and offer a structural and mechanistic basis of supramolecular
82 LdcI assembly, which will aid in the design of future experiments linking LdcI stack formation to *E.*
83 *coli* acid stress fitness.

85 Results

86 Fluorescent protein fusions affect Ldcl structure without modifying localisation of the 87 overexpressed fusion constructs

88 Because of the small size of the *E. coli* cell, we opted for super-resolution microscopy imaging
89 ^{>21} and set out to localise Ldcl inside the cell upon acid stress by either Photoactivation
90 Localisation Microscopy (PALM) or Stochastic Optical Reconstruction Microscopy (STORM)^{22–24}. *A*
91 *priori*, PALM seemed more relevant because this technique relies on genetically-encoded FPs fused
92 to the protein of interest, and is therefore unmatched in terms of labelling specificity and efficiency.
93 In addition, PALM does not require delivery of fluorescent molecules across the cell wall, and
94 enables live cell imaging and single molecule tracking.

95 Considering that Ldcl is an acid stress response enzyme, we opted for either mGeosM²⁵ or
96 Dendra2_{T69A}²⁶ as FP markers because of their relatively low pKa values, high monomericity and high
97 fluorescent quantum yields. In the Ldcl decamer structure, the N-termini are oriented inwards and
98 towards the central pore of the double ring, while the C-termini point to the ring periphery and are
99 readily accessible from the outside. Therefore, on the one side, one could assume that an N-terminal
100 labelling with an FP would be likely to interrupt the Ldcl tertiary structure. On the other side, the only
101 well characterised binding partner of Ldcl, the AAA+ ATPase RavA, is known to interact precisely
102 with the C-terminal β -sheet of Ldcl²⁷. Assembly of two copies of double-pentameric rings of Ldcl and
103 five copies of hexameric RavA spirals results in a huge 3.3 MDa macromolecular cage of intricate
104 architecture and largely unknown function^{27–29}. The Ldcl-RavA cage is proposed to assist assembly
105 of specific respiratory complexes in *E. coli* and to counteract acid stress under starvation conditions
106 by alleviating ppGpp-dependent inhibition of Ldcl^{30–32}. Thus, although these functions still require
107 further investigation, preservation of the RavA-binding propensity should be one of the criteria for
108 assessing the suitability of an Ldcl-FP fusion. Therefore, both N- and C-terminal fusion constructs
109 with either mGeosM or Dendra2_{T69A} attached to Ldcl via an appropriate linker were cloned into
110 dedicated plasmids and overexpressed at conditions optimised for Ldcl overproduction (see
111 Methods for details). Expression of fusion proteins was immediately detected by wide-field
112 fluorescence imaging that showed a similar distribution for the four fusions (Figure 1A). Each
113 construct was then purified in order to assess its structural integrity and RavA binding capacity *in*
114 *vitro*, with a goal of defining the most suitable construct for the subsequent creation of a
115 corresponding chromosomal fusion.

116 The N-terminal Dendra2_{T69A}-Ldcl fusion formed exclusively dimers, confirming the structure-
117 based prediction that the fluorescent tag at this position would perturb the dimer-dimer interaction
118 (Figure 1B, Supplementary Figure 1). Admittedly, native Ldcl was shown to dissociate into dimers *in*
119 *vitro* at pH above approximately 7.5, but in this pH range Ldcl is not supposed to be expressed in
120 the cell and therefore this dissociation may be irrelevant. Surprisingly, in contrast to Dendra2_{T69A}-
121 Ldcl, mGeosM-Ldcl assembled into regular symmetric non-native higher-order oligomers, with a
122 dramatically altered quaternary structure (Figure 1B, Supplementary Figure 1). These oligomers
123 were built of three Ldcl tetramers, bridged together by additional densities. Noteworthy, mEos2, the
124 fluorescent protein from which mGeosM was derived, crystallises as a tetramer (PDB ID: 3S05) that
125 can be straightforwardly fitted into the Ldcl-bridging densities (Figure 1B). This illustrates that despite
126 the fact that mGeosM was designed as a monomeric FP (with the first “m” explicitly standing for

127 monomeric), some residual oligomeric tendency is still maintained. This propensity of mGeosM to
128 oligomerise when bound to Ldcl may be driven by avidity effects – as Ldcl dimers begin to assemble
129 into a decamer, the local concentration of mGeosM increases to such a point that oligomerisation
130 becomes energetically favourable, despite the apparent monomer behaviour of mGeosM in gel
131 filtration²⁵. This is also in line with the known propensity of mEos2, from which mGeosM is derived,
132 to form tetramers at high concentration³³. To conclude, both N-terminal fusions induced non-native
133 assembly of Ldcl, and therefore neither were appropriate for determining the native localisation of
134 this enzyme in the cell.

135 As expected, C-terminal Ldcl fluorescent fusions formed decamers with protruding densities
136 that can be attributed to flexibly attached FPs (Figure 1B, Supplementary Figure 1). Nevertheless,
137 despite the long flexible linker between Ldcl and the FP, these constructs were unable to interact
138 with RavA as shown by Bio-Layer Interferometry (BLI) binding studies (see Methods and
139 Supplementary Figure 1 for details). This means that the functionality of these fusions cannot be
140 considered as entirely retained. Thus, none of the four fusions were suitable for an in-depth imaging
141 analysis under conditions of native Ldcl expression upon acid stress that we planned to undertake.
142 Altogether, we demonstrated that structural integrity and unaltered interaction with known partners
143 are useful read-outs for functional preservation. Based on this result we propose that, when feasible,
144 purification and structure-function analysis of FP fusions should be performed prior to interpretation
145 of the protein localisation inferred from observation of FP fusions by optical methods. This workflow
146 can be used for example in cases where chromosomal manipulation for assessment of intact function
147 is difficult and/or the phenotype is condition-dependent or unclear, whereas basic *in vitro* biochemical
148 and ns-EM characterisation are efficiently set up.

149 **Immunofluorescence of Ldcl in *E. coli* under acid stress reveals its supramolecular** 150 **organisation**

151 Because none of the FP fusions possessed the properties of native Ldcl, we decided to turn
152 to STORM imaging of the endogenous enzyme. One of the drawbacks of STORM is the requirement
153 for exogenous labelling and therefore the fixation and permeabilisation of cells. These procedures
154 are notoriously known to potentially affect cell morphology, and therefore, when possible, the usage
155 of live cell imaging with FP fusions (as for example in PALM), or observation of unfixed samples
156 under cryogenic conditions would be ideal. Moreover, fixation, permeabilisation and specific
157 exogenous labelling in bacteria present unique challenges because of the complex, often multi-
158 layered cell wall^{34,35}. A significant advantage of STORM however is the possibility of direct imaging
159 of the wild-type (WT) protein, which should circumvent the dangers associated with FP fusions
160 described above. The three prerequisites for imaging of WT systems by immuno-based labelling in
161 general and of Ldcl in particular are (i) availability of an antibody or nanobody (antibody fragment
162 derived from heavy-chain-only camelid antibody) coupled to an organic fluorescent dye and directed
163 towards the native protein, (ii) precise knowledge of endogenous expression conditions, (iii)
164 validation of an efficient permeabilisation and immunolabelling technique that enables the
165 antibody/nanobody to enter the cell and specifically target the protein of interest without altering the
166 native organization of the cell ultrastructures.
167

168 As a first step towards STORM imaging of endogenous LdcI, we probed an anti-LdcI
169 nanobody, hereafter called anti-LdcI-Nb. The complex between purified *E. coli* LdcI and the
170 nanobody was purified by size exclusion chromatography (see Methods), and imaged by ns-EM.
171 The resulting 3D map of the LdcI/anti-LdcI-Nb complex demonstrates binding of anti-LdcI-Nb to each
172 LdcI monomer in the decamer and reveals the location of the interaction site which is clearly distinct
173 from the RavA binding site (Figure 2A, Supplementary Figure 2). This nanobody was thus identified
174 as a suitable labelling agent for LdcI imaging in *E. coli* cells, and labelled with the fluorescent dye
175 AlexaFluor647 (AF647) or AlexaFluor488 (AF488) (see Methods).

176 Consistently with published data, LdcI expression could be induced by a pH shift experiment
177 (i.e. transfer of bacteria from pH 7.0 into a pH 4.6 growth medium) in the presence of lysine under
178 oxygen-limiting conditions (Supplementary Figure 3A). While the WT strain grew well under these
179 conditions and efficiently buffered the extracellular medium up to pH 6.2 in approximately 1.5 to two
180 hours (Supplementary Figure 3A) concomitantly with the increase in the level of LdcI expression
181 (Supplementary Figure 3B, C), the growth and the acid stress response of the Δ/dcl mutant strain
182 were severely impaired (Supplementary Figure 3A). Since the peak of LdcI expression by the wild
183 type cells was achieved between one and two hours after exposure to acid stress (Supplementary
184 Figure 3B, C), a time point of 90 minutes was chosen for the subsequent labelling and imaging
185 experiments. The specificity and performance of anti-LdcI-Nb in immunofluorescence labelling of
186 permeabilised *E. coli* cells were characterised by flow cytometry and wide-field fluorescence imaging
187 (Supplementary Figure 3D, E). Both techniques demonstrated that in the absence of LdcI expression
188 no specific fluorescence is seen, confirming the suitability of anti-LdcI-Nb for immunofluorescence
189 studies. Thus, the above-mentioned prerequisites for immuno-based imaging of cellular LdcI have
190 been fulfilled. Noteworthy, the LdcI expression profile highlighted a considerable asset of the usage
191 of STORM instead of PALM for visualisation of the endogenously expressed LdcI: indeed, the
192 transient nature of the expression and the necessity of work under oxygen-limiting conditions may
193 have created difficulties due to the longer maturation time of the FPs under these conditions.

