

1 *Agrobacterium tumefaciens* divisome proteins regulate the transition from polar growth to cell
2 division

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11 **Abstract**

12 The mechanisms that restrict peptidoglycan biosynthesis to the pole during elongation and re-
13 direct peptidoglycan biosynthesis to mid-cell during cell division in polar-growing
14 Alphaproteobacteria are largely unknown. Here, we demonstrate that although two of the three
15 FtsZ homologs localize to mid-cell, exhibit GTPase activity and form co-polymers, only one,
16 FtsZ_{AT}, is required for cell division. We find that FtsZ_{AT} is required not only for constriction and
17 cell separation, but also for the termination of polar growth and regulation of peptidoglycan
18 synthesis at mid-cell. Depletion of FtsZ in *A. tumefaciens* causes a striking phenotype: cells are
19 extensively branched and accumulate growth active poles through tip splitting events. When cell
20 division is blocked at a later stage, polar growth is terminated and ectopic growth poles emerge
21 from mid-cell. Overall, this work suggests that *A. tumefaciens* FtsZ makes distinct contributions
22 to the regulation of polar growth and cell division.

23

24 **Introduction**

25 The spatial and temporal regulation of cell division is a vital process across bacterial species with
26 implications in the development of antimicrobial therapies [1]. The cell division process must
27 coordinate membrane invagination(s), peptidoglycan (PG) biosynthesis and remodeling, and the
28 physical separation of the two daughter cells, all while maintaining cellular integrity.
29 Furthermore, cell division must be precisely regulated to be orchestrated with other key cell
30 cycle processes including cell elongation, DNA replication, and chromosome segregation to
31 ensure that each daughter cell is of sufficient size and contains a complete genome [2, 3].

32

33 To initiate bacterial cell division, the tubulin-like GTPase, FtsZ, polymerizes and forms a
34 discontinuous ring-like structure at the future site of cell division [4-10]. The presence of FtsZ at
35 mid-cell leads to the recruitment of many proteins that function in cell division, collectively
36 called the divisome [11-14]. The divisome includes cell wall biosynthesis proteins, such as the
37 penicillin-binding protein, PBP3, and FtsW, which contribute to PG biosynthesis and remodeling
38 necessary to form new poles in daughter cells [11]. Once the divisome is fully assembled, FtsZ
39 filaments treadmill along the circumference of the mid-cell, driving the Z-ring constriction [9,
40 10]. The movement of FtsZ filaments is correlated with the movement of enzymes that function
41 in septal PG biogenesis. These findings are consistent with the notion that FtsZ not only recruits
42 enzymes that function in PG biogenesis to mid-cell but also regulates their activities to promote
43 proper cell wall biogenesis [15-17].

44

45 In most rod-shaped model organisms used to study cell division, a block in cell division leads to
46 the production of long, smooth filamentous cells. This phenotype suggests that assembly or
47 activation of some divisome components is necessary not only to enable the cells to divide but
48 also to stop cellular elongation. Indeed, in *Escherichia coli*, FtsZ (along with the Z-ring
49 stabilizing proteins FtsA, ZipA, and ZapA) has been proposed to have an early function in the
50 switch from lateral PG biogenesis to mid-cell PG biosynthesis [18]. Following maturation of the
51 divisome by recruitment of additional PG remodeling enzymes and cell division proteins, PG
52 biosynthesis is coordinated with membrane invagination, enabling cells to constrict and separate
53 [19].

54

55 Conversely to e.g. *E. coli*, polar growing rods in the alphaproteobacterial clade Rhizobiales
56 exhibit branched morphologies when cell division is blocked [20-27]. Examination of the cell
57 morphologies resulting from the block in cell division suggests that different types of branched
58 morphologies arise [28]. Drug treatments that block DNA replication cause an early block in cell
59 division, resulting in a “Y” morphology in which the branches are formed from existing growth
60 poles [25, 26]. In contrast, antibiotics that target PBP3 cause mid-cell bulges and branches with
61 some cells adopting a “T” or “+” morphologies [25, 27]. These observations suggest that polar-
62 like PG synthesis is redirected to mid-cell when cell division is blocked at a later stage. The
63 manifestation of two distinct phenotypes during early and late blocks in cell division suggests
64 that divisome assembly and activation may contribute to termination of polar growth, onset of
65 mid-cell PG biosynthesis, cell constriction, and ultimately cell separation.

66

67 In *Agrobacterium tumefaciens*, homologs of FtsZ and FtsA fused to fluorescent proteins localize
68 at the growth pole during elongation and at mid-cell during division [27, 29, 30]. FtsZ was found
69 to arrive at mid-cell considerably earlier than FtsA [30], indicating that FtsZ may be able to
70 initiate Z-ring formation prior to FtsA recruitment to the divisome. This observation is consistent
71 with the described order of divisome assembly in *Caulobacter crescentus* [31] and suggests that
72 a distinctive time-dependent role of these proteins in cell division.

73

74 Here, we take advantage of the ability to deplete essential proteins in *A. tumefaciens* [32] to
75 explore the function of cell division proteins FtsZ, FtsA, and FtsW in a polar growing
76 alphaproteobacterium. Although the genome of *A. tumefaciens* encodes three FtsZ homologs, we

77 find that only one, henceforth referred to as FtsZ_{AT}, is essential for cell survival. FtsZ_{AT} is
78 required to recruit division proteins to mid-cell and likely regulates the activity of PG
79 biosynthesis enzymes at mid-cell. In the absence of FtsZ_{AT}, cells not only fail to divide but are
80 also unable to terminate polar growth. Depletion of either FtsA or FtsW also causes a block in
81 cell division, but unlike FtsZ_{AT} depletion, growth at the poles is halted and instead, polar-like PG
82 synthesis is redirected to mid-cell. These observations suggest that only FtsZ is required to
83 terminate polar growth and initiate cell division-specific PG biosynthesis at mid-cell, whereas
84 FtsZ, FtsA, and FtsW are exclusively required for cell division. Together these findings suggest
85 that *A. tumefaciens* uses sequential regulation of cell division, a theme that is broadly conserved
86 in bacteria.

87

88 **Results and Discussion**

89 **FtsZ_{AT} is required for cell division and termination of polar growth.** *Agrobacterium*
90 *tumefaciens* contains three homologs of *Escherichia coli*'s FtsZ, Atu_2086, Atu_4673, and
91 Atu_4215 (Figure 1A) [27]. *E. coli* FtsZ is comprised of three regions: the conserved N-terminal
92 tubulin-like GTPase domain, a C-terminal linker (CTL), and a conserved C-terminal peptide
93 (CTP), which anchors FtsZ to the membrane via interactions with FtsA [33]. Atu_2086 contains
94 each of these domains out of which the GTPase domain and CTP share 52% and 67% identity to
95 their respective domain in *E. coli* FtsZ, whereas the CTL is extended in length [27]. The gene
96 encoding Atu_2086 is found in a putative operon with genes encoding DdlB, FtsQ, FtsA [34, 35]
97 and is predicted to be essential for cell survival based on saturating transposon mutagenesis [36].
98 Atu_2086 localizes to mid-cell in wildtype (WT) pre-divisional cells (Figure 1B) [27, 29];
99 consistent with a role in cell division. Atu_4673 (called FtsZ₁; consistent with the genome

100 annotation) contains a complete GTPase domain with 49% identity to tubulin domain of *E. coli*
101 FtsZ but lacks both the CTL and CTP [27]. Although Atu_4673 is not predicted to be required
102 for cell survival based on saturating transposon mutagenesis [36], it localizes to mid-cell in pre-
103 divisional cells, suggesting a possible role in cell division (Figure 1B). Atu_4215 (termed FtsZ₃
104 in this work) contains a partial GTPase domain with 48% identity to the N-terminal portion of
105 the *E. coli* FtsZ tubulin domain and lacks both the CTL and CTP [27]. FtsZ₃ is not essential for
106 survival of *A. tumefaciens* based on saturating transposon mutagenesis [36] and exhibits a diffuse
107 localization pattern (Figure 1B). Together, these data suggest that Atu_2086 is the canonical
108 FtsZ protein required for cell division, and this protein will be referred to as FtsZ_{AT} throughout
109 this work (although it is annotated as FtsZ₂ in the *A. tumefaciens* C58 genome [34, 35]).

110

111 To characterize the function of each FtsZ homolog, we constructed deletions of *ftsZ₁* and *ftsZ₃*
112 and a depletion strain of *ftsZ_{AT}*. Since we were unable to construct a deletion of *ftsZ_{AT}*, we used a
113 depletion strategy in which *ftsZ_{AT}* is present as a single copy under the control of an isopropyl β-
114 D-1-thiogalactopyranoside (IPTG) inducible promoter at a neutral site in the chromosome [21,
115 32]. Using western blot analysis, we have confirmed the depletion of FtsZ_{AT} in the absence of
116 IPTG (Figure 1- Figure Supplement 1A).

117

118 Deletion of *ftsZ₁* or *ftsZ₃* does not impact cell viability (Figure 1C), cell morphology (Figure 1D;
119 Table 1; Figure 1-Figure Supplement 1B), microcolony formation (Figure 1D), constriction rate
120 or position (Table 1) when compared to WT cells. Similarly, when FtsZ_{AT} is expressed in the
121 depletion strain (labeled in Figures as +FtsZ_{AT}) the cells remain viable (Figure 1C), are similar in

122 size to WT cells (Table 1), properly position constrictions (Table 1), and form microcolonies
 123 (Figure 1D). In contrast, depletion of FtsZ_{AT} (labeled in Figures as -FtsZ_{AT}) causes a marked
 124 decrease in cell viability (Figure 1C) and triggers the formation of large cells with complex
 125 branched morphologies (Table 1; Figure 1D). To quantify changes in morphology during
 126 depletion of FtsZ_{AT}, the cell area of at least 100 cells was calculated based on phase contrast
 127 images of cells acquired immediately after removal of the inducer (-FtsZ_{AT} 0 h), 8 h after
 128 removal of the inducer (-FtsZ_{AT} 8 h), and 14 h after removal of the inducer (-FtsZ_{AT} 14 h) (Table
 129 1, Figure 1- Figure Supplement 1C). Initially, the FtsZ_{AT} depleted cells are similar to WT in cell
 130 size, but after 8 h of FtsZ_{AT} depletion the cell area has nearly doubled (Table 1, Figure 1- Figure
 131 Supplement 1C). Within 14 h of FtsZ_{AT} depletion, the average cell area has dramatically
 132 increased (Table 1, Figure 1- Figure Supplement 1C). Together, these results demonstrate that
 133 only the FtsZ_{AT} homolog is required for proper cell growth and division.

		Average Cell Length^a (μm +/- SD^b)	Average Cell Area^a (μm^2 +/- SD)	Average Constriction Rate^c (nm/min +/- SD)	Relative Constriction Position^d +/- SD
	WT	2.31 +/- .50	1.66 +/- .35	6.82 +/- 3.19	.49 +/- .05
	$\Delta ftsZ_1$	2.25 +/- .49	1.52 +/- .33	6.99 +/- 3.58	.46 +/- .05
	$\Delta ftsZ_3$	2.24 +/- .47	1.44 +/- .30	6.77 +/- 2.77	.46 +/- .05
	$\Delta ftsZ_1$ $\Delta ftsZ_3$	2.25 +/- .51	1.47 +/- .34	6.61 +/- 3.75	.46 +/- .04
<i>ftsZ_{AT}</i> depletion	- FtsZ _{AT} 0 h	2.71 +/- .70	1.56 +/- .39	6.38 +/- 2.81	.49 +/- .07
	- FtsZ _{AT} 8 h	ND ^e	2.95 +/- 1.12	ND	ND
	- FtsZ _{AT} 14 h	ND	11.37 +/- 4.69	ND	ND

134 **Table 1. Quantitation of cell size and constriction of *ftsZ* mutants**

135 ^aAt least 100 cells were used to quantify the cell length and area for each strain

136 ^bSD – standard deviation.

137 ^cAt least 30 cells were used to quantify the constriction rates for each strain

138 ^dRelative constriction position for at least 40 cells is shown. A value of 0 corresponds to the new
139 pole, 0.5 corresponds to mid-cell, and a value of 1 corresponds to the old pole.

140 ^eND – not determined.

141

142 **Deletion of *ftsZ₁* and *ftsZ₃* does not change the FtsZ_{AT} depletion phenotype.** Since the *ftsZ₁*
143 and *ftsZ₃* single deletions do not have an obvious impact on cell morphology, growth, or division,
144 we constructed double and triple mutants to determine if there is an increasing effect when
145 removing multiple *ftsZ* homologs. Double deletion of *ftsZ₁* and *ftsZ₃* does not cause a decrease in
146 cell viability (Figure 2A, top panel), cell morphology (Table 1), or microcolony formation
147 (Figure 2A, bottom panel). Furthermore, Δ *ftsZ₁* Δ *ftsZ₃* cells properly place constrictions and have
148 an average constriction rate similar to WT (Table 1). Next, we introduced the Δ *ftsZ₁*, Δ *ftsZ₃*, and
149 Δ *ftsZ₁* Δ *ftsZ₃* mutations into the *ftsZ_{AT}* depletion strain to determine if loss of multiple *ftsZ*
150 homologs further aggravated the *ftsZ_{AT}* depletion phenotypes. The combination of the *ftsZ_{AT}*
151 depletion strain with Δ *ftsZ₁*, Δ *ftsZ₃*, or Δ *ftsZ₁* Δ *ftsZ₃* mutations did not result in a further decrease
152 in cell viability (Figure 2B, top panel) or a worsening of cell morphology (Figure 2B, bottom
153 panel) when compared to FtsZ_{AT} depletion alone. Together, these results suggest that the FtsZ₁
154 and FtsZ₃ homologs do not have a major impact on cell division under the conditions tested.

155

156 *ftsZ* gene duplications have occurred independently in several alphaproteobacterial lineages and
157 in chloroplasts and some mitochondria [37]. In most of the cases that have been studied, one

158 FtsZ homolog plays a canonical role in cell or organelle division while the other plays a
159 regulatory or specialized role. However, little is known about the roles of multiple ftsZs in
160 certain alphaproteobacteria species. In both *Rhizobium meliloti* and *Magnetospirillum*
161 *gryphiswaldense*, one of the FtsZs (containing a CTL and CTP similar to FtsZ_{AT}) is essential and
162 the other (truncated after the GTPase domain similar to FtsZ₁) is dispensable [38, 39]. In the case
163 of *M. gryphiswaldense*, the truncated *ftsZ* is dispensable for division but important for
164 biomineralization in this magnetotactic species under certain growth conditions. Similarly, it is
165 possible that FtsZ₁ or FtsZ₃ may have important contributions to cell growth or division of *A.*
166 *tumefaciens* in different environments as e.g. in its plant-associated life-style.

167

168 **FtsZ₁ requires FtsZ_{AT} to localize to mid-cell and to polymerize *in vitro*.** Since FtsZ₁ localizes
169 to mid-cell (Figure 1B), we hypothesized that FtsZ₁ may be a nonessential divisome component.
170 To test this, we examined the localization of FtsZ₁-sfGFP in both WT and the *ftsZ_{AT}* depletion
171 strain (Figure 3). In WT and FtsZ_{AT} induced cells, FtsZ₁-sfGFP does not localize in newborn
172 cells but forms FtsZ-like rings at the future site of division in pre-divisional cells (Figure 3A, top
173 and middle panel). This Z-like ring constricts to form a single focus in dividing cells. These
174 observations suggest that FtsZ₁ may be a divisome component despite the absence of a cell
175 division phenotype in the Δ *ftsZ₁* strain. To explore the possibility of interactions arising due to
176 the loss of FtsZ₁ and FtsZ_{AT}, we next visualized FtsZ₁-sfGFP localization during the depletion of
177 FtsZ_{AT} (Figure 3A, bottom panel). We pre-depleted FtsZ_{AT} for 4 h in liquid to avoid cell
178 crowding caused by division events prior to sufficient FtsZ_{AT} depletion. Early during the
179 depletion of FtsZ_{AT}, FtsZ₁-sfGFP localizes in a FtsZ-like ring near mid-cell. However, as the

180 FtsZ_{AT} depletion continues, FtsZ₁-sfGFP rings and foci progressively fade away, demonstrating
181 that localization of FtsZ₁-sfGFP to mid-cell requires the presence of FtsZ_{AT}.