194 Initial characterisation of the cellular distribution of LdcI 90 minutes after exposure of *E. coli*
195 cells to acid stress was carried out by wide-field fluorescence imaging. Based on these images, it
196 appeared that natively expressed LdcI did not display a homogeneous cytoplasmic distribution but
197 rather showed a patchy localisation pattern (Figure 2B). 3D STORM imaging subsequently provided
198 a more detailed view of this patchy distribution. As shown in Figure 3, Supplementary Figure 4, and
199 Supplementary Movies 1-5, the labelling density was lower in the centre of the bacterium. This
200 indicates a propensity for LdcI to cluster near the cell periphery, towards the inner membrane and
201 the cell poles, rather than being distributed homogeneously through the volume. Recent
202 investigations argue that, in most cases, *in vivo* clustering of metabolic enzymes corresponds to their
203 polymerised states, and represents an efficient means of regulation of enzymatic activity and
204 metabolic homeostasis in response to a stimulus¹. Thus, structure determination of these polymers
205 is a crucial step towards the understanding of regulation mechanisms. Using ns-EM, we previously
206 documented that at low pH and high concentration, LdcI decamers tend to stack⁶. Therefore, the
207 high local concentration of LdcI, clustered in patches in the *E. coli* cell under acid stress conditions
208 boosting LdcI expression, is likely to induce enzyme polymerisation via stack formation. Remarkably,
209 the images convey a visual impression that individual patches tend to be further arranged into bands

210 or helix-like turns, showing a circumferential distribution with a sort of a long-range stripy or pseudo-
211 helical organisation (Figure 3, Supplementary Figure 4, Supplementary Movies 1-5).

212 213 **Structural determinants of Ldcl stack formation revealed by cryo-EM**

214 In order to understand the molecular determinants of Ldcl polymerisation at low pH and to
215 provide a framework for the analysis of Ldcl function under acid stress, we solved the 3D structures
216 of the Ldcl decamer at pH 7.0 and of Ldcl stacks at pH 5.7 by cryo-EM (See Methods, Figure 4,
217 Supplementary Figures 5-6, Supplementary Table 1). The 2.8 Å resolution structure of the Ldcl
218 decamer at pH 7.0 (Figure 4A) is extremely similar (Supplementary Table 2 & 3) to the Ldcl crystal
219 structure solved at pH 8.5 in an inhibited ppGpp-bound state⁶. However, contrary to pH 7.0 and even
220 pH 6.2 where Ldcl is still predominantly decameric¹⁰, Ldcl forms straight rigid filaments on the cryo-
221 EM grid at pH 5.7, which corresponds to the pH of maximum Ldcl enzymatic activity (Figure 4B).
222 The structure of a three-decamer stack was solved to a resolution of 3.3 Å, revealing the structural
223 details of acid stress-induced Ldcl polymerisation (Figure 4C). Ldcl decamers stack tightly on top of
224 one other, with negligible rotation between decamers along the stack. Each dimer fits snugly in the
225 inter-dimer groove of decamers above and below.

226 A comparison of Ldcl decamer structures taken from the Ldcl stack cryo-EM map (at pH 5.7),
227 the Ldcl decamer cryo-EM map (at pH 7.0) and the crystal structure of decameric Ldcl crystallised
228 with bound ppGpp at pH 8.5 (PDB ID: 3N75) reveals some remarkable differences between the stack
229 structure and the two decamer structures (Figure 4D, E, Supplementary Table 2 & 3). While the three
230 structures do not show any major differences at the monomer level, a structural alignment of an Ldcl
231 dimer extracted from the Ldcl decamer structures with a dimer extracted from the Ldcl stack structure
232 uncovers a rigid body-like rotation between monomers around a hinge region located at the
233 monomer-to-monomer interface (Supplementary Table 2 & 3, Figure 4DE). This rotation results in a
234 5° tilt when comparing the N-terminal wing-domains in Ldcl dimers, and an overall slightly decreased
235 diameter of the central cavity inside the stacked Ldcl rings (Figure 4D, E), which may contribute to
236 the tight packing of each dimer into the grooves of an opposing decamer in the stack.

237 A careful examination of the Ldcl stack structure shows that two major inter-decamer
238 interfaces situated at a two-fold symmetry axis perpendicular to the stack direction contribute to stack
239 formation (Figure 5A). In particular, the first interface (Figure 5B) is formed between residues K422,
240 D460, R468, D470, and E482 situated in the ppGpp-binding domain (amino acids 418-564), and
241 residues N314, D316, and G352 from the PLP-binding domain (amino acids 184-417). D460 from
242 one decamer makes an electrostatic interaction with K422' of a neighbouring decamer in the stack.
243 R468 is sandwiched between D316' and E344' from a neighbouring decamer, and makes
244 electrostatic interactions through its η 1 and η 2 nitrogen atoms with D316'. In addition, D470 interacts
245 with the backbone of G352', and E482 forms hydrogen bonds with N314'. The second interface
246 (Figure 5B) is formed between residue N94 of the wing domain (amino acids 1-130) of one set of
247 opposing dimers, and a stretch of four residues in the ppGpp-binding domain – T444, E445, S446
248 and D447 - at the end of helix α 16 from a second set of opposing dimers. The wing domain residue
249 N94 makes hydrogen bonds with E445' of an opposing dimer. A second charged residue, D447,
250 interacts with the backbone of T444', and is held in place by R97 from the wing domain of a
251 neighbouring dimer in the same decamer.

252 Considering that the Ldcl polymerisation is induced by acid stress, we wondered which
253 residues in the interface would be sensitive to pH changes. Surprisingly, most of the side chains
254 involved in the inter-decamer interface are charged arginine (pKa 13), aspartate and glutamate
255 residues (pKa of 4 and 3 respectively), which do not change protonation state in the pH window
256 relevant for Ldcl activity (pH 5-7)³⁶. Nonetheless, other residues, situated outside the interaction
257 interfaces, may drive stack formation through pH-dependent interactions that would in turn lead to
258 the observed inter-monomer rotation and the associated constriction of the Ldcl central cavity,
259 coupled to the alignment of complementary contacts at interfaces one and two (Figure 4D, E). We
260 note for example that H694 should be protonated in the Ldcl stack structure at pH 5.7 but
261 deprotonated in the two ring structures at pH 7.0 and pH 8.5, and that an electrostatic interaction
262 between H694 and D192 situated in the linker region is present in the stack structure only (Figure
263 5E).

264 To validate the observed interactions at the inter-decamer interface, and to assess the
265 individual importance of key residues involved in Ldcl stack formation, we constructed four Ldcl point
266 mutants (R468E, R97E, H694A and H694N), two double mutants (R97E/R468E and E445A/D447A)
267 and one triple mutant (E445A/D447A/R468E). The mutants were purified following the protocol for
268 wild-type Ldcl (see Methods), diluted into a buffer at pH 5.7 and observed by ns-EM (Figure 6,
269 Supplementary Figure 7). Although the grid preparation procedure for ns-EM yields stacks that are
270 shorter and more curved and distorted when compared to the cryo-EM data (Figure 4B, Figure 6),
271 our previous observations of the five paralogous *E. coli* amino acid decarboxylases justify the validity
272 of this approach for a qualitative comparative analysis^{6,36}. Ns-EM images make immediately
273 apparent that the E445A/D447A double mutant does not have a significantly altered capability of
274 stack formation at pH 5.7 when compared to WT Ldcl, whereas the single R468E mutation is
275 sufficient to completely abolish stack formation. The R97E mutant has a moderate destabilising
276 effect and displays fewer and smaller stacks than the WT Ldcl. Consistently with the major role of
277 R468 in the Ldcl stack formation, the R97E/R468E double mutant and E445A/D447A/R468E triple
278 mutant exclusively occur as decamers at low pH. Altogether, these results reveal that R468 is one
279 of the key determinants of Ldcl stack formation. Finally, similarly to the R97E mutant, a modest
280 destabilisation of stack formation is observed for the two histidine mutants, H694A and H694N,
281 favouring our hypothesis that H694 may have an influence on the propensity of Ldcl polymerisation
282 at low pH (Figure 6, Supplementary Figure 7).

283 284 Discussion

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

293 The high resolution cryo-EM structures of the stack at pH 5.7 and of the decamer at pH 7.0
294 complement the previously solved crystal structure of the ppGpp-bound Ldcl decamer. Previous
295 serendipitous co-crystallisation of Ldcl with the nutrient stress response alarmone ppGpp led to
296 assessment of the effect of ppGpp on Ldcl activity, and to a proposal that ppGpp would act as an
297 Ldcl inhibitor that prevents excessive lysine consumption upon nutrient limitation during acid stress⁶.
298 In addition, a ppGpp-dependent disassembly of the Ldcl stacks had been previously observed but
299 could not be structurally explained since the ppGpp binding site is situated between two neighbouring
300 dimers inside the Ldcl ring⁶. Our cryo-EM structures show that one of the residues involved both in
301 ppGpp binding and in the stack formation is arginine 97 (R97). In the crystal structure of ppGpp-
302 bound Ldcl, R97 makes a stacking interaction with the guanosine imidazole ring of ppGpp, while in
303 Ldcl stacks R97 is involved in a key interaction at the second interface (Figure 5D), where it locks
304 D447 in a conformation allowing interactions between helices α 16 from opposing Ldcl decamers.
305 Binding of ppGpp to Ldcl interferes with the R97-D447 interaction, thereby most likely prohibiting
306 correct positioning of D447 at the tip of helix α 16, and resulting in a disruption of the second stack
307 interface, leading to a moderate yet notable stack destabilisation (Figure 6, Supplementary Figure
308 7). Furthermore, our cryo-EM structure of ppGpp-free Ldcl decamers at neutral pH enables to rule
309 out the effect of ppGpp on the differences observed between the Ldcl stack structure at pH 5.7 and
310 the ppGpp-Ldcl crystal structure at pH 8.5. Indeed, despite the absence of ppGpp, R97 is still
311 oriented towards the ppGpp binding site and away from the inter-decamer interface in the pH 7.0
312 decamer cryo-EM map (Figure 5D). This suggests that the conformational changes in Ldcl driving
313 stack formation are mostly driven by low pH and not by the absence of ppGpp, although the D192
314 and H694 hinge residues are similarly far apart in the pH 8.5 and pH 7.0 structures (Figure 5E). Our
315 current work provides a solid experimental and structural basis for a future closer evaluation of the
316 hypothesised role of ppGpp in Ldcl regulation *in vitro* and *in vivo*⁶

317 The clusters of endogenous Ldcl at the bacterial periphery observed by optical imaging would
318 presumably correspond to Ldcl assembled into stacks. Possible reasons for such assembly may be
319 to provide an efficient way to locally increase the Ldcl concentration and to enhance its activity. But
320 why would Ldcl, an apparently highly soluble protein, be driven towards the inner membrane? What
321 would be the advantage for acid-stressed *E. coli* cells to increase the concentration of Ldcl via stack
322 formation in these particular peripheral locations? Localisation of proteins to specific sites in the
323 bacterial membrane was shown to be generally driven by chemical factors such as the phospholipid
324 composition of the lipid microdomains, and by physical factors such as the degree of local curvature
325 or the electric potential of the membrane³⁷. An attractive hypothesis would be that as an acid stress
326 response protein performing a proton-consuming enzymatic reaction, Ldcl may be attracted to proton
327 sinks formed by anionic phospholipids which compartmentalise oxidative phosphorylation
328 (OXPHOS) complexes for efficient functioning in bacterial respiration and adaptation to
329 environmental changes. Indeed, OXPHOS complexes were often described to be unevenly
330 distributed in the membrane in the form of mobile patches³⁸⁻⁴², providing evidence for highly dynamic
331 and spatially organised bioenergetic membranes in *E. coli* cells⁴². In addition, certain bacterial
332 flotillins, which are essential scaffold proteins of the functional membrane microdomains, equivalent
333 to the lipid rafts of eukaryotic cells, also show a patchy distribution and were shown to interact with
334 specific OXPHOS complexes^{42,43}. In this regard, two different lines of evidence would be interesting

335 to note. First, Ldcl was described to co-purify with a partially assembled Complex I⁴⁴, whereas the
336 Ldcl-binding partner RavA, as well as ViaA, the second protein expressed from the *ravAviaA* operon
337 and which also interacts with RavA, were shown to interact with both Complex I and fumarate
338 reductase^{31,32}. Second, the other *E. coli* PLP-dependent lysine decarboxylase LdcC, exercising the
339 role of cadaverine biosynthesis irrespectively of acid stress⁴⁵, neither binds RavA nor forms
340 stacks^{28,36}, in spite of its 69% of identity with Ldcl. It is interesting to note that in LdcC the key stack-
341 forming residue R468 has been substituted for an alanine while the interacting D316 has been
342 preserved.

343 The reason for the arrangement of the Ldcl clusters into stripes or pseudo-helical patterns is
344 also intriguing. While similar distributions have been documented for bacterial cytoskeletal, cell
345 division, chromosome partitioning, RNA degradation and secretion machineries, an eventual impact
346 of labelling on these distributions, demonstrated specifically for the YFP-MreB⁴⁶, warrants caution in
347 their interpretation^{12,14}. Here however, we observed endogenous wild-type Ldcl in cells fixed prior to
348 their permeabilisation and for Ldcl labelled with anti-Ldcl-Nb, which means that the resulting pattern
349 is likely real and not an artefact. In addition, examination of some published images of OXPPOS
350 patches (for example Llorente-Garcia et al., 2014) also seem to hint to a possible pseudo-helical
351 organisation. Excitingly, anionic phospholipid-specific dyes and fluorescently-labelled antibiotics
352 specific for nascent peptidoglycan synthesis upon cell elongation were also described to be
353 distributed on helical or stripe patterns^{37,47-50}. It is therefore tempting to imagine that inside the
354 bacterial cell, Ldcl has a tendency to follow a general path upon polymerisation governed by the
355 underlying patterns in the cell envelope, which are proposed to follow a global right-handed chiral
356 order⁵¹.

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

363 **Methods**

364 **Expression constructs**

365 For fluorescence studies, several FP fusion constructs were generated starting from an
366 available plasmid containing the coding sequence of Ldcl (Uniprot entry P0A9H3), cloned in the
367 pET22b(+) vector with a C-terminal 6xHis-tag⁶. All constructs were generated using the Gibson
368 cloning strategy and verified by sequencing analysis. The Gibson assembly was performed using
369 0,4U T5 exonuclease (NEB M0363S), 2,5U Phusion polymerase (NEB M0530S) and 400U Taq
370 ligase (NEB M0208L) in 1X ISO buffer (100mM Tris-HCl pH 7,5, 10 mM MgCl₂, 0.8 mM dNTP mix,
371 10mM DTT, 50 mg PEG-8000, 1 mM NAD). 7.5 µL of the GIBSON Master Mix was mixed with 2.5
372 µL DNA, containing circa 100 ng of vector. The mix was incubated for 60 min at 50°C.
373 Transformations were performed in Top10 competent bacteria (One Shot™ TOP10 Chemically
374 Competent *E. coli*, Invitrogen C404003) and selected using 100 µg/mL ampicillin or 50 µg/mL
375 kanamycin sulphate from Euromedex (Ampicillin ref EU0400-D, Kanamycin ref EU0010-D). Agarose
376

377 Gel purification and DNA plasmid extraction kits were purchased from Macherey-Nagel (NucleoSpin
378 Plasmid ref 740588-250 and NucleoSpin Gel and PCR Clean-up ref 740609-250).

379 Dendra2_{T69A}-Ldcl and mGeosM-Ldcl were both cloned in the pET-TEV vector containing an
380 N-terminal 6xHis-tag, and a TEV cleavage site between Dendra2_{T69A} or mGeosM and the Ldcl gene.
381 Ldcl-Dendra2_{T69A} and Ldcl-mGeosM were both cloned in the pET22b(+) vector with Ldcl followed by
382 either Dendra2_{T69A} or mGeosM containing an uncleavable C-terminal 6XHis-tag.

383 The sequence of a anti-Ldcl-Nb was cloned in the pHEN6 vector containing the pelB leader
384 sequence from *Erwinia carotovora* for secretion into the periplasm, and a C-terminal 6xHis-tag
385 (kindly provided by Dr. Aline Desmyte, AFMB Marseille).

386 Plasmids, primers and cloning strategy are summarised in Supplementary Table 4.

387

388 **Protein purification**

389 Ldcl-FP fusions and Ldcl mutants were expressed in BL21(DE3) cells grown in LB medium
390 supplemented with 100 µg/mL ampicillin or 50 µg/mL kanamycin sulphate. Protein expression was
391 induced using 40 µM IPTG (Euromedex EU008-C) and carried out overnight at 18°C. The Ldcl-FP
392 fusions and Ldcl mutants were purified as previously described for wild-type Ldcl^{27,28,36}, in a final
393 buffer containing 25 mM Tris (pH 7.5), 0.3 M NaCl, 5% glycerol, 1 mM DTT and 0.1 mM PLP.

394 The anti-Ldcl-Nb was expressed in *E. coli* WK6 cells following the protocol described by⁵²,
395 and purified with immobilised metal affinity chromatography (IMAC, using a Ni-NTA column) followed
396 by Size exclusion chromatography (SEC) using a a superdex 75 Increase 10/300GL column (GE-
397 Healthcare, ref: 29-1487-21) equilibrated with a buffer containing 25 mM Tris pH 7.4 and 0.3 M NaCl.