182

183 Since FtsZ₁ is recruited to mid-cell by FtsZ_{AT}, we hypothesized that FtsZ_{AT} and FtsZ₁ may form
184 co-polymers. To first test the ability of FtsZ_{AT} and FtsZ₁ to independently form polymers, each
185 protein was purified and subjected to polymerization studies. Right angle light scattering assays
186 of wildtype FtsZ_{AT} revealed that this protein exhibits a GTP-dependent increase in light
187 scattering at concentrations above 2 μM , consistent with its polymerization (Figure 3B, blue
188 lines). Negative stain transmission electron microscopy (TEM) confirmed that FtsZ_{AT} forms
189 gently curved protofilaments in the presence of GTP (Figure 3C, left panel) and it rapidly
190 releases inorganic phosphate suggesting that GTP is hydrolyzed (Figure 3D, blue lines; 4.7 ± 0.2
191 $\text{GTP min}^{-1} \text{FtsZ}^{-1}$ at 8 μM FtsZ_{AT}, n=3). Surprisingly, we did not observe polymerization of
192 wildtype FtsZ₁, even at high protein concentrations either in light scattering (Figure 3B, red line),
193 TEM (Figure 3C, center panel), or GTP hydrolysis assays (Figure 3D, red line).

194

195 In light of the dependence of FtsZ₁ on FtsZ_{AT} for mid-cell localization, we next sought to
196 determine if FtsZ_{AT} and FtsZ₁ can form co-polymers. To conduct these experiments, FtsZ₁-
197 L71W and FtsZ_{AT}-L72W were purified to enable monitoring of protein polymerization using
198 tryptophan fluorescence. The leucine to tryptophan mutation introduces a tryptophan on the
199 surface of FtsZ that increases in fluorescence when it is buried in the subunit interface upon
200 polymerization [40]. While wildtype FtsZ_{AT} (with no tryptophan) does not change in
201 fluorescence on addition of GTP (Figure 3E, solid blue line), FtsZ_{AT}-L72W fluorescence

202 increases rapidly after GTP addition reflecting polymerization (Figure 3E, dashed blue line).
203 When wildtype FtsZ_{AT} is added to FtsZ_{AT}-L72W, bringing the total FtsZ concentration to 8 μ M,
204 fluorescence again increases, but then drops back to baseline upon complete consumption of
205 GTP by this high concentration of FtsZ (Figure 3E, dotted blue line). Conversely, on its own or
206 combined with wildtype FtsZ_I, FtsZ_I-L71W maintains a constant tryptophan fluorescence level
207 before and after addition of GTP, consistent with our conclusion that it does not polymerize on
208 its own (Figure 3E, red lines). Remarkably, tryptophan fluorescence increases when FtsZ_I-L71W
209 and FtsZ_{AT} are mixed, indicating that the FtsZ_I-L71W is incorporated into polymers in the
210 presence of FtsZ_{AT} (Figure 3C, purple dashed line). When FtsZ_{AT}-L72W is mixed with FtsZ_I,
211 fluorescence increases above the level observed for FtsZ_{AT}-L72W alone and drops to baseline
212 faster than FtsZ_{AT}-L72W on its own, again indicating co-polymerization. Finally, equimolar
213 concentrations of FtsZ_{AT} alone or mixtures of FtsZ_{AT} and FtsZ_I exhibit similar rates of GTP
214 hydrolysis (Figure 3D) and form qualitatively similar polymers by TEM (Figure 3C, right panel).
215 Together, these observations indicate that FtsZ_I cannot polymerize independently, but that
216 FtsZ_{AT} and FtsZ_I form co-polymers with similar structure and GTP hydrolysis rates as FtsZ_{AT}
217 polymers.

218

219 Though multiple FtsZs are present in a number of bacterial and chloroplast lineages, their co-
220 assembly properties have only begun to be characterized. In contrast to our observations, each of
221 the FtsZs of *M. gryphiswaldense* was able to independently polymerize *in vitro*, but they also
222 appeared to directly interact, perhaps reflecting an ability to co-polymerize [39]. Chloroplast
223 FtsZs from *Arabidopsis thaliana* are also able to co-polymerize and, at least under some
224 conditions, to independently polymerize [41]. Conversely, one of the FtsZs from tobacco

225 chloroplasts cannot polymerize on its own but promotes polymerization of its partner homolog
226 [42]. Finally, the FtsZ pair from the chloroplasts of representative green and red algae co-
227 polymerize into polymers with altered assembly dynamics from either homopolymer [43]. It is
228 likely that in each of these cases, the assembly or co-assembly properties of the duplicated FtsZs
229 have evolved to suit a niche regulatory function. We hypothesize the FtsZ₁ from *A. tumefaciens*
230 has low affinity for itself, but higher affinity for FtsZ_{AT}, limiting its homopolymerization but
231 allowing for co-polymerization both *in vitro* and in cells. Since FtsZ₁ cannot polymerize
232 independently, FtsZ_{AT} must first polymerize at mid-cell after which FtsZ₁ can be recruited by co-
233 polymerization. The biological relevance of these biochemical and cell biological properties
234 awaits further study.

235

236 **FtsZ_{AT} depletion results in tip splitting events.** Once we identified FtsZ_{AT} as the primary
237 homolog involved in cell division we next analyzed the growth phenotype during FtsZ_{AT}
238 depletion more carefully. Compared to FtsZ_{AT} induced cells (Figure 4A, top), observation of
239 cells during FtsZ_{AT} depletion by time-lapse microscope reveals remarkable changes in cell
240 morphology (Figure 4A, bottom; Movie 1). Early during the depletion of FtsZ, an ectopic pole
241 forms near mid-cell. We hypothesize that this occurs due to the ability of the remaining FtsZ to
242 identify the mid-cell and recruit PG biosynthesis machinery to that site. Both the original growth
243 pole and the ectopic pole are growth-active, resulting in the presence of multiple growth poles.
244 These growth poles are unable to terminate cell elongation and ultimately most growth active
245 poles are split, leading to the accumulation of many growth active poles (Figure 4A, bottom;
246 Movie 1) and the rapid increase in cell area until the cell lyses.

247

248 The branched morphology observed during FtsZ_{AT} depletion is in stark contrast to FtsZ depletion
249 observed in other organisms. In species like *E. coli* and *B. subtilis*, which utilize laterally
250 localized peptidoglycan biosynthesis during elongation, depletion of FtsZ results in long, smooth
251 filamentous cells. We hypothesize that the branching morphology of the *A. tumefaciens* FtsZ_{AT}
252 depletion strain can be attributed to polar elongation. During the block in cell division, the
253 growth pole continues to grow and presumably recruits additional peptidoglycan biosynthesis
254 proteins. This could lead to an over-accumulation of elongasome proteins causing the pole to
255 split into two poles. A similar branching pattern has been characterized during typical growth of
256 *Streptomyces coelicolor* [44]. In this polar growing bacterium, the established elongasome splits,
257 leaving a small portion of the elongasome behind as growth continues. With time, the subpolar
258 elongasome accumulates in size and eventually forms a new growth pole. Although the polar
259 growth molecular mechanisms are not conserved between *A. tumefaciens* and *S. coelicolor*, the
260 fundamental principle of tip splitting as a consequence of polar growth appears to be shared.

261

262 **PopZ-YFP accumulates at growth poles in the absence of FtsZ_{AT}.** In WT *A. tumefaciens*,
263 deletion of *popZ* has been shown to cause ectopic poles and cells devoid of DNA, demonstrating
264 a role in coordinating cell division with chromosome segregation [45, 46]. We hypothesize that
265 PopZ-dependent coordination of cell division likely involves FtsZ. In WT, PopZ-YFP localizes
266 to the growing pole during elongation and is recruited to mid-cell just prior to cell separation
267 (Figure 4B, top panel) [45-47]. When FtsZ_{AT} is expressed in the *ftsZ_{AT}* depletion strain, PopZ-
268 YFP has a similar localization pattern as in WT cells (Figure 4B, middle panel). When FtsZ_{AT} is
269 depleted, PopZ-YFP stays at the growth poles and as tip splitting events lead to the production of
270 new growth poles, PopZ-YFP appears to be split and retained at all growth active poles (Figure

271 4B, bottom panel). These observations indicate that FtsZ is required for mobilizing PopZ from
272 the growth pole to mid-cell. Remarkably, both FtsZ and FtsA are mislocalized in the absence of
273 PopZ, leading to the establishment of asymmetric constrictions sites and a broad range of cell
274 lengths [45]. Together, these data suggest that the presence of both PopZ and FtsZ are important
275 for proper positioning and functioning of the divisome.

276

277 In addition to its function in maintaining proper cell division, *A. tumefaciens* PopZ is also
278 required for chromosome segregation and tethers the centromere of at least one chromosome to
279 the growth pole [46]. Thus, we examined the DNA content of cells depleted of FtsZ_{AT}. In both
280 WT cells and in conditions where *ftsZ_{AT}* is induced in the *ftsZ_{AT}* depletion strain, DNA labeled
281 with Sytox orange is diffuse throughout most cells (Figure 4C, top two panels). In late divisional
282 cells, true separation of nucleoids is observed indicating successful completion of chromosome
283 segregation (Figure 4C, marked with an asterisk in the top two panels). In cells depleted of
284 FtsZ_{AT} for both 8 and 14 h, DNA is diffuse throughout the elongated branches (Figure 4C,
285 bottom two panels). The absence of distinct nucleoids may suggest that final stages of
286 chromosome segregation are coordinated with cell separation as has been described for other
287 bacteria including *E. coli* and *C. crescentus* [48].

288

289 To look more carefully at genomic content, we visualized YFP-ParB1, which serves as a proxy
290 to track centromere partitioning in *A. tumefaciens* [46], in WT and *ftsZ_{AT}* depletion cells. In both
291 WT cells and cells expressing FtsZ_{AT} in the *ftsZ_{AT}* depletion strain, a single YFP-ParB1 focus is
292 present at the old pole in new cells generated by a recent cell division event (Figure 4D, top and

293 middle panels). As the cells elongate, a second focus appears and translocates across the
294 longitudinal axis to the growing pole (Figure 4D, top and middle panels). After 4 h of FtsZ_{AT}
295 pre-depletion, YFP-ParB1 foci can be seen at both poles, but when the cell fails to divide, a third
296 focus of YFP-ParB1 appears and translocates along the longitudinal axis of the cell before taking
297 a rapid turn toward a new ectopic pole formed from near mid-cell (Figure 4D, bottom panel).
298 Next, we quantified the number of YFP-ParB1 foci relative to cell area (Figure 4E). In WT and
299 FtsZ_{AT} expressing cells in the *ftsZ_{AT}* depletion strain, small cells have only a single focus of YFP-
300 ParB1. This is followed by a transition period in which elongating cells accumulate a second
301 focus of YFP-ParB1. Finally, the largest, pre-divisional cells have two YFP-ParB1 foci (Figure
302 4E). Cells depleted of FtsZ_{AT} for 8 h accumulate YFP-ParB1 foci as they increase in area (Figure
303 4E). Cells with an area larger than 3 μm^2 all have at least 2 YFP-ParB1 foci, suggesting that
304 chromosome replication is not blocked during FtsZ depletion. Furthermore, in larger cells
305 additional YFP-ParB1 foci accumulate. These data suggest that cell division is not strictly
306 required for the initiation of DNA replication in *A. tumefaciens*, although completion of
307 chromosome segregation may be coordinated with cell division.

308

309 **Loss of septal PG synthesis results in altered total PG composition.** Since polar growth
310 appears to continue in the absence of FtsZ (Figure 4A, bottom panel), we used fluorescent-D-
311 amino acids (FDAAs), to probe sites enriched in peptidoglycan synthesis [49] during depletion
312 of FtsZ_{AT}. In WT cells, FDAAs localize at a single pole in elongating cells and at mid-cell in pre-
313 divisional cells (Figure 5A) [49]. As FtsZ_{AT} is depleted, FDAAs are targeted strictly to the poles,
314 confirming that polar peptidoglycan synthesis is responsible for the observed increase in cell
315 biomass after 8 h and 14 h of depletion (Figure 5A).

316

317 Since cells depleted of FtsZ_{AT} fail to terminate polar growth and do not produce septal
318 peptidoglycan, we hypothesized that the peptidoglycan composition may reveal chemical
319 signatures of peptidoglycan derived from polar growth. Thus, we characterized the
320 peptidoglycan composition of both WT cells and the *ftsZ_{AT}* depletion strain in both the presence
321 and absence of IPTG using ultra-performance liquid chromatography (UPLC) [50]. The major
322 muropeptides found in WT *A. tumefaciens* PG and their quantification are shown in (Figure 5-
323 Figure Supplement 1) and include monomeric (M), dimeric (D), and trimeric (T) muropeptides.
324 The muropeptide composition and abundance is similar between WT cells, WT cells grown in
325 the presence of IPTG, and the *ftsZ_{AT}* depletion strain grown in the presence of IPTG such that
326 FtsZ_{AT} is expressed (Figure 5- Figure Supplement 1). These findings suggest that there are no
327 major changes in PG composition due to IPTG and that the presence of IPTG leads to
328 complementation in the *ftsZ_{AT}* depletion strain. In contrast, when the *ftsZ_{AT}* depletion strain is
329 grown in the absence of IPTG for 14 h, marked changes in muropeptide composition are
330 observed (Figure 5B-D). While the overall abundance of monomeric, dimeric, and trimeric
331 muropeptides are not dramatically impacted (Figure 5C), the abundance of specific muropeptides
332 is modified. When FtsZ_{AT} is depleted, there is a significant increase in monomeric disaccharide
333 tripeptide (M3) and a decrease in the abundance of the monomeric disaccharide tetrapeptide M4
334 (Figure 5B). This observation is consistent with the possibility that the absence of FtsZ_{AT} leads to
335 an increase in LD-carboxypeptidase activity, which would remove the terminal peptide from M4,
336 leading to both a reduction in the levels of M4 and an increase in the abundance in M3.
337 Following FtsZ_{AT} depletion, the overall degree of muropeptide crosslinking decreases (Figure
338 5D). In particular, there is a marked decrease in DD-crosslinkages, which are formed by the DD-

339 transpeptidase activity associated with penicillin-binding proteins (PBPs). The dominant dimeric
340 muropeptide formed in the presence of FtsZ_{AT} is D44, which contains a DD-crosslink; in contrast,
341 the dominant dimer formed in the absence of FtsZ_{AT} is D33, which contains an LD-crosslink
342 (Figure 5B). These data suggest that the activity of LD-transpeptidases is increased and the
343 activity of PBP-mediated DD-transpeptidases is decreased during FtsZ_{AT} depletion. The increased
344 pool of M3 may provide additional acceptor substrate for LD-transpeptidases to increase the
345 production of D33 relative to D44. In addition, increased LD-carboxypeptidase activity could
346 contribute to increase further the levels of D33 using D34 as a substrate.

347

348 The *A. tumefaciens* genome contains 14 LD-transpeptidases, 7 of which are specific to
349 Rhizobiales. The Rhizobiales-specific LD-transpeptidase encoded by *Atu_0845* (referred to here
350 as LDTP₀₈₄₅) has been shown to localize to the growing pole in WT cells and has been
351 hypothesized to contribute to polar growth [30]. This localization pattern was confirmed in both
352 WT and FtsZ_{AT} induced cells (Figure 5E, top and middle panels). We find that LDTP₀₈₄₅
353 localizes at growth poles during depletion of FtsZ_{AT} (Figure 5E, bottom). This observation
354 suggests that this LDTP₀₈₄₅ may contribute to changes in PG composition during FtsZ_{AT}
355 depletion and supports a potential role for LD-transpeptidases in polar growth during elongation.
356 The localization and function of putative periplasmic LD-carboxypeptidases in *A. tumefaciens*
357 remain to be explored. Overall, these findings suggest that LD-carboxypeptidase and LD-
358 transpeptidase activities are increased during FtsZ_{AT} depletion, indicating that these classes of
359 enzymes may contribute to polar growth of *A. tumefaciens*.

360

361 **The C-terminal Conserved Peptide (CTP) of FtsZ_{AT} is required for proper termination of**
362 **polar growth.** To better understand the mechanism by which FtsZ_{AT} terminates polar growth,
363 we constructed truncated proteins to analyze the function of the C-terminal conserved peptide
364 (CTP) and the C-terminal linker (CTL) (Figure 1A). The CTP is a highly conserved domain
365 which binds to proteins such as FtsA, that tether FtsZ to the membrane [37, 51, 52]. The CTL is
366 an intrinsically disordered region of variable length found in FtsZ proteins, which functions in
367 the regulation of PG biosynthesis and protofilament assembly [17, 53-55]. To probe the function
368 of the FtsZ_{AT} CTP and CTL domains, we expressed FtsZ_{AT}ΔCTP and FtsZ_{AT}ΔCTL in both WT
369 and FtsZ_{AT} depletion backgrounds.

370
371 In order to execute these experiments, we constructed a vector with an alternative “inducible”
372 promoter system, which is compatible with the chromosomal IPTG depletion system. We
373 modified pSRKKm [56] by replacing *lacI^q* with the gene encoding the cumate responsive
374 repressor CymR [57, 58] and replacing the *lacO* operator sites with *cuO* operator sites (Figure 6-
375 Figure Supplement 1A). This approach allows the same promoter to drive expression of both
376 chromosomal full-length *ftsZ_{AT}* using IPTG and plasmid-encoded *ftsZ* variants using cumate. For
377 simplicity, henceforth we referred to IPTG induction as mediated by P_{lac} and cumate induction as
378 mediated by P_{cym}. Expression of sfGFP from P_{cym} requires the presence of cumate (Figure 6-
379 Figure Supplement 1B) and is comparable to expression of sfGFP from P_{lac} (Figure 6- Figure
380 Supplement 1C). Although higher concentrations of cumate inhibit growth of WT *A.*
381 *tumefaciens*, 0.01 mM cumate does not impair growth of WT cells (Figure 6- Figure Supplement
382 1D; left) and is sufficient to complement growth of the *ftsZ_{AT}* depletion strain in the absence of
383 IPTG (Figure 6- Figure Supplement 1D; right).

384

385 In the *ftsZ_{AT}* depletion strain, we introduced 4 vectors: an empty vector with P_{cym} (pEmpty), P_{cym}-
386 *ftsZ_{AT}* (pFtsZ_{AT}), P_{cym}-*ftsZ_{AT}ΔCTP* (pFtsZ_{ΔCTP}) or P_{cym}-*ftsZ_{AT}ΔCTL* (pFtsZ_{ΔCTL}). When full-length
387 *ftsZ_{AT}* is expressed from the chromosome, the viability of cells is not impacted by the presence of
388 the P_{cym} vectors (Figure 6A, top left panel). In the absence of induction of *ftsZ* from the
389 chromosome, the presence of the uninduced P_{cym} vectors, including pFtsZ_{AT}, is not sufficient to
390 rescue viability of the FtsZ-depleted cells (Figure 6A, top right panel); however, viability is
391 significantly restored by expression of plasmid-encoded FtsZ_{AT} in the presence of cumate
392 (Figure 6A, bottom left panel). Expression of plasmid-encoded FtsZ_{AT}ΔCTP partially rescues the
393 depletion of FtsZ_{AT} (Figure 6A, bottom left panel). In contrast, expression of plasmid-encoded
394 FtsZ_{AT}ΔCTL does not rescue the depletion of FtsZ_{AT} (Figure 6A, bottom left panel) and when
395 both chromosomal full-length FtsZ_{AT} and FtsZ_{AT}ΔCTL are expressed, viability is impaired,
396 suggesting that FtsZ_{AT}ΔCTL may have a dominant negative phenotype (Figure 6A, bottom right
397 panel).

398

399 Next, we observed cell morphology of the *ftsZ_{AT}* depletion strain carrying each of the four
400 vectors under conditions where the chromosomal FtsZ_{AT} is depleted and the plasmid-encoded
401 FtsZ variants are expressed for 6 or 14 h (Figure 6B). The presence of pEmpty does not impact
402 the FtsZ_{AT} depletion phenotype: branched cells with multiple growth active poles are observed
403 (Figure 6B). Plasmid-encoded FtsZ_{AT} rescues the chromosomal FtsZ_{AT} depletion, resulting in the
404 production of typical rod-shaped cells with PG biosynthesis occurring at a single pole or mid-cell
405 (Figure 6B, middle left). The partial rescue of FtsZ_{AT} depletion in viability by expression of
406 FtsZ_{AT}ΔCTP was matched by a less severe defect in cell morphology (Figure 6B, middle right).

407 Although cells are branched, they are much shorter and have fewer branches than FtsZ_{AT}
408 depletion. FDAA labeling reveals that the expression of FtsZ_{AT}ΔCTP enables mid-cell labeling
409 (Figure 6B, middle left), suggesting that PG is synthesized at mid-cell and that some cells may
410 undergo division. Indeed, time-lapse microscopy of the FtsZ_{AT} depletion strain expressing only
411 FtsZΔCTP reveals that the cells are capable of cell division events (Figure 6C, top panel).
412 Remarkably, the sites of cell constriction and cell division are often asymmetric, giving rise to a
413 cell population with a broad length distribution. Furthermore, polar growth is not terminated
414 efficiently and both polar elongation and tip splitting events are evident. Together, these
415 observations suggest that the FtsZ CTP contributes to proper termination of polar growth and
416 divisome assembly. Expression of plasmid-encoded FtsZ_{AT}ΔCTL in the absence of chromosome-
417 encoded FtsZ_{AT} gives rise to a distinct cell morphology (Figure 6B, far right panel). After 6
418 hours of FtsZ_{AT}ΔCTL expression, some cells contain mid-cell bulges. Remarkably, in these cells,
419 FDAA labeling reveals that PG biosynthesis is occurring in the bulges and not at either pole.
420 After 14 h, most cells have mid-cell swelling and multiple ectopic poles. Time-lapse microscopy
421 reveals that polar growth is terminated and growth appears to be directed to mid-cell (Figure 6C,
422 bottom panel, 320 min). When cell division fails, ectopic growth poles emerge from the mid-cell
423 bulges (Figure 6C, bottom panel 520 min). The ectopic poles elongate, polar growth is
424 terminated, and new ectopic growth poles are formed near the initial bulge site (Figure 6C,
425 bottom panel). These observations suggest that the CTL of FtsZ_{AT} is required for proper cell
426 division but is not required for the termination of polar growth.

427

428 **The CTL of FtsZ_{AT} is required for proper PG composition.** The mid-cell bulges observed
429 during FtsZΔCTL expression are reminiscent of those observed when FtsZ_{CC}ΔCTL is expressed

430 in *C. crescentus* [17]. In *C. crescentus*, the CTL was shown to be required for robust PG
431 biosynthesis [17]. We therefore hypothesized that the altered PG composition observed during
432 depletion of FtsZ_{AT} could be due to absence of the CTL. To test this hypothesis, we introduced
433 plasmids containing no FtsZ (empty vector control, pEmpty), full-length FtsZ_{AT} (pFtsZ_{AT}), or
434 FtsZ_{AT}ΔCTL (pFtsZ_{AT}ΔCTL) into the *ftsZ* depletion strain. Each strain was grown under
435 conditions in which expression of FtsZ_{AT} from the chromosomal copy is depleted and expression
436 of the FtsZ variant (if present) from the plasmid is induced. PG was isolated from these strains
437 following induction/depletion and analyzed. Induction of full-length FtsZ_{AT} from the plasmid
438 yields lower levels of monomeric muropeptides compared to other strains, especially M3, and
439 increased levels of dimeric and trimeric muropeptides, including D44 and T444 (Figure 7A-B).
440 Overall the expression of full-length FtsZ_{AT} leads to an increased level of muropeptides with DD-
441 crosslinks (Figure 7C). These observations indicate that expression of plasmid-encoded full-
442 length FtsZ_{AT} compensates for the loss of FtsZ_{AT} from the chromosome. In contrast, the
443 expression of FtsZ_{AT}ΔCTL did not compensate for the loss of full-length FtsZ as the PG
444 composition is more similar to the PG profile of FtsZ-depleted cells (Figure 7A-C). This
445 observation suggests that the CTL of FtsZ_{AT} likely function in the regulation of proper PG
446 biosynthesis at mid-cell.

447

448 **FtsZ CTL regulates protofilament assembly.** Work in *C. crescentus* has shown that the
449 FtsZ_{CC}CTL directly regulates protofilament structure and dynamics [53]. To determine if the
450 CTL of FtsZ_{AT} similarly regulates its assembly, we purified FtsZ_{AT}ΔCTL and a control
451 FtsZ_{AT}+CTL protein containing the same restriction sites at the junctions with the GTPase
452 domain and CTP as the ΔCTL construct, but with the CTL in place. FtsZ_{AT}+CTL formed mostly

453 single, gently curved protofilaments when visualized by TEM under all conditions tested (Figure
454 7D), similar to those observed for wildtype FtsZ_{AT} (Figure 3C). In contrast, under high salt
455 conditions we observed extended bundles of FtsZ_{AT}ΔCTL (Figure 7D). Furthermore, we saw a
456 decreased rate of GTP hydrolysis by FtsZ_{AT}ΔCTL under conditions that promote bundling
457 (Figure 7D; 3.3 ± 0.2 GTP min⁻¹ FtsZ⁻¹ for FtsZ_{AT}+CTL and 2.1 ± 0.1 GTP min⁻¹ FtsZ⁻¹ for
458 FtsZ_{AT}ΔCTL with 300 mM KCl, n =3). Together, these results suggest an important role for the
459 CTL in limiting lateral interactions between protofilaments and promoting polymer turnover.
460 These results in *A. tumefaciens* are consistent with effects of the CTL on polymer bundling
461 reported in *C. crescentus* [17, 53] and *E. coli* [59]. Moreover, in light of our observations that
462 FtsZ_{AT}ΔCTL does not restore proper PG chemistry to FtsZ_{AT}-depleted cells (Figure 7A,B), these
463 data are in line with the growing body of evidence linking FtsZ dynamics and polymer
464 superstructure to the regulation of PG biosynthesis.

465

466 **FtsA is required for cell division but not termination of polar growth.** FtsA is an actin-like
467 protein that associates with the membrane through an amphipathic helix and binds the FtsZ CTP
468 to anchor FtsZ polymers to the membrane [51, 60]. In *C. crescentus*, recruitment of FtsA to mid-
469 cell occurs well after the establishment of the FtsZ-ring and is dependent on the presence of FtsZ
470 [13, 61]. In *A. tumefaciens*, FtsA-sfGFP is retained at the growth pole prior to appearing at mid-
471 cell just before cell division [27, 30]. Here, we confirm that FtsA-sfGFP is observed as a focus at
472 the growth pole until transitioning to a ring-like structure at mid-cell (Figure 8A, top panel). In
473 fact, at some timepoints, both a polar focus and a mid-cell ring of FtsA are observed. Eventually,
474 the polar focus disappears as the FtsA-sfGFP ring becomes more intense just prior to cell
475 division. During the depletion of FtsZ_{AT}, a focus of FtsA-sfGFP can be found at the growing

476 pole, and at a newly formed ectopic pole near mid-cell (Figure 8A, bottom panel). FtsA-sfGFP
477 remains associated with each growth pole, and as the poles undergo tip splitting events, each
478 focus of FtsA-sfGFP is also split, resulting in the presence of FtsA-sfGFP in each of the 4
479 growth-active poles. These observations suggest that FtsZ_{AT} is required not only for proper mid-
480 cell localization of FtsA to mid-cell prior to cell division but also contributes to release of FtsA-
481 sfGFP from the growth pole.

482 Since FtsA tethers FtsZ to the membrane and enables divisome assembly [37, 51, 52] in *E. coli*,
483 we expected that the depletion of FtsA would phenocopy the depletion of FtsZ. Although a
484 saturating transposon mutagenesis screen indicated that *ftsA* is not essential for *A. tumefaciens*
485 cell survival [36], we were unable to construct a Δ *ftsA* mutant. Thus, we constructed a depletion
486 strain in which expression of *ftsA* is controlled by P_{lac}. Under conditions where FtsA is present in
487 the *ftsA* depletion strain, cells maintain proper rod-shaped morphology, polar growth, and cell
488 division occurs from constrictions formed near mid-cell (Figure 8B-C, top panels). In contrast,
489 when FtsA is depleted, cells exhibit a marked change in morphology (Figure 8B, bottom panel;
490 Movie 2). During the depletion of FtsA, rod-shaped cells initially elongate from a growth pole
491 (Figure 8B, bottom panel, 0 min). Polar growth is terminated and growth is re-initiated from near
492 mid-cell, typically resulting in the formation of two ectopic poles perpendicular to the original
493 longitudinal axis of the cell (Figure 8B, bottom panel, 170 min). Cells depleted of FtsA continue
494 multipolar growth (Figure 8B, bottom panel, 360 min), terminate growth from both poles and
495 reinitiate growth from near mid-cell resulting in the formation of a new pair of ectopic growth
496 poles (Figure 8B, bottom panel, 510 min). This pattern of multipolar growth, polar growth
497 termination, and new branch formation is continued until cells eventually bulge at the mid-cell
498 and lyse. Overall these observations indicate that the phenotypes caused by FtsZ and FtsA

499 depletion are distinct from one another and suggest that only FtsZ is required for proper
500 termination of polar growth.

501

502 To confirm that polar growth occurs and is terminated during FtsA depletion, cells were labeled
503 with FDAAs (Figure 8C, bottom panel). Indeed, FDAAs label the tips of two poles, which are
504 emerging from near mid-cell consistent with the re-initiation of polar growth. To further confirm
505 that polar growth is terminated during FtsA depletion, we observed the localization of PopZ-YFP
506 (Figure 8D, top panel). PopZ marks the growth poles [47] and becomes trapped at growth poles
507 during depletion of FtsZ (Figure 4B). During FtsA depletion, PopZ-YFP is initially present at the
508 growth pole (Figure 8D, top panel, 0 min). Next, PopZ-YFP disappears from the growth poles
509 and reappears near mid-cell (Figure 8D, top panel, 80 min) indicating that polar growth is
510 terminated. Throughout the FtsA depletion, PopZ-YFP continues to disappear from growth poles
511 and reappears at the tips of newly emerging growth poles. Overall, these observations clearly
512 indicate that FtsA is not necessary for termination of polar growth; however, FtsA has an
513 essential function at a later stage of cell division since the cells fail to divide and are prone to
514 lysis.

515

516 The ability of cells to target growth to near mid-cell during FtsA depletion suggests that FtsZ-
517 rings may form, enabling the termination of polar growth. Indeed, FtsZ_{AT}-sfGFP-rings form near
518 mid-cell early during FtsA depletion (Figure 8D, bottom panel). FtsZ_{AT}-sfGFP is briefly retained
519 at new growth poles before reappearing to mark the site where a new growth pole will emerge.
520 These observations are consistent with the finding the FtsA is retained at the growth pole longer

521 than FtsZ [27, 62], and suggest that FtsA arrives at mid-cell after Z-ring assembly and the
522 initiation of FtsZ-dependent cell wall biogenesis. The results observed here in *A. tumefaciens* are
523 consistent with the observation that FtsA arrives to mid-cell after FtsZ and the onset of mid-cell
524 cell wall biogenesis in *C. crescentus* [13, 61]. In both *A. tumefaciens* and *C. crescentus*, the late
525 arrival of FtsA to the divisome suggests that other proteins contribute to proper tethering of FtsZ
526 to the membrane. In *C. crescentus*, the FtsZ-binding protein, FzlC, functions as a membrane
527 anchor early during the establishment of the divisome [31, 63]. A homolog of FzlC is readily
528 found in the *A. tumefaciens* genome (Atu2824) and may contribute to the ability of FtsZ-rings to
529 form in the absence of FtsA.