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

401

402 **Ns-EM on Ldcl-FP fusions, Ldcl/anti-Ldcl-Nb and Ldcl mutants**

403 FP fusion samples after gel filtration were diluted to a concentration of approximately 0.025
404 mg/mL. 3 µL was applied to the clean side of carbon on a carbon–mica interface and stained with
405 2% uranyl acetate (mGeosM-Ldcl, Ldcl-mGeosM, Ldcl-Dendra2_{T69A}, Ldcl/anti-Ldcl-Nb, Ldcl
406 mutants) or 2% sodium silicotungstate (Dendra2_{T69A}-Ldcl). Images were collected on a 120 kV
407 Tecnai T12 microscope with an Orius 1000 camera (Gatan) or on a 200 kV Tecnai F20 electron
408 microscope with either a OneView camera (Gatan) or a Ceta camera (Thermo Scientific). All images
409 were collected with a defocus range of approximately -1.0 µm to -2.0 µm and with pixel sizes between
410 2.29 Å/pixel and 3.42 Å/pixel.

411

412 **Image processing – Ldcl-FP and Ldcl/anti-Ldcl-Nb**

413 36 micrographs of Dendra2_{T69A}-Ldcl with a pixel size of 2.73 Å/pixel, 368 micrographs of
414 mGeosM-Ldcl with a pixel size of 2.29 Å/pixel, 23 micrographs of Ldcl-Dendra2_{T69A}, 92 micrographs
415 of Ldcl-mGeosM with a pixel size of 3.42 Å/pixel, and 124 micrographs of Ldcl/anti-Ldcl-Nb with a
416 pixel size of 2.82 Å/pixel were used for image analysis.

417 CTF estimation was performed with CTFFIND3⁵³. Semi-automatic particle selection was
418 carried out with BOXER⁵⁴, with box sizes of 98 x 98 pixels for Dendra2_{T69A}-Ldcl, 180 x 180 pixels for

419 mGeosM-Ldcl, 128 x 128 pixels for Ldcl-Dendra2_{T69A} and Ldcl-mGeosM, and 112 x 112 pixels for
420 Ldcl/anti-Ldcl-Nb respectively. Particle extraction followed by several rounds of cleaning by 2D
421 classification in RELION-1.4⁵⁵, resulted in the following number of particles for each dataset:
422 Dendra2_{T69A}-Ldcl = 7140, mGeosM-Ldcl = 5514, Ldcl-Dendra2_{T69A} = 832, Ldcl-mGeosM = 12,211
423 and Ldcl/anti-Ldcl-Nb = 14,075.

424 For Dendra2_{T69A}-Ldcl, initial model generation was carried out in RELION-2.1⁵⁶ without any
425 symmetry applied. For mGeosM-Ldcl, initial model generation was carried out in RELION-2.1 with
426 either C1, C3 or D3 symmetry applied. The results of all three calculations being very similar, the
427 model with applied D3 symmetry was selected. For Ldcl-Dendra2_{T69A}, Ldcl-mGeosM and Ldcl/anti-
428 Ldcl-Nb, the previously-determined Ldcl decamer structure (PDB ID: 3N75) was filtered to 60 Å and
429 used as an initial model for 3D refinement.

430 3D refinement was carried out for each dataset with applied C2 symmetry for Dendra2_{T69A}-
431 Ldcl, D3 symmetry for mGeosM-Ldcl, and D5 symmetry for Ldcl-Dendra2_{T69A}, Ldcl-mGeosM and
432 Ldcl/anti-Ldcl-Nb. Rigid body fitting of Ldcl (PDB ID: 3N75), mEos2 (PDB ID: 3S05) and Dendra2
433 (PDB ID: 2VZX) crystal structures was then carried out in Chimera⁵⁷ for the five datasets.

434

435 **Cryo-EM on Ldcl stacks (pH 5.7)**

436 Wild-type Ldcl was purified as previously described¹⁰ from an *E. coli* strain impaired in the
437 production of ppGpp (MG1655 Δ relA Δ spoT) in order to avoid any serendipitous ppGpp binding.
438 Purified Ldcl was diluted to a final concentration of approximately 0.25 mg/mL in a buffer containing
439 25 mM MES (pH 5.7), 0.3 M NaCl, 5% glycerol, 1 mM DTT and 0.1 mM PLP. 3 μ L of the sample
440 was applied to a glow-discharged R2/1 300 mesh holey carbon copper grid (Quantifoil Micro Tools
441 GmbH) and plunge-frozen in liquid ethane using a Vitrobot Mark IV (FEI) operated at 100% humidity.
442 Datasets were recorded at the European Synchrotron Radiation Facility (ESRF) in Grenoble,
443 France⁵⁸, on a Titan Krios microscope (Thermo Scientific) equipped with a BioQuantum LS/967
444 energy filter (Gatan) and a K2 summit direct electron detector (Gatan) operated in counting mode. A
445 total of 2564 movies of 30 frames were collected with a total exposure of 6 s, total dose of 29.3 e⁻/Å²
446 and a slit width of 20 eV for the energy filter. All movies were collected at a magnification of 130,000x,
447 corresponding to a pixel size of 1.052 Å/pixel at the specimen level. A summary of cryo-EM data
448 collection parameters can be found in Supplementary Table 1.

449

450 **Image Processing – Ldcl stacks (pH 5.7)**

451 Motion correction and dose-weighting of the recorded movies were performed using
452 MotionCor2⁵⁹. CTF parameters were determined on the aligned and dose-weighted sums using
453 CTFFIND4⁶⁰. After manual inspection of the dose-weighted sums, the best 558 (21.7%) micrographs
454 were selected for further processing. Ldcl stacks were manually picked using e2helixboxer in
455 EMAN2⁶¹. A total of 15,165 Ldcl-stack particles were extracted in RELION-3.0⁶² with an extract size
456 of 320 pixels, resulting in boxes containing three Ldcl decamers, and with the --helix option with an
457 outer diameter of 160 pixels and a helical rise of 77 pixels. After particle extraction, per-particle CTF
458 correction was performed using Gctf⁶³. Extracted particles were subjected to 2D classification in
459 RELION-3.0, resulting in a cleaned dataset containing 15,157 particles. Initial 3D refinement with
460 imposed D5 symmetry was carried out in RELION-3.0, using an initial model generated by manually

stacking three Ldcl decamers (PDB ID: 3N75) in Chimera⁵⁷ and low-pass filtering the resulting Ldcl stack to 40 Å. The resulting 4.3 Å resolution 3D reconstruction, along with the cleaned particle stack, was subsequently imported into CryoSPARC⁶⁴. A final homogeneous 3D refinement in CryoSPARC, using a dynamic mask and imposing D5 symmetry, resulted in a map with a resolution of 3.28 Å based on the 0.143 gold-standard Fourier shell correlation (FSC) criterion⁶⁵. 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 image processing steps for Ldcl stacks can be found in Supplementary Table 1 and Supplementary Figure 4, with local resolution and FSC curves shown in Supplementary Figure 5.

Cryo-EM on Ldcl decamers (pH 7.0)

Purified Ldcl was dialysed into a buffer at pH 7.0 and diluted to a concentration of 0.25 mg/mL. 3 µL of diluted sample was applied to a glow-discharged (R2/1 300 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. Images were recorded on a Glacios microscope (Thermo Scientific) equipped with a Falcon II direct electron detector (Thermo Scientific). A total of 2772 movies of 29 frames were collected with a total exposure of 1.5 s and a total dose of 45 e⁻/Å². All movies were collected at a magnification of 116,086x, corresponding to a pixel size of 1.206 Å/pixel at the specimen level.

Image Processing – Ldcl decamers (pH 7.0)

Motion correction was carried out using patch motion correction in CryoSPARC, discarding the first two frames. Initial CTF estimation was then carried out on summed frames using CTFFIND4. A subset of 600 micrographs were subjected to automatic picking using the blob picker in CryoSPARC, resulting in ~238,000 picked particles. Particles were then extracted with a box size of 256 x 256 and subjected to 2D classification. Particles from the best classes showing clear secondary structural features for Ldcl were selected for homogeneous refinement (with applied D5 symmetry) against EMD-3204 low-pass filtered to 30 Å, resulting in a reconstruction with a resolution of 4.2 Å (FSC = 0.143). This reconstruction was then used to create templates for picking the entire dataset using the template picker in CryoSPARC, after filtering to 12 Å. ~796,000 particles were extracted and subjected to 2D classification, and the best ~428,000 particles were subjected to heterogeneous refinement with applied D5 symmetry against the 4.2 Å map, resulting in one higher-resolution class corresponding to ~229,000 particles. These particles were subjected to homogeneous refinement with applied D5 symmetry, resulting in a map with a resolution of 2.76 Å (FSC = 0.143) which was then sharpened with a B-factor of -173 Å². A summary of cryo-EM data collection parameters and image processing steps for Ldcl decamers can be found in Supplementary Table 1 and Supplementary Figure 4, with local resolution and FSC curves shown in Supplementary Figure 5.

Fitting of structures and refinement

For fitting of atomic models in the 3D reconstructions of Ldcl at pH 5.7 (stack) or pH 7.0 (decamer), two (for the stack) or one copy (for the decamer) of the Ldcl X-ray crystal structure (PDB

503 ID: 3N75) were first rigid-body fitted in the corresponding 3D reconstructions using Chimera.
504 Refinement was performed using the Phenix software package⁶⁶ and was identical for both 3D
505 reconstructions. A first round of real space refinement was carried out with enabled rigid-body, global
506 minimization, local grid search and ADP refinement parameters, and imposing rotamer,
507 Ramachandran, NCS and reference model (PDB ID: 3N75) restraints. A final round of real space
508 refinement was then performed using the same settings, but without rigid body refinement and
509 without applying reference restrains, setting the 'nonbonded_weight' parameter to 4000 and
510 disabling 'local_grid_search'. A summary of refinement and model validation statistics can be found
511 in Supplementary Table 1.

512

513 **pH shift experiment**

514 Stationary phase cultures, which were grown overnight from single colonies in LB medium,
515 were diluted to OD₆₀₀ ~ 0.01 and re-grown at 37°C within approximately 1h 45 min in fresh LB
516 medium to an OD₆₀₀ of 0.1. From this culture, 14 mL were transferred to 15 mL falcon tubes and
517 pelleted by centrifugation at RT for 5 min. The supernatant was decanted, whereby systematically
518 around 200 µL LB remained in the falcon tube. The pellet of bacteria was resuspended and
519 afterwards, LB-4.6 containing 30 mM L-lysine was added to the cells up to 14 mL. To prepare LB-
520 4.6 medium, LB powder (Sigma-Aldrich) was completely dissolved in distilled water by stirring for 30
521 min. The pH of 4.6 was then adjusted using HCl. After autoclaving, sterile filtered L-lysine was added
522 to LB-4.6. For this, L-lysine was dissolved in an aliquot of LB-4.6, sterile filtered and mixed with the
523 remaining LB-4.6. 30 mM L-lysine were used. In order to grow the culture under oxygen-limiting
524 conditions, the lid of the falcon tube was closed and the tubes were placed at 37°C on a shaker (150
525 rpm). After defined time-points, aliquots were taken for OD₆₀₀ measurement, pH measurement,
526 SDS-PAGE or immunofluorescence. For each time-point, a new tube was opened and not reused
527 further.

528

529 **Western Blotting**

530 SDS-PAGE was performed with a Biorad electrophoresis chamber using standard 12% reducing
531 SDS-PAGE gels. Proteins were transferred to a nitrocellulose membrane (Biorad) using a Trans-
532 Blot Turbo Transfer System (Biorad). The membrane was blocked for 1h using 5% BSA in TBS
533 supplemented with 0.1% Tween (TBS-T). Afterwards, the membrane was incubated for 1h with an
534 anti-Ldcl antibody (Qalam-Antibodies) in BSA/TBS-T (1:5000). The membrane was subsequently
535 washed 3 x 10 min in TBS-T and incubated for 1 h with HRP-coupled anti-rabbit antibody (1:10000
536 in BSA/TBS-T). Finally the membrane was washed 3 x 10 min in TBS-T. The membrane was rinsed
537 once with TBS, prior to detection of antibody-labelled proteins using ECL reagent (GE Healthcare).
538 Incubation and detection of antibody was performed at 25 °C.

539

540 **Nanobody labelling**

541 For the labelling reaction, 50 µL nanobody (i.e. 200 µg) was pipetted into a 1.5 mL tube,
542 placed on ice and supplemented with 5.5 µL of 1 M bicarbonate buffer at pH 8.3. 100 µg of Alexa-
543 647 or Alexa-488 NHS ester dye (Life Technologies, A37573 and A20000) were dissolved in 10 µL
544 DMSO to final concentration of 10 mg/mL. 5 µL (i.e. 40 nmol) dye in DMSO was added to the protein

545 and incubated for 1 h at room temperature in a shaking block, covered with an aluminium foil to
546 protect the dye from the light. Excess dye was removed by iterative buffer exchange using a 3K spin
547 column (Amicon-Ultra-4 Centrifugal Filters Ultracell 3K, Millipore UFC800396) to PBS. The degree
548 of labelling was inferred from measuring the OD.

549 **Cell preparation for immunofluorescence staining of *E. coli* cells with nanobodies**

550 For immunofluorescence staining, the OD600 of the cell culture was measured, and the
551 volume of cells corresponding to OD = 4, with OD = 1 corresponding to about 8×10^8 cells per mL,
552 were collected by centrifugation. After removal of LB by pipetting, cells were resuspended in 2 mL
553 of 4 % FA in PBS (made from 16% Formaldehyde solution, Methanol-free from Thermo Scientific).
554 Falcons were placed on a rotor for constant agitation for 45 min at room temperature. After fixation,
555 cells were collected by centrifugation and the solution was removed by pipetting. Cells were then
556 resuspended in 14 mL PBS (Gibco, Thermo Scientific) to remove and dilute the fixative. Cells were
557 permeabilised for 10 min using 2 mL 0.1 % Triton X-100 in PBS and subsequently washed three
558 times with 10 mL PBS. Finally, cells were transferred to 1.5 mL tubes, centrifuged and resuspended
559 in 200 μ L 1% BSA/PBS (BSA/PBS solution was dissolved for 30 min and sterile-filtered to avoid
560 clumps). After 30 min of incubation, 0.5 μ g anti-Ldcl-Nb, labelled with the dedicated dye, was added
561 to the 200 μ L bacteria-BSA/PBS suspension. Cells were incubated with the labelled anti-Ldcl-Nb for
562 16 h at 4°C. The next day, cells were washed three times with 1 mL PBS, centrifuged to remove
563 antibody solution and resuspended in 250 μ L PBS. When needed, Hoechst 33342 (Sigma-Aldrich)
564 was added to a final concentration of about 100 ng/mL.

565 **Wide-field imaging and flow cytometry**

566 For epifluorescence imaging, 2 μ L of cells were placed between a glass slide and a coverslip, which
567 have been carefully pressed together, and observed using an inverted IX81 microscope, with a
568 UPLFLN 100X oil immersion objective (N.A. 1.3) (Olympus), using the appropriate specific excitation
569 and emission filters for AF488 (GFP-3035B set, Semrock) and DAPI (DAPI-5060B set, Semrock).
570 Acquisitions were performed with Volocity software (QuorumTechnologies™) with a sCMOS
571 2048x2048 camera (Hamamatsu ORCA Flash 4, 16 bits/pixel) achieving a final magnification of 64
572 nm per pixel. For flow cytometry, 50 μ L of cells suspensions were injected in a MACSQuant VYB
573 flow cytometer (Miltenyi Biotech, Bergish Gladbach, Germany) using the 488 nm excitation and
574 525(50) nm emission channel (B1). AF488 positive populations were estimated after forward scatter
575 (FSC) and side scatter (SSC) gating on the cells. Data were further processed with MACSQuantify
576 software (Miltenyi Biotech).

577 **STORM imaging**

578 For Single Molecule Localization Microscopy, cells were transferred to a glucose buffer containing
579 50 mM NaCl, 150 mM Tris (pH 8.0), 10% Glucose, 100 mM MEA (Mercaptoethylamine) and 1x Glox.
580 Glox was prepared as a 10x stock and contained 1 μ M catalase and 2.3 μ M glucoseoxidase. Menzel
581 glass slides (Thermo Scientific) and precision coverslips (1.5H from ThorLabs) were cleaned for 30
582 min using an UV-ozone cleaning device (HELIOS□500, UVOTECH Systems). 2 μ L of immuno-
583 labelled cells were placed onto a glass slide and covered with the coverslip, then cells were carefully
584
585
586

587 spread by pressing glass slides firmly together and the sides were sealed with transparent nail polish
588 to avoid evaporation. Mounted samples were imaged on a homemade SMLM set up based on an
589 IX81 microscope (Olympus). STORM was performed by focusing a 643 nm excitation laser beam
590 (Toptica Diode laser) to the back focal plane of an oil immersion UAPON100X (N.A. 1.49) objective.
591 The intensity of the laser was tuned using an Acousto-Optical Tunable Filter (OATF, Quanta Tech).
592 Acquisition was obtained with a 16 bits/pixel Evolve 512 EMCCD (Photometrics) using Metamorph
593 (Molecular Devices), for a final pixel size of 123 nm. 3D-STORM based on point-spread-function
594 astigmatism⁶⁷ was performed using a cylindrical lens (LJ1516L1-A, Thorlabs) placed in the detection
595 light path. STORM datasets consisting of about 30,000 frames, using a 643 nm laser power density
596 of 3 kW/cm² and a 405 nm laser power density of up to 1 W/cm² and a frametime of 50 ms, were
597 acquired with an EMCCD gain set to 200. 3D point spread function calibration was achieved using
598 tetraspec beads. Finally, data were processed with the Thunderstorm plugin⁶⁸ in ImageJ⁶⁹. 3D-
599 images were rendered with Visp⁷⁰ using a minimum neighbour density threshold of 20 to 28.