530

531 **Depletion of the downstream divisome component FtsW phenocopies depletion of FtsA.**

532 Having observed a distinct effect on cell morphology in the absence of *ftsA*, we wondered if the
533 phenotype observed during *ftsA* depletion could be recapitulated in the absence of another late-
534 arriving divisome protein. To test this hypothesis, we constructed a depletion strain of FtsW,
535 which is recruited to mid-cell after FtsA in both *E. coli* and *C. crescentus* divisome assembly
536 models [11, 13]. Depletion of FtsW results in a phenotype which is strikingly similar to the
537 depletion of FtsA (Figure 9). When FtsW is induced normal growth is observed (Figure 9A, top
538 panel). During FtsW depletion, polar growth is terminated, resulting in the establishment of
539 growth-active poles from near mid-cell (Figure 9A, bottom panel; Movie 3). Multiple rounds of
540 termination of polar growth followed by reinitiation of growth from near mid-cell occur until the
541 mid-cell bulges and the cells ultimately lyse (Figure 9A, bottom panel). Labeling of growth
542 active poles with FDAAs (Figure 9B) or by tracking PopZ-YFP localization (Figure 9C, top
543 panel) confirmed that new branches which emerge from mid-cell are formed by polar growth.

544 Finally, we confirmed that FtsZ-rings form during the depletion of FtsW and the presence of an
545 FtsZ_{AT}-sfGFP-ring typically marks the site where an ectopic growth pole will form (Figure 9C,
546 bottom panel). Together, these observations suggest that FtsZ-rings are formed in the absence of
547 FtsW, enabling the initiation of cell wall biogenesis. Given that FtsW drives septal PG
548 biosynthesis, [64] these findings indicate that the cell wall biogenesis that occurs during
549 depletion of FtsA or FtsW may require the elongation machinery, which typically functions in
550 polar growth. Since the elongation machinery for *A. tumefaciens* remains to be identified, it is
551 possible that there is considerable overlap between the machineries that contribute to polar and
552 septal PG biosynthesis.

553

554 **Concluding Remarks**

555 While many questions remain unanswered about the regulation of cell wall biogenesis in *A.*
556 *tumefaciens*, our work sheds light on the transition from polar growth to mid-cell growth. We
557 find that FtsZ_{AT}, FtsA, and FtsW are required for constriction and cell separation, but FtsZ_{AT} is
558 also required to terminate polar growth and initiate mid-cell peptidoglycan synthesis. How might
559 the formation of an FtsZ_{AT}-ring at mid-cell cause the termination of polar growth? We find that
560 PopZ, and LDTP₀₈₄₅ become trapped at the growth poles during FtsZ depletion (Figure 5). It is
561 possible that one or more of these proteins contributes to both polar peptidoglycan biosynthesis
562 and mid-cell peptidoglycan synthesis and that the FtsZ-dependent targeting of these proteins (and
563 likely others) to mid-cell triggers the termination of polar growth. While the mid-cell localization
564 of PopZ is dependent on the presence of FtsZ_{AT} (Figure 4), FtsZ_{AT}-ring stability and placement
565 are impacted by the absence of PopZ [45]. Furthermore, deletion of *popZ* impairs termination of
566 polar growth and results in cell division defects [45, 46, 65]. The apparent co-dependence of
567 FtsZ and PopZ for localization may suggest that these proteins function together during the early

568 stages of cell division, particularly the termination of polar growth and onset of mid-cell PG
569 biosynthesis.

570

571 Overall, our results are consistent with a general model, which is highly conserved in bacteria, in
572 which the establishment of a FtsZ-ring leads to the recruitment of many other cell division
573 proteins to mid-cell [11], though many mechanistic questions remain. How is FtsZ_{AT} targeted to
574 mid-cell? A variety of mechanisms that contribute to the proper placement of FtsZ at mid-cell
575 have been described (for review see [66, 67]). The most well studied mechanisms of FtsZ
576 positioning include negative regulation by the Min system and nucleoid occlusion. While genes
577 encoding components of the Min system are readily identifiable in the *A. tumefaciens* genome,
578 the deletion of *minCDE* has a minimal impact on placement of constriction sites and cell division
579 efficiency [68]. Furthermore, FtsZ_{AT}-GFP rings form over DNA prior to nucleoid separation in
580 *A. tumefaciens*. These observations indicate that additional regulatory mechanisms must
581 contribute to proper division site selection in *A. tumefaciens*. Following the appearance of FtsZ at
582 mid-cell, how is the FtsZ_{AT}-ring stabilized? In *E. coli*, the FtsZ-ring is stabilized by interactions
583 with FtsA and ZipA, which tether FtsZ filaments to the membrane [51, 52, 69, 70]. In *A.*
584 *tumefaciens*, FtsZ_{AT} appears at mid-cell well before FtsA [30] and we observe that FtsZ_{AT} rings
585 form even when FtsA is depleted (Figure 7C, bottom panel). Furthermore, the position of
586 FtsZ_{AT}-GFP rings marks the site of ectopic pole formation. These observations suggest the
587 FtsZ_{AT} is stabilized, at least early during cell division by other proteins. While there are no
588 obvious ZipA homologs encoded in the *A. tumefaciens* genome, a homolog of FzlC, which
589 functions to stabilize FtsZ in *C. crescentus* [31, 63], is encoded in the genome.

590

591 The observation that FtsZ is necessary for the initiation of mid-cell PG biosynthesis suggests that
592 FtsZ is necessary for recruitment of PG biosynthesis enzymes to mid-cell. Septal PG
593 biosynthesis is likely mediated by FtsW, a putative PG glycosyltransferase [71-73], and PBP3
594 (FtsI), a PG DD-transpeptidase [74]. In *A. tumefaciens*, depletion of FtsW does not cause a
595 complete block of PG synthesis at mid-cell (Figure 8). This observation suggests that mid-cell
596 PG biosynthesis is mediated by other cell wall biogenesis enzymes while the activity of FtsW
597 contributes to later stages of cell division, consistent with the inability of cells to form
598 constrictions and separate in the absence of FtsW. These observations may indicate that the
599 initial PG biosynthesis at mid-cell comprises the final stage of cell elongation, consistent with
600 descriptions of FtsZ-dependent mid-cell elongation in *C. crescentus* [16]. The observation that
601 growth-active, ectopic poles emerge from near mid-cell during FtsW depletion (Figure 8B)
602 provides evidence in support of this possibility. Thus, FtsZ-dependent PG biosynthesis may
603 contribute to both elongation and cell division in *A. tumefaciens*. For a polar growing bacterium,
604 it is tempting to speculate that the retention of PG biosynthesis machinery dedicated to
605 elongation at the site of cell division may prime the newly formed poles to become growth active
606 following cell separation.

607

608 **Materials and Methods**

609 **Bacterial strains and culture conditions.** All bacterial strains and plasmids used are listed in
610 Table S4.1. *A. tumefaciens* strains were grown in ATGN minimal medium with .5% glucose [75]
611 at 28°C. *E. coli* strains were grown in Luria-Bertani medium at 37°C. When indicated,
612 kanamycin (KM) was used at 300 µg/ml for *A. tumefaciens*, 50 µg/ml for *E. coli* DH5α, and 25
613 µg/ml for *E. coli* S17-1 λ *pir*. Gentamicin was used when indicated at 200 µg/ml for *A.*

614 *tumefaciens* and 20 µg/ml for *E. coli* DH5α. IPTG was added at a concentration of 1 mM when
615 indicated. Cumate was added at a concentration of 0.1 mM when indicated.

616

617 **Construction of expression plasmids and strains.** All strains and plasmids used are listed in
618 Table S4.1, while primers used are listed in Table S4.2. For amplification of target genes, primer
619 names indicate the primer orientation and added restriction sites. To construct expression vectors
620 containing *ftsZ_{AT}-sfgfp*, *ftsZ₁-sfgfp*, *ftsZ₃-sfgfp*, and *ldtp₀₈₄₅-sfgfp* the respective coding sequence
621 was amplified from purified C58 genomic DNA using primers indicated in Table S4.2. The
622 amplicons were digested overnight and ligated into cut pSRKKM-P_{lac}-*sfgfp* using NEB T4 DNA
623 ligase at 4°C overnight. The newly formed *sfgfp* fusion of each gene was excised from the
624 plasmid by overnight digestion with NdeI and NheI. Fragments containing *ftsZ_{AT}-sfgfp*, *ftsZ₁-*
625 *sfgfp*, *ftsZ₃-sfgfp*, and *ldtp₀₈₄₅-sfgfp* were then ligated into cut pRV-MCS2 to give constitutive
626 expression vectors containing the fusions. To construct the *popZ-yfp* expression vector, *popZ*
627 along with the upstream promoter sequence were amplified from purified C58 genomic DNA,
628 digested and ligated into pMR10.

629

630 To construct pSRKKM-P_{cym}, a synthesized gBlock from IDT Integrated DNA Technologies was
631 made containing the regulatory elements of the cumate system similar to previously described
632 plasmid constructs [76, 77]. The P_{cym} promoter region is annotated in Table S4.2. The sequence
633 encoding the cumate repressor was codon optimized for *A. tumefaciens* and placed under the
634 control of the constitutive kanamycin promoter from pSRKKM-P_{lac}-*sfgfp*. The synthesized
635 gBlock was digested overnight with EcoRI and NdeI. The resulting fragment was then ligated

636 into cut pSRKKm-P_{lac}-*sfgfp* thereby replacing the original *lac* promoter and repressor with the
637 cumate repressor and cumate regulated promoter.

638

639 Next, *yfp-parB* was excised from pSRKKM-P_{lac}-*yfp-parB* [46] and ligated into pSRKKM-P_{cym} to
640 create an expression vector compatible with the depletion strains. To create expression vectors
641 for *ftsZ_{AT}*, *ftsZ_{AT}ΔCTP*, and *ftsZ_{AT}ΔCTL* the respective target gene was amplified utilizing
642 indicated primers, digested overnight with NdeI and BamHI and ligated into pSRKKM-P_{cym}.

643 All expression vectors were verified by sequencing. All vectors were introduced into *A.*
644 *tumefaciens* strains utilizing standard electroporation protocols [78] with the addition of IPTG in
645 the media when introducing plasmids into in depletion backgrounds.

646

647 **Construction of deletion/depletion plasmids and strains.** Vectors for gene deletion by allelic
648 exchange were constructed using recommended methods for *A. tumefaciens* [78]. Briefly, 500 bp
649 fragments upstream and downstream of the target gene were amplified using primer pairs P1/P2
650 and P3/P4 respectively. Amplicons were spliced together by SOEing using primer pair P1/P4.
651 The amplicon was digested and ligated into pNTPS139. The deletion plasmids were introduced
652 into *A. tumefaciens* by mating using an *E. coli* S17 conjugation strain to create KM resistant,
653 sucrose sensitive primary exconjugants. Primary exconjugants were grown overnight in media
654 with no selection. Secondary recombinants were screened by patching for sucrose resistance and
655 KM sensitivity. Colony PCR with primers P5/P6 for the respective gene target was used to
656 confirm deletion. PCR products from P5/P6 primer sets were sequenced to further confirm
657 deletions.

658 For depletion strains, target genes (*ftsZ_{AT}*, *ftsA*, and *ftsW*) were amplified, digested and ligated
659 into either pUC18-mini-Tn7T-GM-P_{lac} or pUC18-mini-Tn7T-GM-P_{lac}. The mini-Tn7 vectors,
660 along with the pTNS3 helper plasmid, were introduced into C58Δ*tetRA*::a-*att*Tn7 as described
661 previously [32]. Transformants were selected for gentamicin resistance and insertion of the target
662 gene into the a-*att* site was verified by colony PCR using the tet forward and Tn7R109 primer.
663 PCR products were sequenced to confirm insertion of the correct gene. Next, the target gene was
664 deleted from the native locus as described above in the presence of 1 mM IPTG to drive
665 expression of the target gene from the engineered site.

666

667 **Construction of plasmids for protein expression and purification.** To construct pET21a
668 FtsZ_{AT}, *ftsZ_{AT}* was amplified from C58 genomic DNA with FtsZ_{AT} For NdeI and FtsZ_{AT} Rev
669 EcoRI, digested with NdeI and EcoRI, and ligated into similarly digested pET21a. To construct
670 pTB146 FtsZ_I, *ftsZ_I* was amplified from C58 genomic DNA with FtsZ_I For SapI and FtsZ_I Rev
671 BamHI, digested with SapI and BamHI, and ligated into similarly digested pTB146. Ligation
672 products were transformed into NEB Turbo (New England Biolabs) and selected for ampicillin
673 resistance. Insertions were verified by colony PCR and Sanger sequencing. Primers FtsZ_{AT}-
674 L72W and FtsZ_I-L71W were used to mutagenize pET21a FtsZ_{AT} and pTB146 FtsZ_I,
675 respectively, using the Quikchange Multi Lightning Mutagenesis Kit (Agilent) and following the
676 manufacturer's protocol to generate pET21a FtsZ_{AT}-L72W and pTB146 FtsZ_I-L71W. Mutations
677 in the targeted sites were verified by Sanger sequencing.

678 pET21c FtsZ_{AT}+CTL and pET21c FtsZ_{AT}ΔCTL were constructed in several steps. First, the
679 GTPase domain of *ftsZ_{AT}* was amplified from C58 genomic DNA using FtsZ_{AT} For NdeI and
680 FtsZ_{AT} GTPase Rev KpnI SacI, split into two aliquots, and digested with NdeI and KpnI or NdeI

681 and SacI. The CTL region of *ftsZ_{AT}* was amplified from C58 genomic DNA using FtsZ_{AT} CTL
682 For KpnI and FtsZ_{AT} CTL Rev SacI and digested with KpnI and SacI. For FtsZ_{AT}+CTL, the
683 GTPase domain amplicon (digested with NdeI and KpnI) was combined with the CTL amplicon
684 (digested with KpnI and SacI) and together they were ligated into pXCFPN-1 [79] digested with
685 NdeI and SacI. For FtsZ_{AT}ΔCTL, the GTPase domain amplicon (digested with NdeI and SacI)
686 was ligated into pXCFPN-1 digested with NdeI and SacI. Each was transformed into NEB
687 Turbo, selected for spectinomycin resistance, and confirmed by colony PCR and Sanger
688 sequencing. Next, the CTP was added to each of the above constructs by annealing oligos FtsZ_{AT}
689 CTP + and FtsZ_{AT} CTP – (engineered with overhangs compatible with SacI and NheI ligation)
690 and ligating into the above constructs digested with SacI and NheI. Each was transformed into
691 NEB Turbo and confirmed as above to generate pX1 FtsZ_{AT}+CTL and pX1 FtsZ_{AT}ΔCTL.
692 Finally, FtsZ_{AT}+CTL and FtsZ_{AT}ΔCTL were subcloned into pET21c by digestion of pX1
693 FtsZ_{AT}+CTL and pX1 FtsZ_{AT}ΔCTL with NdeI and NheI and ligating into similarly digested
694 pET21c. Each was transformed into NEB Turbo, selected for ampicillin resistance, and
695 confirmed by colony PCR and Sanger sequencing.

696

697 **DIC and phase contrast microscopy.** Exponentially growing cells (OD₆₀₀ = ~0.6) were spotted
698 on 1% agarose ATGN pads as previously described [80]. Microscopy was performed with an
699 inverted Nikon Eclipse TiE with a QImaging Rolera em-c² 1K EMCCD camera and Nikon
700 Elements Imaging Software. For time-lapse microscopy, images were collected every ten
701 minutes, unless otherwise stated.