600

601 **Bilayer interferometry measurements**

602 For BLI binding studies, RavA with a biotinylated C-terminal AviTag was expressed and purified as
603 previously described²⁹. BLI experiments were performed in 1x HBS pH at 7.0 (25 mM HEPES, 300
604 mM NaCl, 10 mM MgCl₂, 10% glycerol) supplemented with 1x kinetics buffer (0.1% w/v BSA, 0.02%
605 v/v Tween-20), 1mM ADP, 1mM DTT and 0.1 mM PLP. Experiments were performed using the BLItz
606 System instrument (FortéBio), operated at room temperature. Before the start of each BLI
607 experiment, RavA-AviTag was incubated with 1 mM ADP for 10 min. Streptavidin-coated biosensors
608 (FortéBio) were functionalised with biotinylated RavA-AviTag, then quenched with 10 µg/mL biocytin.
609 Experiments with the wild-type Ldcl are from²⁹. For C-terminal Ldcl-FP fusions, pins were dipped in
610 wells containing a range of Ldcl-FP concentrations from 0 to 1000 nM, with no binding signal
611 recorded at any concentration of Ldcl-FP.

612

613 **Data and code availability**

614 Cryo-EM maps, along with the corresponding fitted atomic structures, have been submitted to the
615 EMDB and PDB with accession codes EMD-10850 and PDB-6YN6 for Ldcl stacks (pH 5.7), and
616 EMD-10849 and PDB-6YN5 for Ldcl decamers (pH 7.0).

617

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636

637 **Author Contributions**

638 C.L., M.B., V.A., A.F. and K.H. cloned constructs, M.J., A.F. and K.H. purified proteins. M.J. and I.G.
639 performed ns-EM imaging and analysis. A.D., G.E. and M.B.-V. performed cryo-EM imaging. J.F.
640 and A.D. performed cryo-EM analysis. M.J. and J.F. built models resulting from EM maps. M.J., J.F.
641 and I.G. analysed structures and interpreted data. C.L. and M.B. performed optical imaging of
642 overexpression constructs with input of V.A., J.-P.K. and D.B. C.L. performed nanobody
643 characterisation for optical imaging with input from J.-P.K. C.L. analysed endogenous expression
644 and performed optical imaging of endogenous LdcI with input of V.A., J.-P.K. and D.B. C.L. and D.B.
645 analysed STORM Images. M.J., C.L., J.F. and I.G. analysed the data and prepared the figures and
646 tables. C.L. and D.B. contributed to the design of the optical imaging part of the project together with
647 I.G. I.G. designed, supervised and funded the overall study. M.J., J.F. and I.G. wrote the manuscript
648 with significant input from C.L. and contributions from all of the authors.

649

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Figure 1.

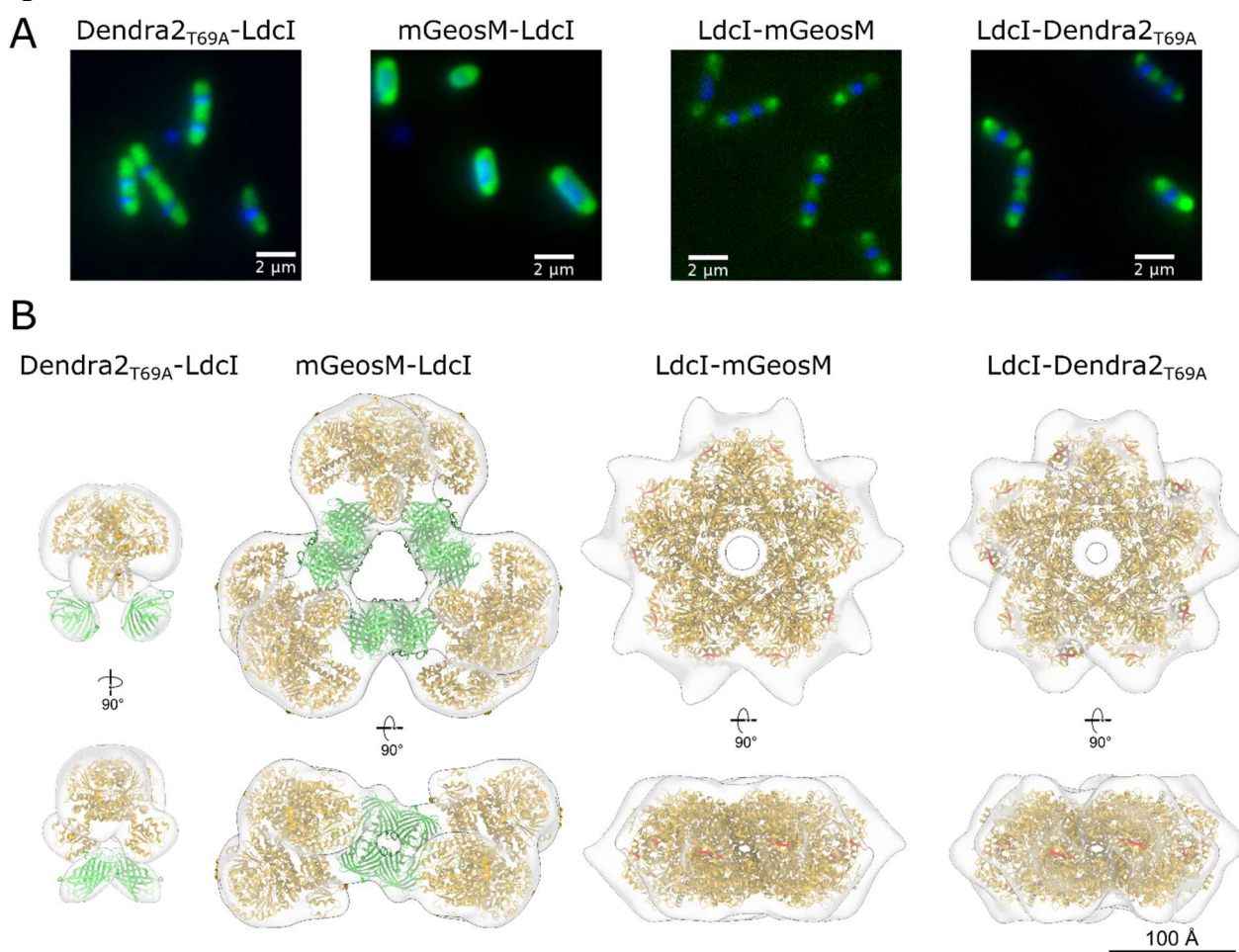


Figure 1. Fluorescent protein fusions affect LdcI structure without significantly altering cellular localisation of overexpressed constructs. A) Wide-field fluorescence microscopy images of *E. coli* cells overexpressing the fluorescent fusion proteins in A). Green fluorescence = FP-LdcI, blue fluorescence = DAPI-stained DNA. **B)** ns-EM 3D maps of fluorescently-tagged LdcI, with fitted models. Left – Dendra2_{T69A}-LdcI forms dimers, with fluorescent barrels located next to the N-terminus of LdcI as expected. Second from left – mGeosM-LdcI forms large non-native oligomers, composed of three LdcI tetramers bridged by tetramers of mGeosM. Second from right and right – both C-terminal fluorescent fusions LdcI-mGeosM and LdcI-Dendra2_{T69A} form decamers, with protrusions at the C-terminus (coloured in red) attributed to flexibly-linked fluorescent proteins. Fitted PBD models are as follows – LdcI: 3N75; Dendra2: 2VZX; mGeosM: 3S05 (mEos2 crystal structure).

Figure 2.

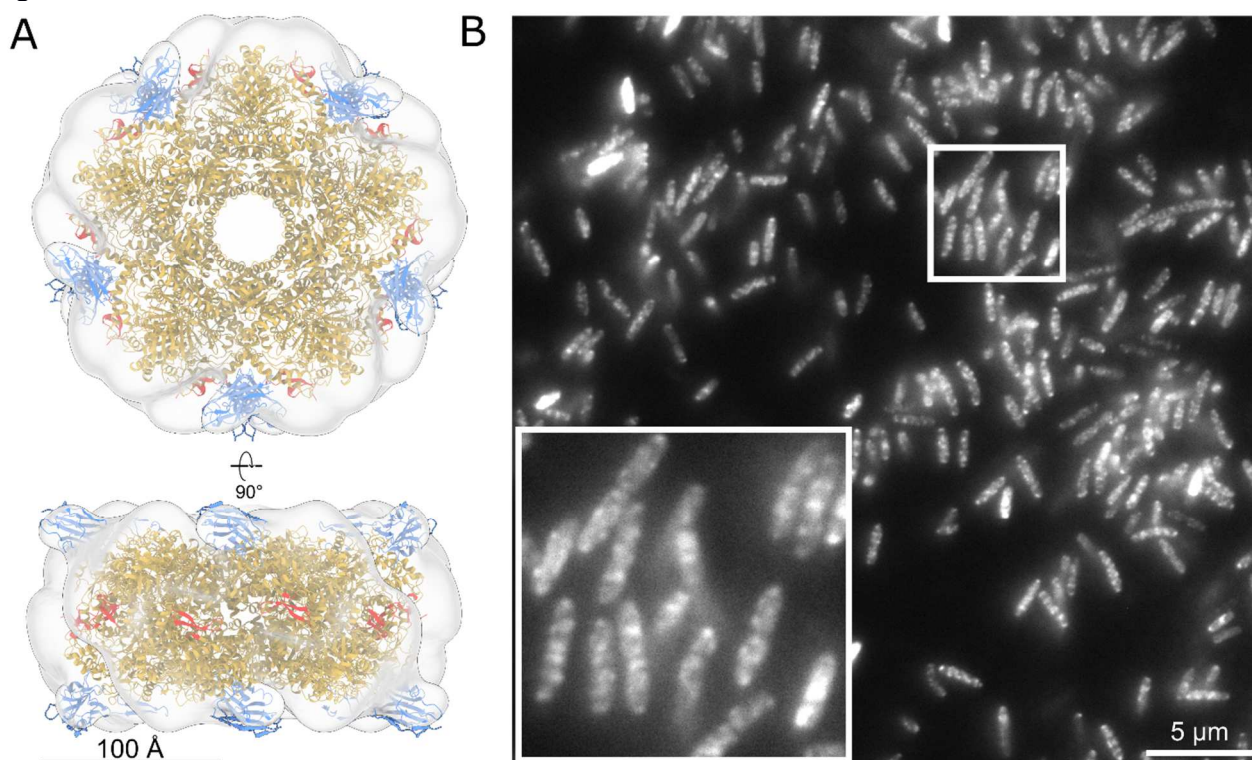


Figure 2. Anti-Ldcl-Nb is a useful tool to probe cellular localisation of endogenous Ldcl under acid stress conditions. **A)** Ns-EM 3D reconstruction of the Ldcl decorated by anti-Ldcl-Nb. An Ldcl decamer (gold, PDB ID: 3N75) and 10 nanobodies (blue, PDB ID: 1MEL, anti-lysozyme nanobody) are fitted in the density, with the nanobody binding at the top and bottom of the decameric ring. The C-terminal RavA binding site is indicated in red, and is in a spatially distinct location from the bound nanobodies. Scale bar = 100Å. **B)** Wide-field fluorescence microscopy image of wild-type *E. coli* MG1655 cells grown at pH 4.6 for 90 minutes stained with anti-Ldcl-Nb labelled with AF488. Inset – zoom of the image showing punctuate fluorescence patterns. Scale bar = 5 μm.

Figure 3.

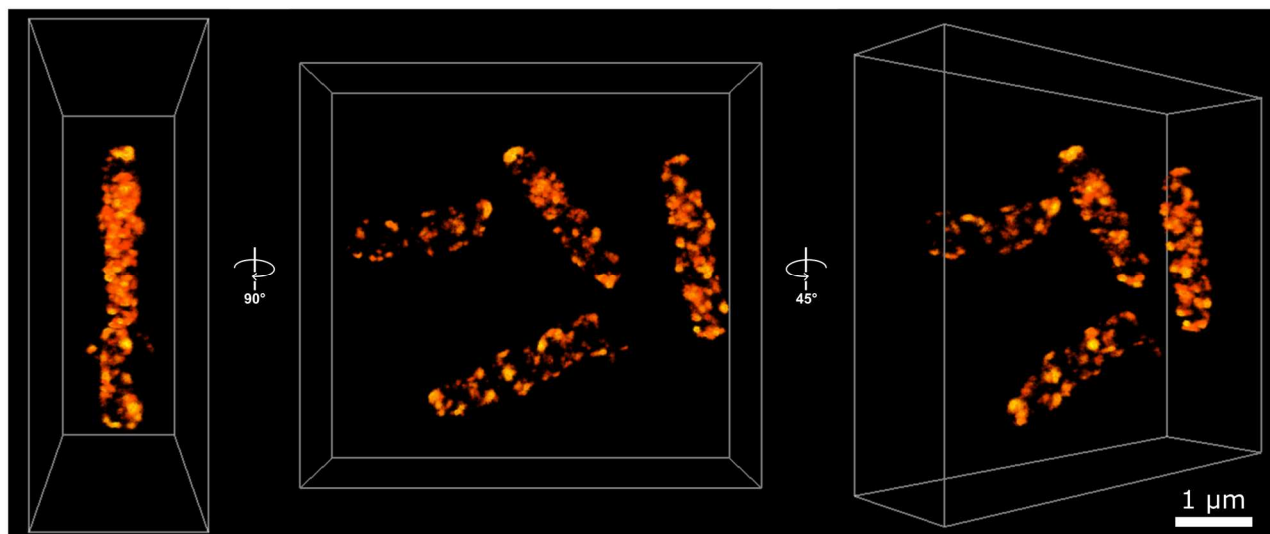


Figure 3. 3D STORM imaging of *E. coli* cells stained with anti-Ldcl-Nb reveal a patchy pseudo-helically arranged distribution of endogenous Ldcl upon acid stress. 3D-STORM imaging of wild-type Ldcl with AF647-conjugated anti-Ldcl-Nb. Points are coloured according to localisation density, with brighter points corresponding to higher localisation density. The centre panel shows four cells in the field of view, looking down the z-axis. Left and right panels show side and tilted views respectively. Scale bar = 1 nm.

Figure 4.

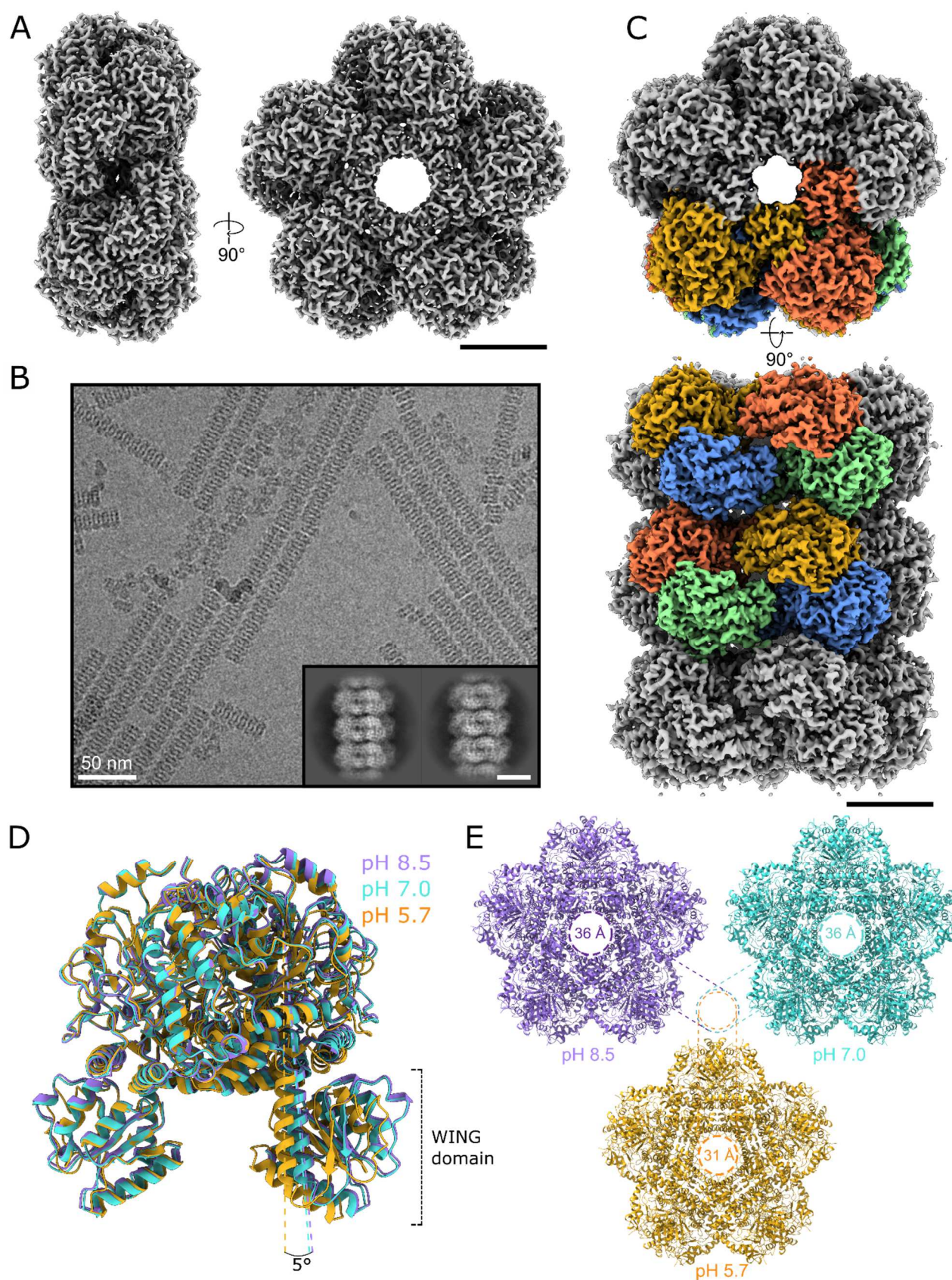


Figure 4. Cryo-EM analysis of the Ldcl decamer at neutral pH and Ldcl stacks formed in acid stress conditions. A) Cryo-EM reconstruction of the Ldcl decamer at pH 7.0 from side (left) and top (right) views. Scale bar = 50 Å. **B)** Cryo-EM micrograph of Ldcl stacks, scale bar = 100 nm. Inset – 2D class 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 Ldcl stack. Four dimers

are coloured either blue/gold or green/coral, corresponding to the colouring of the atomic model presented in Figure 5. Scale bar = 50 Å. **D)** Overlay of Ldcl 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 Ldcl at pH 5.7 and pH 7.0/8.5. **E)** Comparison of the central decamer pore diameter between Ldcl at pH 8.5, 7.0 and 5.7, showing a 5 Å decrease in the pore size upon stack formation at low pH.