702

703 **Fluorescence microscopy.** Plasmid encoded FtsZ_{AT}-sfGFP, FtsZ₁-sfGFP, FtsZ₃-sfGFP, and
704 LDTP₀₈₄₅-sfGFP fusions were expressed from the P_{van} promoter, which provides constitutive low
705 levels of expression (Figure 6- Figure Supplement 1C). Plasmid encoded FtsA-sfGFP and PopZ-
706 YFP fusions were expressed from the native promoters. Expression of plasmid encoded YFP-
707 ParB was induced by the presence of 0.1 mM cumate for 2 hours (h). Cells containing plasmids
708 with fluorescent protein fusions were grown to exponential phase before imaging on agarose
709 pads.

710 To visualize DNA, 1 ml of exponentially growing cells was treated with 1 µl of Sytox Orange
711 for 5 minutes. Cells were collected by centrifugation and washed with PBS 2 times followed by a
712 final resuspension in PBS. Cells were then imaged on agarose pads.

713 To visualize sites of active peptidoglycan synthesis 1 ml of exponentially growing cells was
714 labeled with the fluorescent D-amino acid (FDAA), HCC amino-D-alanine (HADA), as
715 previously described [49, 80].

716

717 **Cell viability and growth curve assays.** For cell viability spot assays, exponentially growing
718 cultures were diluted to OD₆₀₀ = 0.1 and serially diluted in ATGN. 3 µl of each dilution was
719 spotted onto ATGN and incubated at 28°C for 3 days before imaging. When appropriate ATGN
720 plates contained KM 300 µg/ml, IPTG 1mM, and cumate 0.1 mM as indicated in figure legends.
721 For growth curve analysis, exponentially growing cultures were diluted to OD₆₀₀ = .05 in 200 µl
722 of ATGN in 96-well plates. Plates were shaken for 1 minute before OD₆₀₀ readings, which were
723 taken every 10 minutes.

724

725 **Cell morphology and constriction rate analysis.** Exponentially growing cells were imaged
726 using phase contrast microscopy as described above. Cell length, area, and constrictions were
727 detected using MicrobeJ software [81].

728 To calculate constriction rates, cells with detectable constrictions were tracked using time-lapse
729 microscopy. The width of the cell constriction was measured at an initial time-point and the
730 measurement was repeated after 10 minutes. The difference in constriction width was divided by
731 the 10-minute time interval to give a constriction rate.

732

733 **Western blot analysis.** For western blot analysis of FtsZ depletion, the *ftsZ* depletion strain was
734 grown in 40 ml ATGN with 1 mM IPTG to exponential phase. 2 ml of culture was collected
735 prior to depletion (time 0) by centrifugation at 10,000 x g for 3 minutes. The remaining culture
736 was collected by centrifugation at 3500 x g for 10 minutes, and supernatants were discarded.
737 Cells were washed in sterile water and pelleted again. To deplete FtsZ, the pellet was
738 resuspended in fresh ATGN without IPTG and grown under standard culturing conditions. 2-ml
739 samples were collected by centrifugation after 30, 45, 60, 120, and 240 minutes of depletion.
740 OD₆₀₀ was taken for each sample prior to centrifugation so that samples could be normalized to
741 an OD₆₀₀ equivalent to 0.68. The cell pellets were incubated with 100 µl of a master mix
742 containing 1 ml of BugBuster protein extraction reagent (Novagen) and supplemented with 1
743 EDTA-free protease inhibitor cocktail (Sigma), 10 µl of lysonase (Novagen), 2,500 U/ml DNase
744 I (Thermo Scientific), and 1 mM dithiothreitol (DTT) (Thermo Scientific) for 25 minutes with
745 shaking at room temperature to lyse the cell pellets. The whole-cell lysates were clarified by
746 centrifugation at 10,000 rpm for 15 min. A final concentration of 1 X Laemmli buffer was added
747 to the cleared cell lysates. Samples were boiled at 100°C for 5 min prior to loading on a 4-15%

748 Mini-PROTEAN TGX Precast Gel (Bio-Rad). The separated proteins were electroblotted onto
749 polyvinylidene difluoride (PVDF) membranes (Bio-Rad) and blocked overnight in 5% nonfat
750 dry milk powder solubilized in 1% TBST (Tris-buffered saline [TBS], 1% Tween 20). The
751 blocked PVDF membranes were probed with *Escherichia coli* anti-FtsZ (1:3000) monoclonal
752 antibody (gift from Joe Lutkenhaus) for 1.5 h in 5% milk-TBST, followed by incubation with
753 anti-rabbit (1:5000) HRP (Pierce 31460) secondary antibody for 1 h in 5% milk-TBST. The
754 secondary antibody was detected using the ECL Plus HRP substrate (Thermo Scientific Pierce).

755

756 For comparison of expression from P_{van} , P_{lac} , and P_{cym} promoters, strains were grown in 2 ml
757 ATGN with 200 ug/mL KM to exponential phase. P_{lac} and P_{cym} were induced with 1 mM IPTG
758 and 50 μ M cumate, respectively for 4 h. Cell pellets were lysed as described above and clarified
759 whole-cell lysates were boiled with 1 X Laemmli buffer for 5 min prior to loading on 4-15%
760 Mini-PROTEAN TGX Precast Gel (Bio-Rad). The separated proteins were electroblotted onto
761 PVDF membranes (Bio-Rad), blocked as described above, and probed with anti-GFP (1:3,000)
762 monoclonal antibody (Thermo Scientific Pierce) for 1 h in 5% milk-TBST, followed by
763 incubation with a donkey anti-mouse (1:300) horseradish peroxidase-conjugated secondary
764 antibody (Thermo Scientific Pierce) for 1 h in 5% milk-TBST. The secondary antibody was
765 detected using the ECL Plus HRP substrate (Thermo Scientific Pierce).

766

767 **Protein expression and purification.** FtsZ_{AT}, FtsZ_{AT}-L72W, FtsZ_{AT}+CTL (FtsZ_{AT} with
768 restriction sites flanking the CTL), and FtsZ_{AT} Δ CTL (FtsZ_{AT} with the CTL deleted, containing
769 the same restriction sites at the GTPase-CTC junction as in FtsZ_{AT}+CTL) were expressed and
770 purified in untagged form. Each was produced from a pET21 expression vector (pEG1555 –

771 FtsZ_{AT}, pEG1556 - FtsZ_{AT}-L72W, pEG1444 - FtsZ_{AT}+CTL, pEG1445 - FtsZ_{AT}ΔCTL) in
772 *Escherichia coli* Rosetta(DE3)pLysS induced at 37°C for 4 h with 0.5 mM IPTG after OD
773 reached 0.8 to 1.0 OD at 600 nm. Cells were harvested by centrifugation at 6000 x g and
774 resuspended in 30 mL FtsZ QA buffer (50 mM Tris-HCl pH 8, 50 mM KCl, 0.1 mM EDTA,
775 10% glycerol) per liter of culture. Resuspensions were snap frozen in liquid nitrogen and stored
776 at -80°C until purification. To purify, resuspensions were thawed quickly and cells were lysed by
777 incubation with 1 mg/mL lysozyme, 2.5 mM MgCl₂, DNase I, 2 mM PMSF, and a cComplete
778 mini EDTA-free protease inhibitor tablet (Roche) for 45 min to 1 h at room temperature
779 followed by sonication. Lysates were cleared by centrifugation at 15000 x g for 30 min at 4°C
780 and filtered through a 0.45 μm filter before anion exchange chromatography (HiTrap Q HP 5
781 mL, GE Life Sciences). Protein was eluted with a linear KCl gradient (FtsZ QA buffer with 50 to
782 500 mM KCl) and fractions containing FtsZ were verified by SDS-PAGE, pooled, and subjected
783 to ammonium sulfate precipitation. Precipitates (at 17-20% ammonium sulfate saturation
784 depending on the variant) were verified by SDS-PAGE, resuspended in HEK50G (50 mM
785 HEPES-KOH pH 7.2, 0.1 mM EDTA, 50 mM KCL, 10% glycerol, 1 mM β-mercaptoethanol),
786 and further purified by gel filtration (Superdex 200 10/300 GL, GE Life Sciences). Peak
787 fractions were pooled, snap frozen in liquid nitrogen, and stored at -80°C.

788

789 FtsZ₁ and FtsZ₁-L71W were produced as His₆-SUMO fusions and cleaved to yield untagged,
790 scarless proteins. Each was produced from a pTB146 expression vector (pEG1535 - FtsZ₁,
791 pEG1542 - FtsZ₁-L71W) in *E. coli* Rosetta (DE3)pLysS as described above. Cells were
792 harvested by centrifugation as above, resuspended in HK300G (50 mM HEPES-KOH pH7.2,
793 300 mM KCl, 10% glycerol) with 20 mM imidazole, snap frozen in liquid nitrogen, and stored at

794 -80°C until purification. To purify, resuspensions were thawed quickly and cells were lysed by
795 incubation with 1 mg/mL lysozyme, 2.5 mM MgCl₂, and DNase I for 45 min at room
796 temperature followed by sonication. Lysate was cleared and filtered as described above. Protein
797 was isolated by Ni²⁺ affinity chromatography (HisTrap FF 1 mL, GE Life Sciences) and eluted in
798 HK300G with 300 mM imidazole. Fractions containing His₆-SUMO fusions were verified by
799 SDS-PAGE, combined with Ulp1 Sumo protease at a 1:100 (protease:FtsZ) molar ratio, and
800 cleaved by incubation at 30°C for 3.5 h. Cleaved FtsZ₁ or FtsZ₁L71W was purified away from
801 His₆-SUMO by gel filtration (Superdex 200 10/300 GL, GE Life Sciences) in HEK50G. Peak
802 fractions were pooled, snap frozen in liquid nitrogen, and stored at -80°C.

803

804 **Polymerization kinetics assays.** A Fluoromax-3 spectrofluorometer (Jobin Yvon, Inc) was used
805 to monitor FtsZ polymerization by right-angle light scattering and tryptophan fluorescence. FtsZ₁
806 and/or FtsZ_{AT} (wild-type or L71W/L72W mutants, as indicated in figures and text) was
807 polymerized in HEK50 (50 mM HEPES-KOH pH 7.2, 50 mM KCl, 0.1 mM EDTA) with 2.5
808 mM MgCl₂. 2 mM GTP was used to induce polymerization for light scattering and 50 μM GTP
809 was used to induce polymerization for tryptophan fluorescence (GTP is fluorescent at the
810 wavelengths used, so low concentrations must be used). GTP was added after baseline light
811 scatter or fluorescence was established. For light scattering, samples were excited at 350 nm and
812 scatter was detected at 350 nm with slits set to 2 nm. For tryptophan fluorescence, samples were
813 excited at 295 nm and emission was detected at 344 nm, with 2 nm slits.

814

815 **GTPase assay.** FtsZ₁ and/or FtsZ_{AT} was polymerized in HEK50 with 2.5 mM MgCl₂ and 2 mM
816 GTP. FtsZ_{AT}+CTL or FtsZ_{AT}ΔCTL was polymerized in HEK50 or HEK300 (same as HEK50 but
817 with 300 mM KCl) as indicated, with 10 mM MgCl₂ and 2 mM GTP. GTP was added at time 0.
818 Reaction was stopped at 5, 10, 15, 20, and 30 minutes with quench buffer (50 mM HEPES-KOH
819 pH 7.2, 21.3 mM EDTA, 50 mM KCl). Inorganic phosphate in solution (liberated by GTP
820 hydrolysis) over time was measured using SensoLyte MG Phosphate Assay Kit Colorimetric
821 (AnaSpec, Inc, Fremont, California).

822
823 **Negative stain transmission electron microscopy (TEM).** FtsZ₁ and/or FtsZ_{AT} were
824 polymerized in HEK50 with 2.5 mM MgCl₂ and 2 mM GTP. 4 μM FtsZ_{AT}+CTL or FtsZ_{AT}ΔCTL
825 were polymerized in HEK50 or HEK300 as indicated with 10 mM MgCl₂ and 2 mM GTP. After
826 a 15-minute incubation at room temperature, samples were applied to carbon-coated glow-
827 discharged grids with 0.75% uranyl formate staining as previously described [17, 82]. TEM
828 samples were imaged using a Philips/FEI BioTwin CM120 TEM equipped with an AMT XR80 8
829 megapixel CCD camera (AMT Imaging, USA).

830
831 **Peptidoglycan composition analysis**

832 Six cultures of WT and *ftsZ* depletion cells were grown in 10 ml of ATGN with IPTG to
833 exponential phase. The 10 ml cell cultures were added to 40 ml of fresh media. The 50 ml cultures
834 were grown to exponential phase and pelleted by centrifugation at 4000 x g for 10 minutes. Cell
835 pellets were washed three times with ATGN by centrifugation and resuspension to remove IPTG.
836 After the final wash 3 cell pellets were resuspended in 50 ml ATGN and the remaining 3 pellets

837 were resuspended in 50 ml ATGN with 1 mM IPTG. Each culture was grown for 14 h. The optical
838 densities of the cells were monitored to ensure the optical density of the cultures never went above
839 $OD_{600} = 0.7$ to avoid changes to peptidoglycan content due to stationary phase. If necessary, fresh
840 medium was added to dilute the cultures to maintain exponential growth. After 14 h of growth, 50
841 ml of the exponential cultures were collected and pelleted by centrifugation at $4000 \times g$ for 20
842 minutes. Cell pellets were resuspended in 1 mL of ATGN and 2 mL of 6% SDS and stirred with
843 magnets while boiling for 4 h. After 4 h, samples were removed from heat but continued to stir
844 overnight. Samples were then shipped to Dr. Felipe Cava's laboratory for purification and analysis.
845
846 Upon arrival, cells were boiled and simultaneously stirred by magnets for 2 h. After 2 h, boiling
847 was stopped and samples were stirred overnight. Peptidoglycan was pelleted by centrifugation for
848 13 min at 60000 rpm (TLA100.3 Beckman rotor, Optima Max-TL ultracentrifuge; Beckman), and
849 the pellets were washed 3 to 4 times by repeated cycles of centrifugation and resuspension in water.
850 The pellet from the final wash was resuspended in 50 μ l of 50 mM sodium phosphate buffer, pH
851 4.9, and digested overnight with 100 μ g/ml of muramidase at 37°C. Muramidase digestion was
852 stopped by boiling for 4 min. Coagulated protein was removed by centrifugation for 15 min at
853 15000 rpm in a desktop microcentrifuge. The muropeptides were mixed with 15 μ l 0.5 M sodium
854 borate and subjected to reduction of muramic acid residues into muramitol by sodium borohydride
855 (10 mg/ml final concentration, 20 min at room temperature) treatment. Samples were adjusted to
856 pH 3 to 4 with orthophosphoric acid and filtered (0.2- μ m filters).

857
858 Muropeptides were analyzed on a Waters UPLC system equipped with an ACQUITY UPLC BEH
859 C18 Column, 130 Å, 1.7 μ m, 2.1 mm \times 150 mm (Waters) and a dual wavelength absorbance

860 detector. Elution of muropeptides was detected at 204 nm. Muropeptides were separated at 45°C
861 using a linear gradient from buffer A [formic acid 0.1% (v/v)] to buffer B [formic acid 0.1% (v/v),
862 acetonitrile 20% (v/v)] in a 12 min run with a 0.250 ml/min flow. Peptidoglycan compositional
863 analysis on triplicate samples was completed on two separate occasions.

864

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868

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875

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1136 **Figure Legends**

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1138 **Figure 1. Characterization of FtsZ homologs in *A. tumefaciens*.** A) Domain schematic of FtsZ
1139 homologs in *A. tumefaciens*. Note that domains are not drawn to scale. B) Representative image
1140 of localization patterns for each homolog. FtsZ_{AT}-sfGFP and FtsZ₁-sfGFP show mid-cell ring
1141 formation while FtsZ₃-sfGFP fails to localize. C) Cell viability is shown by spotting serial
1142 dilutions. $\Delta ftsZ_1$, $\Delta ftsZ_3$, and $+ftsZ_{AT}$ have similar viability to WT, while $-ftsZ_{AT}$ displays a drastic
1143 decrease in viability. D) Cell morphology and microcolony formation of $\Delta ftsZ_1$ and $\Delta ftsZ_3$ are
1144 similar to WT, while $-ftsZ_{AT}$ results in long branched cells that fail to divide. All scale bars are set
1145 to 2 μ m.

1146 **Figure 1- Figure Supplement 1. Cell growth and morphology of *ftsZ* mutants.** A.) Western
1147 blot analysis showing the depletion of FtsZ_{AT} after the removal of inducer. B) Cell lengths of WT,
1148 $\Delta ftsZ_1$, $\Delta ftsZ_3$, and induced $ftsZ_{AT}$ are indistinguishable. C) Cell area in WT and induced $ftsZ_{AT}$
1149 are the same while cells depleted of FtsZ_{AT} for 8 and 14 hours accumulate cell area.

1150 **Figure 2. Deletion of multiple *ftsZ* homologs does not yield an additive effect.** A) Cell
1151 viability (top) and morphology (bottom) of the double mutant $\Delta ftsZ_1\Delta ftsZ_3$ is indistinguishable
1152 from WT. B) Cell viability (top) and morphology (bottom) of $\Delta ftsZ_1$, $\Delta ftsZ_3$, or $\Delta ftsZ_1\Delta ftsZ_3$
1153 during FtsZ_{AT} depletion are indistinguishable from FtsZ_{AT} depletion alone. All scale bars are set
1154 to 2 μ m. Black bar denotes $ftsZ_{AT}$ depletion strain background.

1155 **Figure 3. FtsZ₁ requires FtsZ_{AT} to polymerize *in vitro* and to localize in cells.** A) FtsZ₁-
1156 sfGFP forms midcell rings which constrict in WT and when $ftsZ_{AT}$ is induced. FtsZ₁-sfGFP fails
1157 to constrict early rings and disassembles during FtsZ_{AT} depletion. Scale bar is set to 2 μ m. B)

1158 Light scattering over time for purified proteins at the indicated concentrations. FtsZ_{AT}
1159 polymerizes at concentrations above 2 μ M, but FtsZ_I does not polymerize. GTP (2 mM) was
1160 added where indicated by the arrow to induce polymerization. Experiments were performed in
1161 triplicate and mean curves are shown. C) Negative stain TEM of the indicated proteins. Co-
1162 polymers of FtsZ_{AT} and FtsZ_I are indistinguishable from FtsZ_{AT} polymers. Scale bar is set to 100
1163 nm. D) Inorganic phosphate concentration in solution over time in the presence of the indicated
1164 proteins and protein concentrations. Co-polymers of FtsZ_{AT} and FtsZ_I consume GTP at the same
1165 rate as FtsZ_{AT} homopolymers at equivalent total FtsZ concentration. Reactions were performed
1166 in triplicate and mean \pm standard error is plotted. E) Tryptophan fluorescence over time for the
1167 indicated proteins. FtsZ_I-L71W (red) shows no polymerization (Trp fluorescence) on its own, but
1168 can co-polymerize with added FtsZ_{AT} (purple). GTP (50 μ M) was added where indicated by the
1169 arrow to induce polymerization. Experiments were performed in triplicate and representative
1170 curves are shown.

1171 **Figure 4. Characterization of genomic content during FtsZ_{AT} depletion.** A) Timelapse
1172 microscopy in minutes demonstrates proper cell division and microcolony formation in +*ftsZ_{AT}*
1173 induced with IPTG (top panel). Timelapse during depletion of FtsZ_{AT} demonstrates branches
1174 forming from tip splitting events (bottom panel). B) Timelapse microscopy shows that PopZ-
1175 YFP maintains polar localization during elongation and dissociates moving to the mid-cell at
1176 division in WT and when *ftsZ_{AT}* is induced. PopZ-YFP becomes trapped at the growth poles
1177 during FtsZ_{AT} depletion. C) Sytox Orange labeled DNA is diffuse throughout young cells and
1178 separated nucleoids are seen in late divisional cells in WT and when FtsZ_{AT} is induced. These
1179 DNA free regions are marked with an asterisk. Nucleoids fail to form, as diffuse DNA labeling is
1180 observed during FtsZ_{AT} depletion at both 8 h and 14 h depletion timepoints. D) Timelapse

1181 microscopy demonstrates YFP-ParB1 form a single focus at the old pole in WT and when *ftsZ_{AT}*
1182 is induced. A second focus then translocates the cell length to the growth pole. Timelapse
1183 microscopy during FtsZ_{AT} depletion reveals a third focus which translocates the cell towards an
1184 ectopic pole. E) Quantitation of YFP-ParB1 foci plotted against cell area. Number of foci
1185 increase as cell area increases. All scale bars are set to 2 μm.

1186 **Movie 1. Growth and morphological changes during FtsZ_{AT} depletion.** Cells were washed to
1187 remove inducer and grown in liquid ATGN for 4 hours before spotting on a ATGN pad. Images
1188 were acquired every ten minutes and movie is played at 10 frames per second for a total of 145
1189 frames.

1190 **Figure 5. Characterization of polar peptidoglycan synthesis during FtsZ_{AT} depletion.** A)
1191 FDAA labels active peptidoglycan synthesis at a single growing pole and septum in WT and
1192 cells depleted of FtsZ_{AT} for 0 h. As FtsZ_{AT} is depleted for 8 and 14 h, multiple growth poles are
1193 labeled, and septum labeling is lost. B) Quantitation of the major muropeptide peaks in *ftsZ_{AT}*
1194 depletion strain induced and depleted. C) Abundance of total monomers, dimers, and trimers in
1195 the muropeptide profile in *ftsZ_{AT}* depletion strain. D) Abundance of total LD and DD
1196 crosslinkage in *ftsZ_{AT}* depletion strain induced. For B,C,and D, data shown are the average
1197 abundance of each muropeptide and are taken from analysis of three independent biological
1198 samples of *ftsZ_{AT}* depletion strain induced (black bars) and depleted for 14 h (gray bars).
1199 Statistical was calculated by t-tests and is indicated with an asterisk (P-value <0.05 (*), <0.005
1200 (**), <0.001 (***)). E) Timelapse microscopy of LDTP₀₈₄₅-sfGFP in WT and when *ftsZ_{AT}* is
1201 induced yields growth pole localization during elongation and mid-cell localization during
1202 septum formation. In cells depleted of FtsZ_{AT}, localization is trapped at the growing poles. All
1203 scale bars are set to 2 μm.

1204 **Figure 5- Figure Supplement 1. Peptidoglycan analysis of control strains.** A) UPLC spectra
1205 of mucopeptides derived from WT cells. Major mucopeptides are labeled. M= monomers, D=
1206 dimers, T= trimers. Numbers indicate the length of the mucopeptide stems and the position of
1207 crosslink in dimers and trimers. B) Quantitation of the major mucopeptide peaks in WT with
1208 IPTG (black), WT without IPTG (black with gray outline), and *ftsZ_{AT}* depletion strain induced
1209 with IPTG (Gray). Data shown is the average abundance of each mucopeptide and is taken from
1210 analysis of three independent biological samples. Statistical significance is indicated with an
1211 asterisk.

1212 **Figure 6. Functional analysis of FtsZ_{AT}ΔCTL and FtsZ_{AT}ΔCTP.** A) Cell viability is measured
1213 by spotting serial dilutions of *ftsZ_{AT}* variants in the *ftsZ_{AT}* depletion background. When
1214 chromosomal *ftsZ_{AT}* is induced by IPTG and plasmid driven *ftsZ_{AT}* variants are uninduced, all
1215 strains have similar viability (top left). When both chromosomal *ftsZ_{AT}* is uninduced and plasmid
1216 driven *ftsZ_{AT}* variants are uninduced, all strains exhibit an equal decrease in viability (top right).
1217 When chromosomal *ftsZ_{AT}* is uninduced and plasmid driven *ftsZ_{AT}* variants are induced by
1218 cumate, FtsZ_{AT} expression rescues viability, FtsZ_{AT}ΔCTP partially rescues, and FtsZ_{AT}ΔCTL
1219 fails to rescue viability (bottom left). When both chromosomal *ftsZ_{AT}* is induced with IPTG and
1220 plasmid driven *ftsZ_{AT}* variants are induced by cumate, FtsZ_{AT}ΔCTL expression reduces viability
1221 while other variants have no impact (bottom right). B) Representative images displaying
1222 morphology and FDAA labeling while chromosomal *ftsZ_{AT}* is uninduced and plasmid driven
1223 *ftsZ_{AT}* variants are induced for 6 and 14 h. C) Timelapse microscopy while chromosomal *ftsZ_{AT}*
1224 is uninduced and plasmid driven FtsZ_{AT}ΔCTP is expressed reveal that polar growth fails to
1225 terminate and undergoes tip splitting, although septum formation and cell division also take place
1226 (top panel). Timelapse microscopy while chromosomal *ftsZ_{AT}* is uninduced and plasmid driven

1227 FtsZ_{AT}ΔCTL is expressed shows termination of polar growth and new pole formation near mid-
1228 cell (bottom panel).

1229 **Figure 6- Figure Supplement 1. Validation of a cumate inducible vector in *A. tumefaciens*.**

1230 A) Sequence schematic of the cumate operon modified for use is pSRKKM-*sfGFP*. Regions are
1231 color coded to match sequence found in Table S2. B) Representative image of WT cells
1232 harboring pSRKKM-Pcym-*sfGFP* uninduced (left) and induced (right). C) Western blot analysis
1233 comparing expression levels of *sfGFP* expressed from pRV, pSRKKM-Plac, and pSRKKM-
1234 Pcym. D) Growth curve analysis of WT cells harboring pSRKKM-Pcym-empty induced with
1235 different levels of cumate (left). pSRKKM-Pcym-*ftsZ_{AT}* rescues chromosomal FtsZ_{AT} depletion
1236 with 0.01mM cumate (right).

1237 **Figure 7. FtsZ_{AT} requires the CTL for robust PG biosynthesis and proper polymerization.**

1238 A.) Quantitation of the major muropeptide peaks in *ftsZ_{AT}* depletion strain expressing an empty
1239 plasmid, full length FtsZ_{AT}, or FtsZ_{AT}ΔCTL. B) Abundance of total monomers, dimers, and
1240 trimers in the muropeptide profile. C) Abundance of total LD and DD crosslinkage. For A, B,
1241 and C, data shown are the average abundance of each muropeptide and are taken from analysis
1242 of three independent biological samples. Statistical significance is indicated with an asterisk. D.)
1243 Negative stain TEM of the 4 μM of the indicated protein in the presence of 50 or 300 mM KCl.
1244 FtsZ_{AT}ΔCTL shows increased propensity to bundle at high salt. Scale bar is set at 100 nm. E.)
1245 Phosphate in solution over time in the presence of indicated proteins in solution with 50 or 300
1246 mM KCl. The rate of GTP hydrolysis by FtsZ_{AT}ΔCTL is reduced under high salt conditions that
1247 promote bundling.

1248 **Figure 8. FtsA is not required for termination of polar growth.** A) FtsA-*sfGFP* persists at
1249 growth poles and forms mid-cell rings which constrict in WT (top panel). FtsA-*sfGFP* becomes

1250 trapped at the growth poles and foci split as the growth poles split during FtsZ_{AT} depletion
1251 (bottom panel). B) Timelapse microscopy shows cells expressing FtsA grow and divide normally
1252 forming microcolonies (top panel). Cells depleted of FtsA terminate polar growth and form new
1253 growth poles near the mid-cell (bottom panel). C) FDAAs label a single growth pole when FtsA
1254 is present (top) and label multiple poles emerging from the mid-cell when FtsA is absent
1255 (bottom). D) Timelapse microscopy during FtsA depletion shows PopZ-YFP localizes to the
1256 growth poles and dissociates as growth is terminated. It then reappears at the new pole sites
1257 (top). During FtsA depletion, FtsZ-sfGFP forms rings marking the future sites of pole formation
1258 (bottom). All scale bars are set to 2 μ m.

1259 **Movie 2. Growth and morphological changes during FtsA depletion.** Cells were washed to
1260 remove inducer and grown in liquid ATGN for 2 hours before spotting on a ATGN pad. Images
1261 were acquired every ten minutes and movie is played at 10 frames per second for a total of 97
1262 frames.

1263 **Figure 9. FtsW is not required for termination of polar growth** A) Timelapse microscopy
1264 shows cells expressing FtsW grow and divide normally forming microcolonies (top panel). Cells
1265 depleted of FtsW terminate polar growth and form new growth poles near the mid-cell (bottom
1266 panel). B) FDAA labels a single growth pole when FtsW is present (top) and labels multiple
1267 poles emerging from the mid-cell when FtsW is absent (bottom). C) Timelapse microscopy
1268 during FtsW depletion shows PopZ-YFP localizes to the growth poles and dissociates as growth
1269 is terminated. It then reappears at the new pole sites (top). During FtsW depletion, FtsZ-sfGFP
1270 forms rings marking the future sites of pole formation (bottom).

1271

1272 **Movie 3. Growth and morphological changes during FtsW depletion.** Cells were washed to
1273 remove inducer and grown in liquid ATGN for 4 hours before spotting on a ATGN pad. Images
1274 were acquired every ten minutes and movie is played at 10 frames per second for a total of 85
1275 frames.