Figure 5.

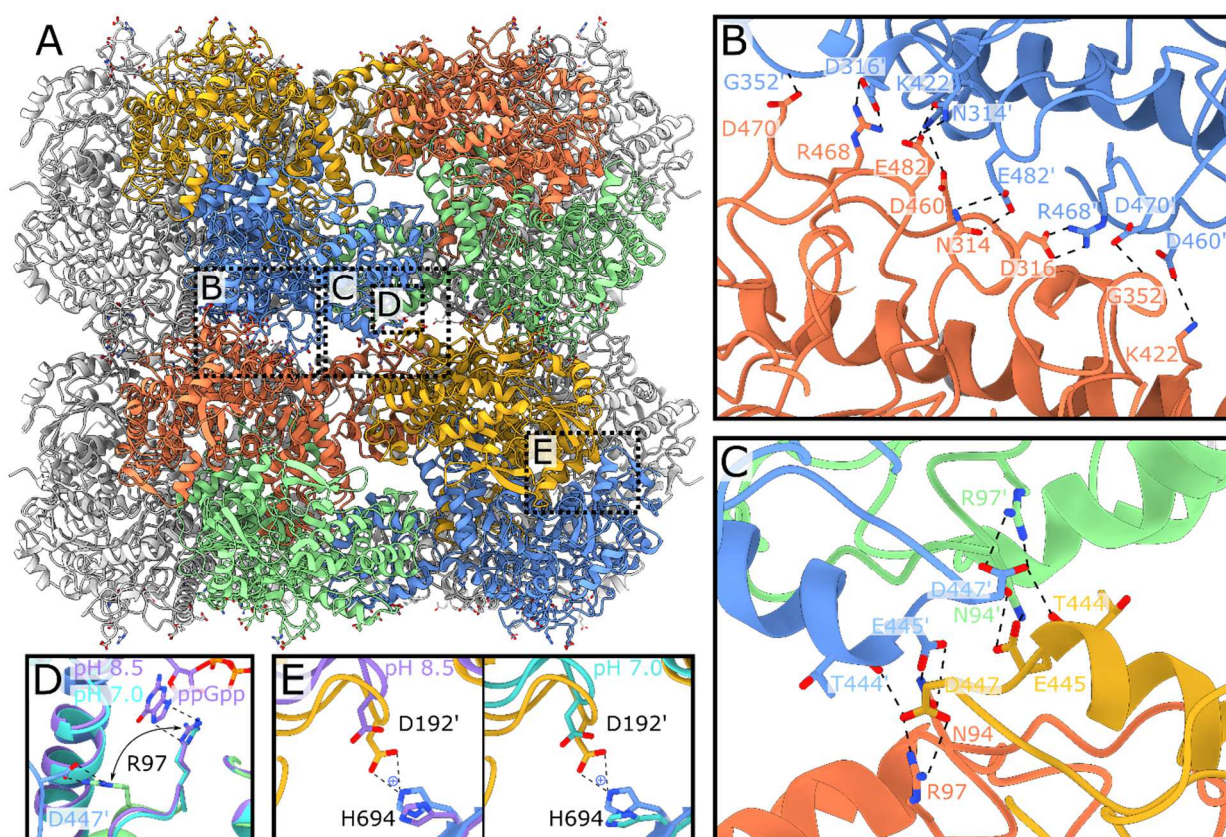


Figure 5. Structural insights into molecular determinants of the Ldcl polymerisation under acid stress conditions. **A)** Atomic model of a two-decamer Ldcl stack at pH 5.7. Dimers are coloured as shown in the cryo-EM map in Figure 4. Dotted boxes on the two-decamer stack indicate the locations of the zooms shown in panels B-E. **B)** Close-up of the first decamer-decamer interface, which includes the key stack-forming residue R468. **C)** Close-up of the second decamer-decamer interface. **D)** Overlay of the Ldcl decamer structures at pH 8.5 and 7.0 with the Ldcl stack structure at pH 5.7, focussed on R97. R97 in the Ldcl 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 towards the ppGpp binding site. **E)** Comparison between the H694-D192' distance in the Ldcl stack at pH 5.7 (coloured gold and blue), the Ldcl decamer at pH 8.5 (left, coloured purple) and pH 7.0 (right, coloured cyan). Key residues are labelled for all panels, and interactions are shown with dotted lines.

Figure 6.

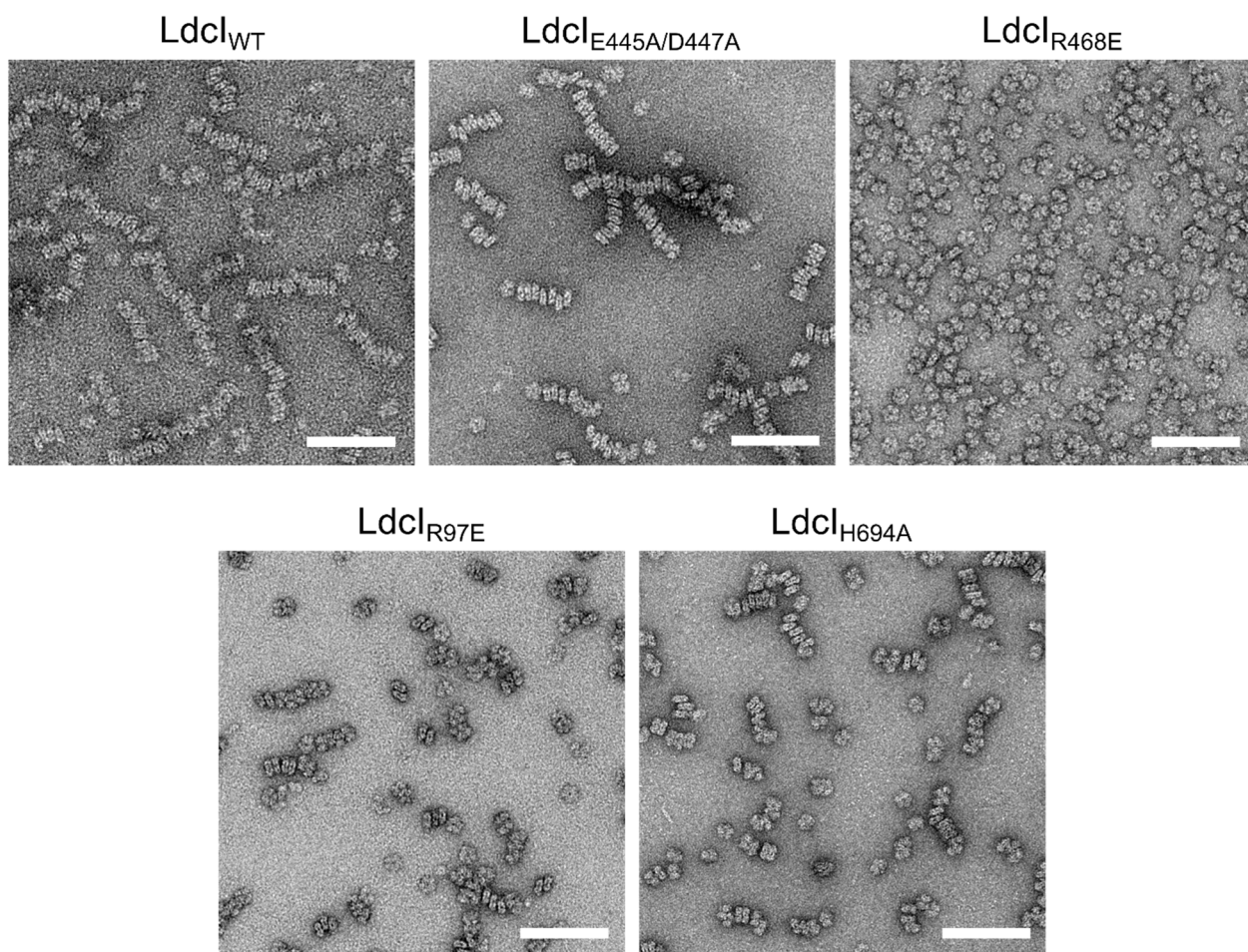


Figure 6. Mutational analysis of the predicted molecular determinants of the Ldcl polymerisation under acid stress conditions Cropped negative stain EM micrographs of wild-type and mutant Ldcl at pH 5.7, scale bar = 100 nm. Ldcl_{WT} polymerises at pH 5.7, as does the double mutant Ldcl_{E445A/D447A}. In contrast, the single mutation R468E abolishes stack formation completely. Both Ldcl_{R97E} and Ldcl_{H694A} are able to polymerise, however the stacks tend to be shorter than for Ldcl_{WT}. Micrographs for Ldcl_{R97E/R468E}, Ldcl_{E445A/D447A/R468E}, and Ldcl_{H694N} were also collected but are not shown here. Ldcl_{E445A/D447A/R468E} and Ldcl_{R97E/R468E} behaved like Ldcl_{R468E} and remained entirely decameric at low pH, while Ldcl_{H694N} displayed similar behaviour to Ldcl_{H694A}.