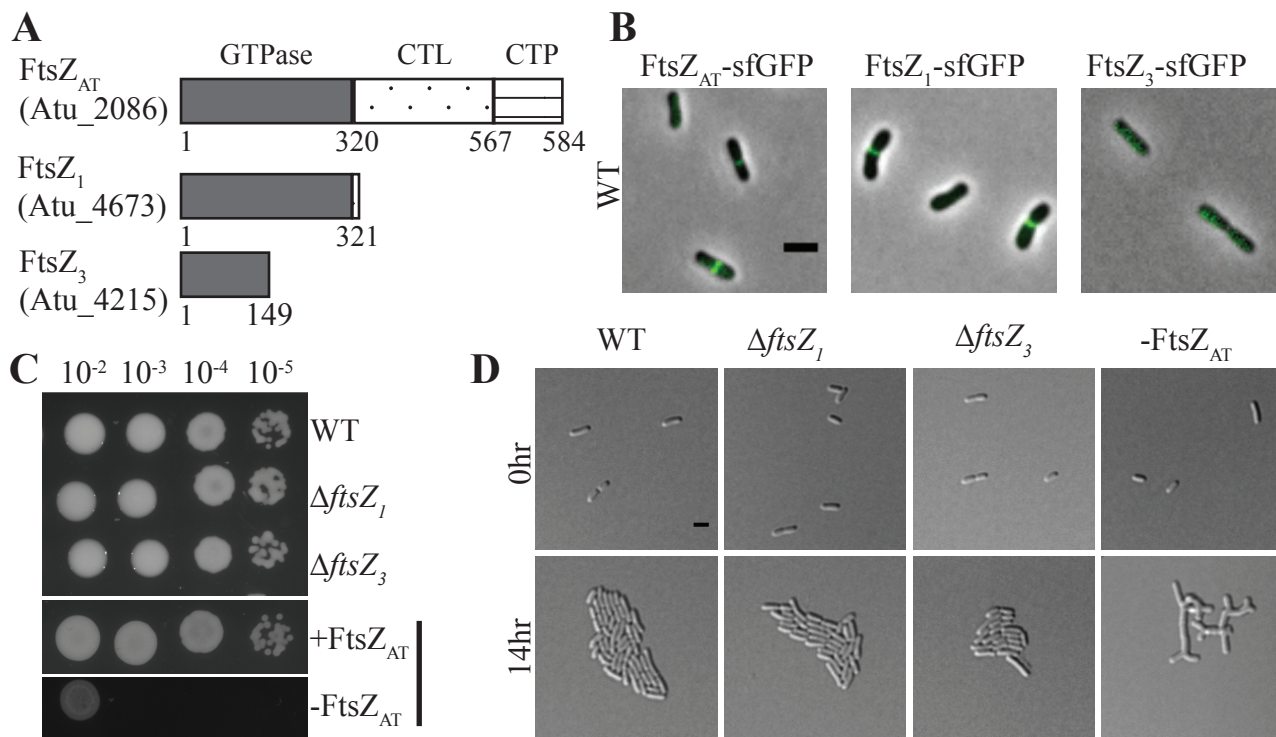


Figure 1

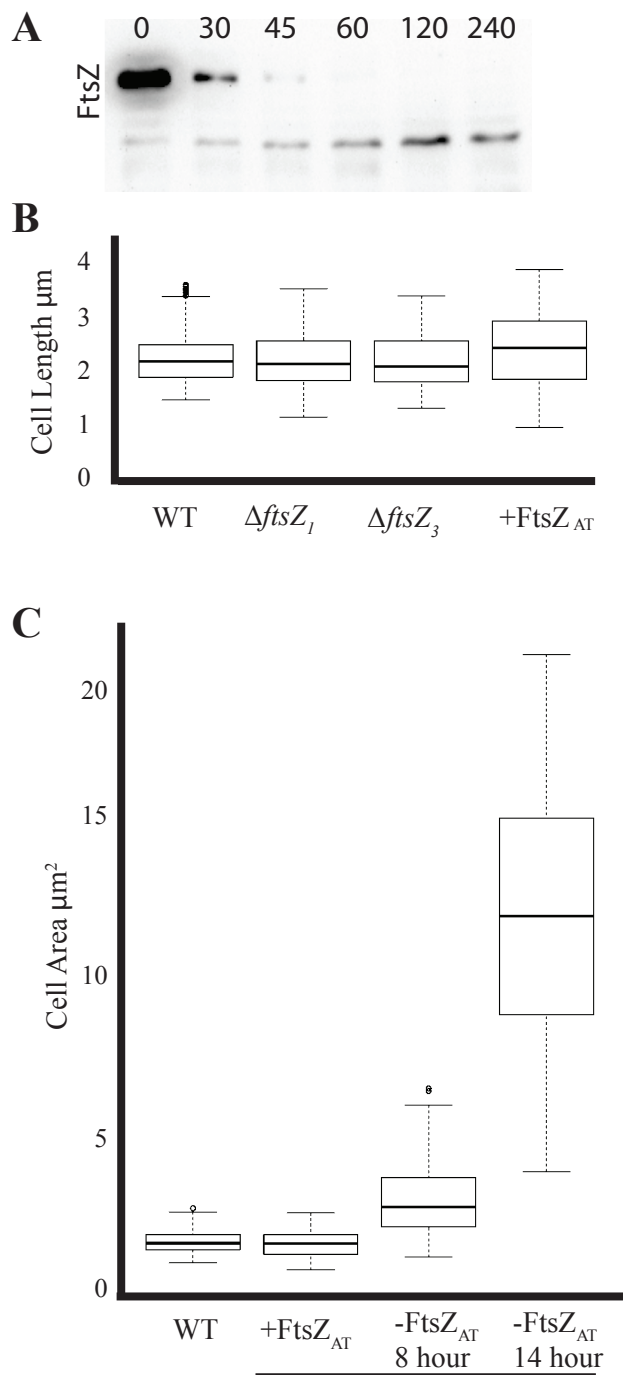


Figure 1- Figure Supplement 1

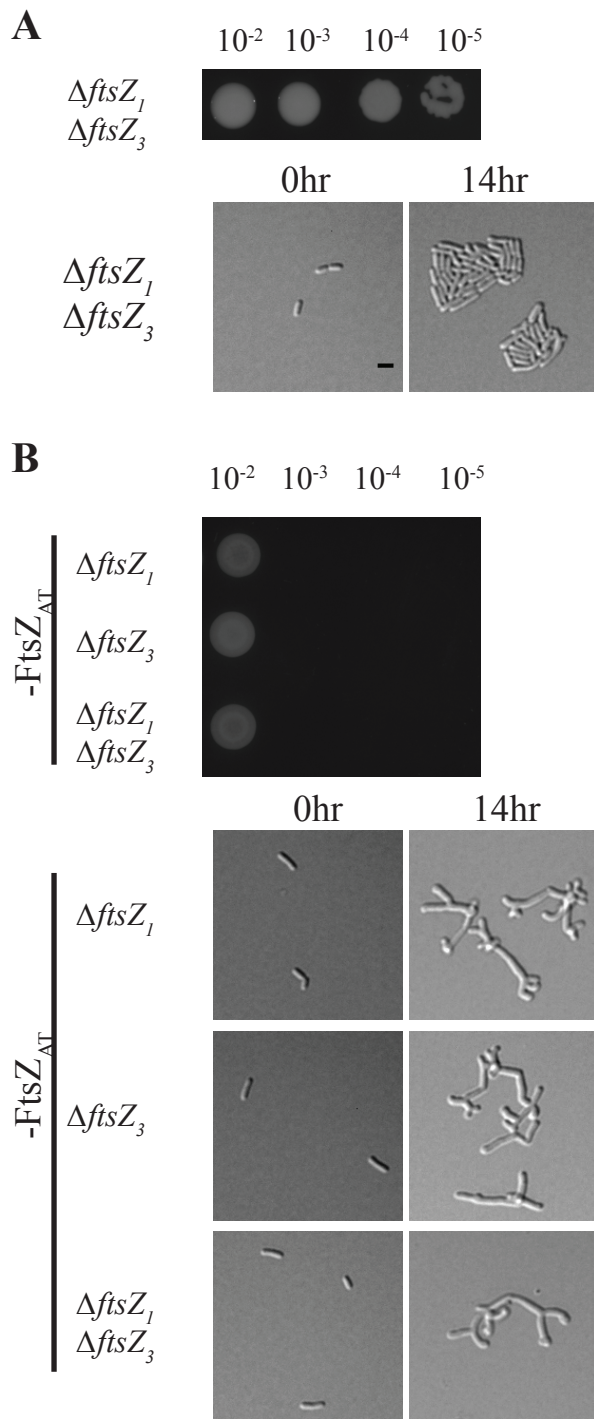


Figure 2

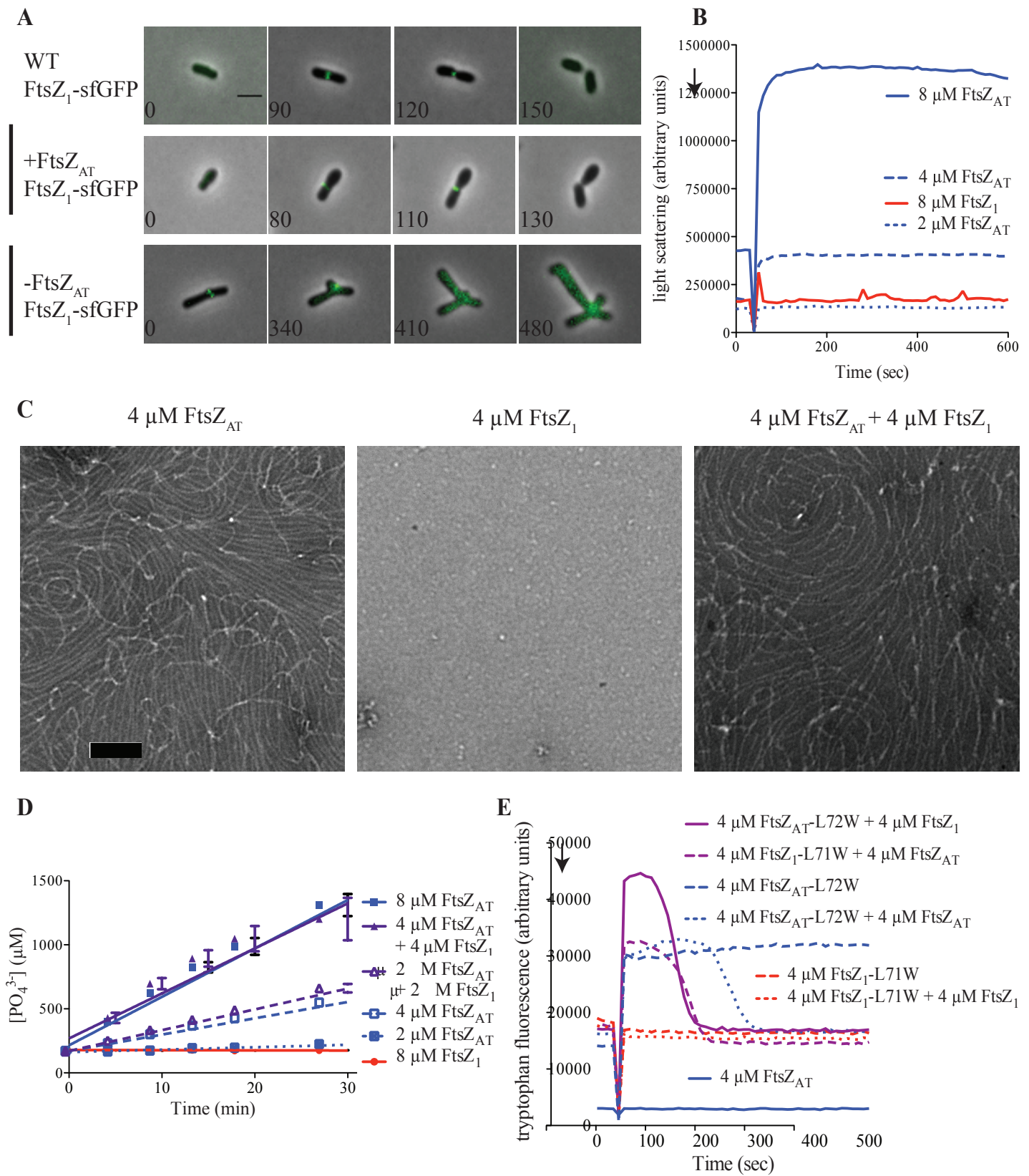


Figure 3

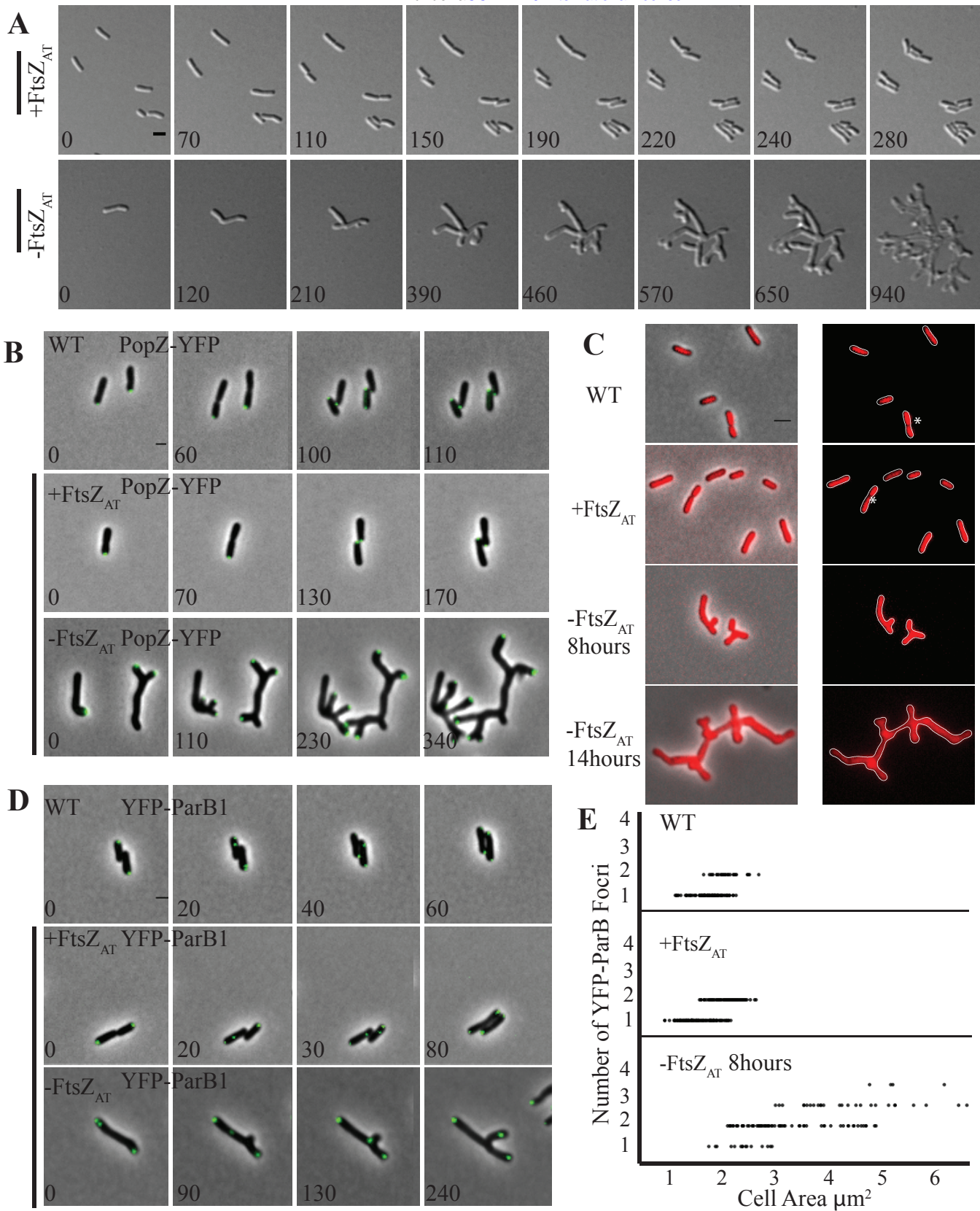


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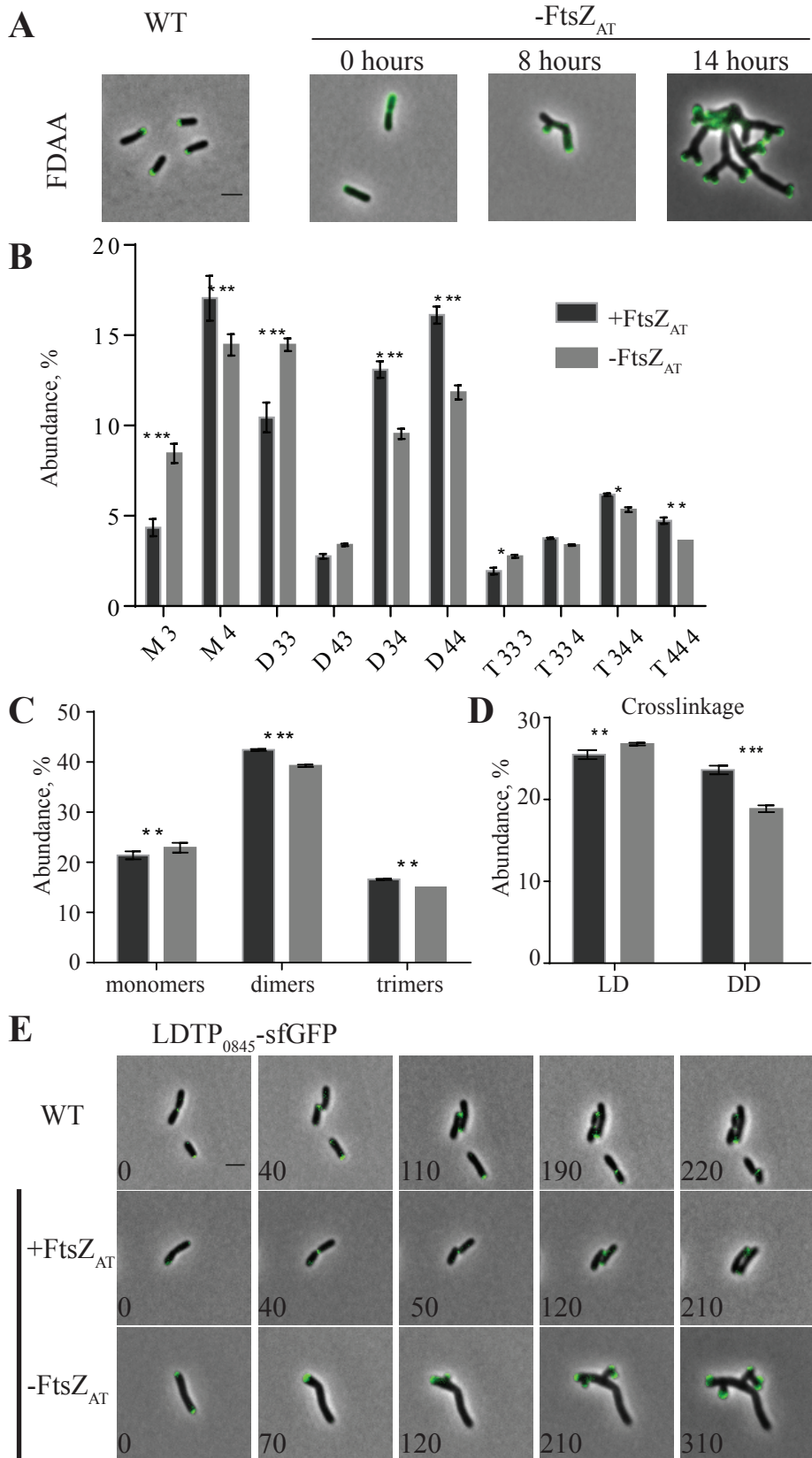


Figure 5

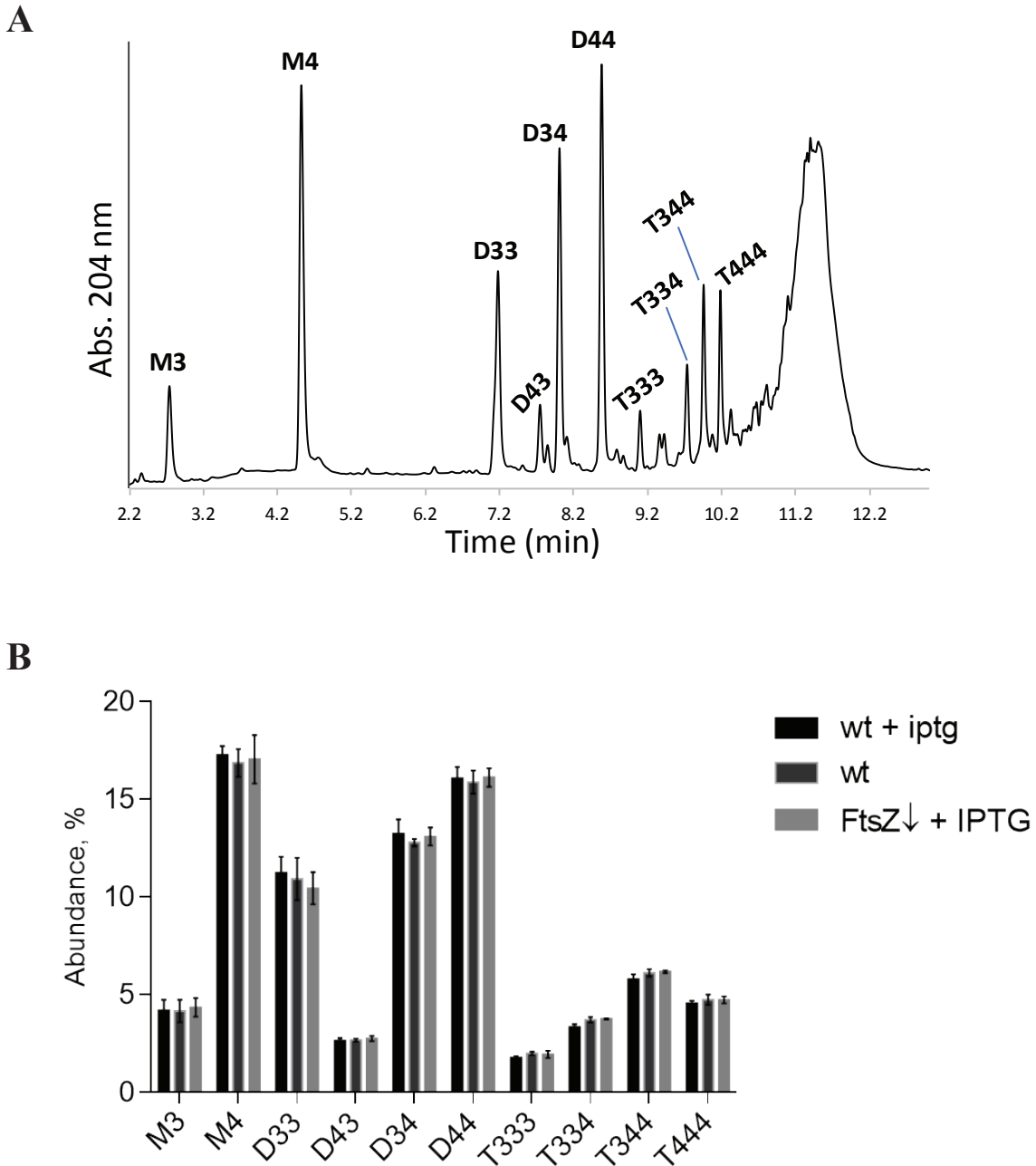


Figure 5- Figure Supplement 1

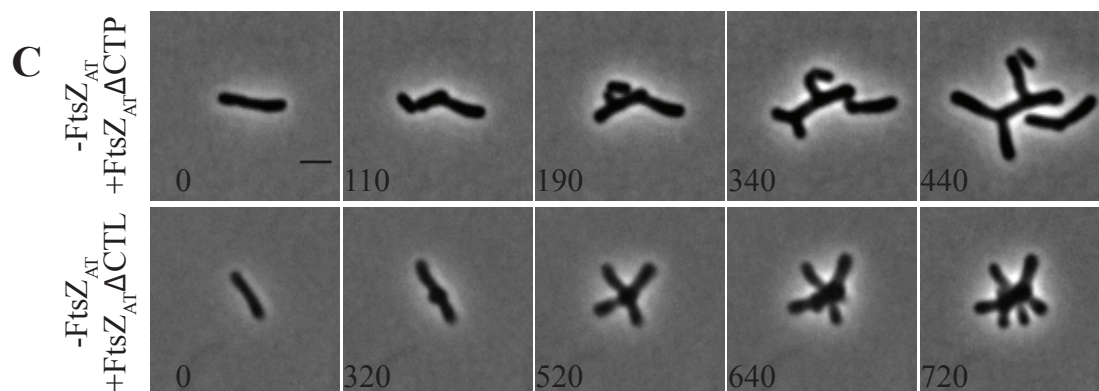
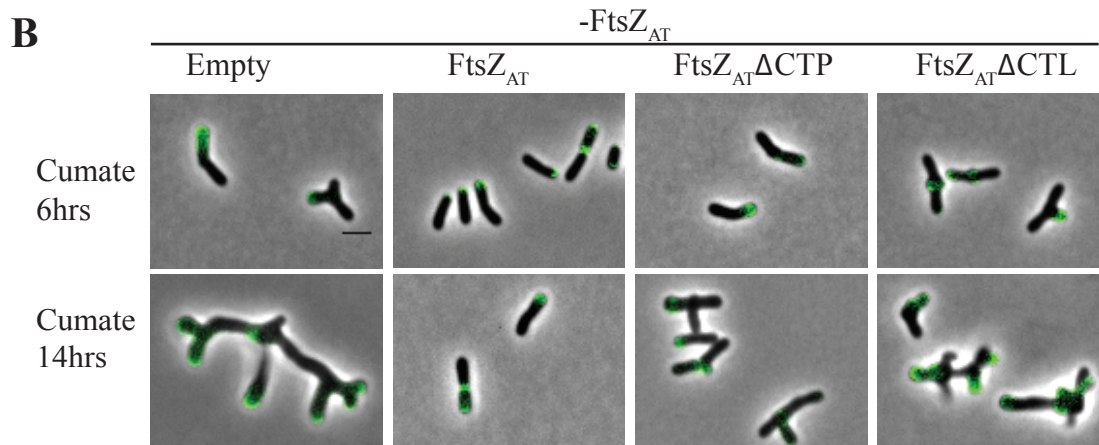
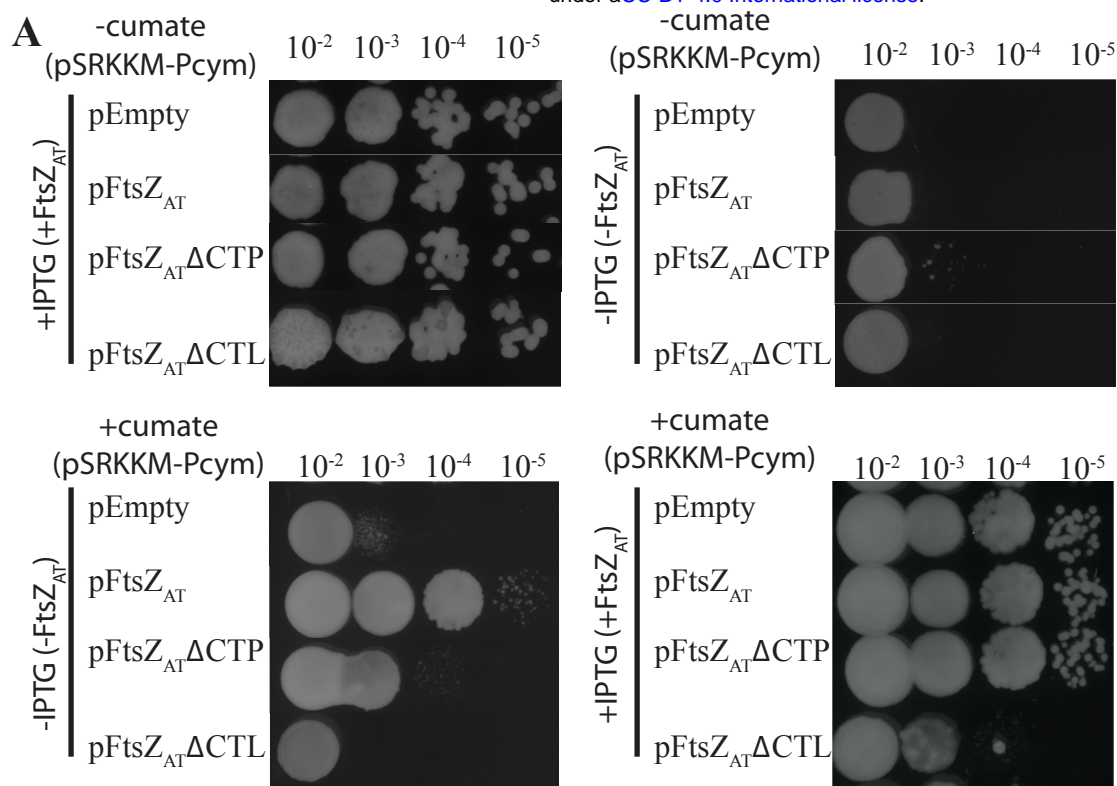


Figure 6

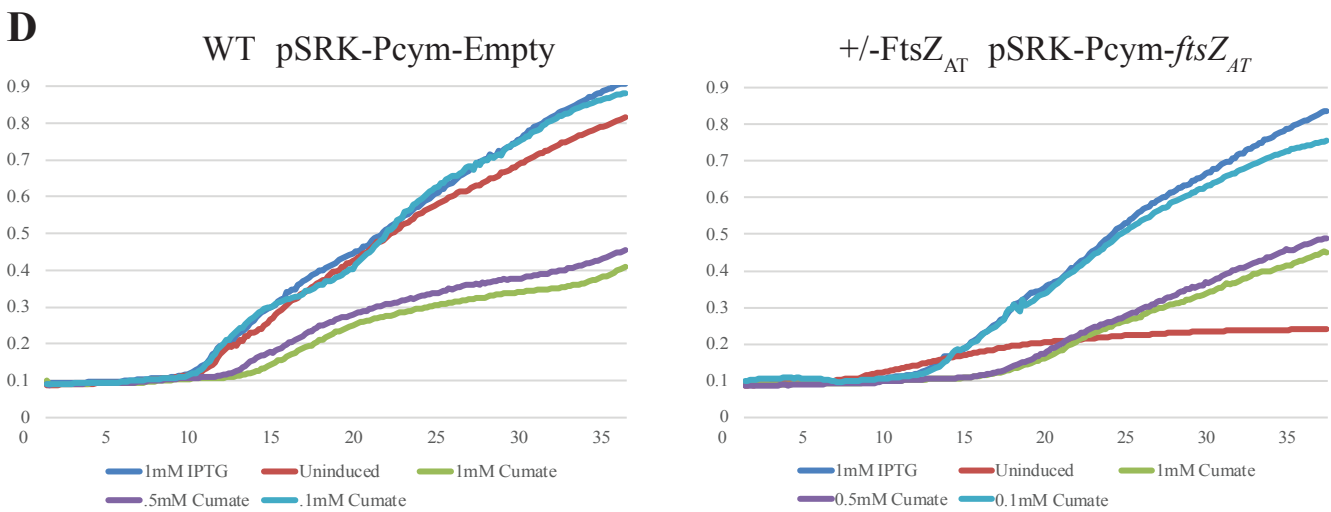
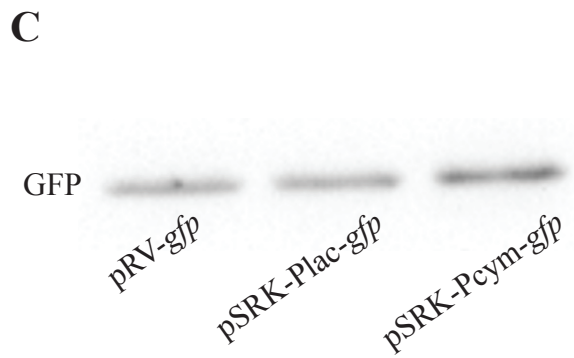
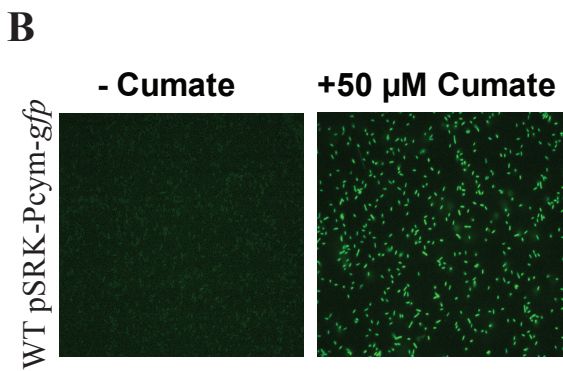
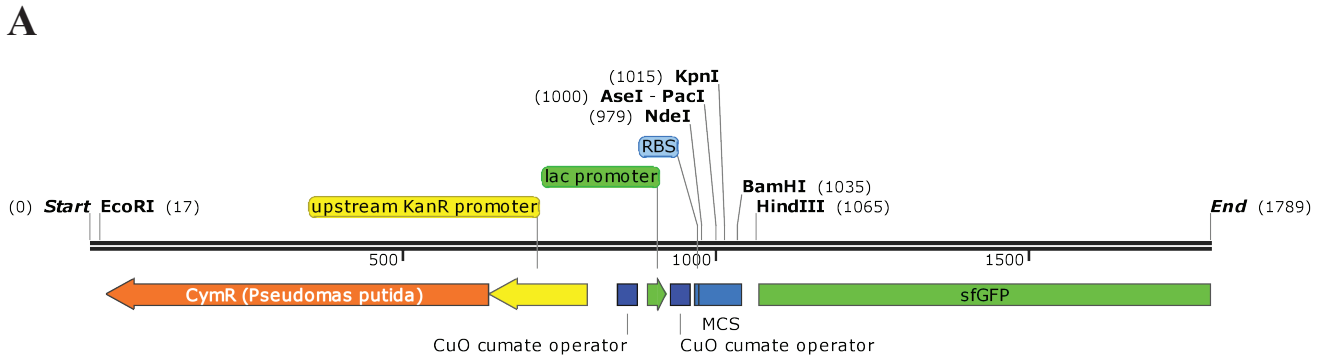


Figure 6- Figure Supplement 1

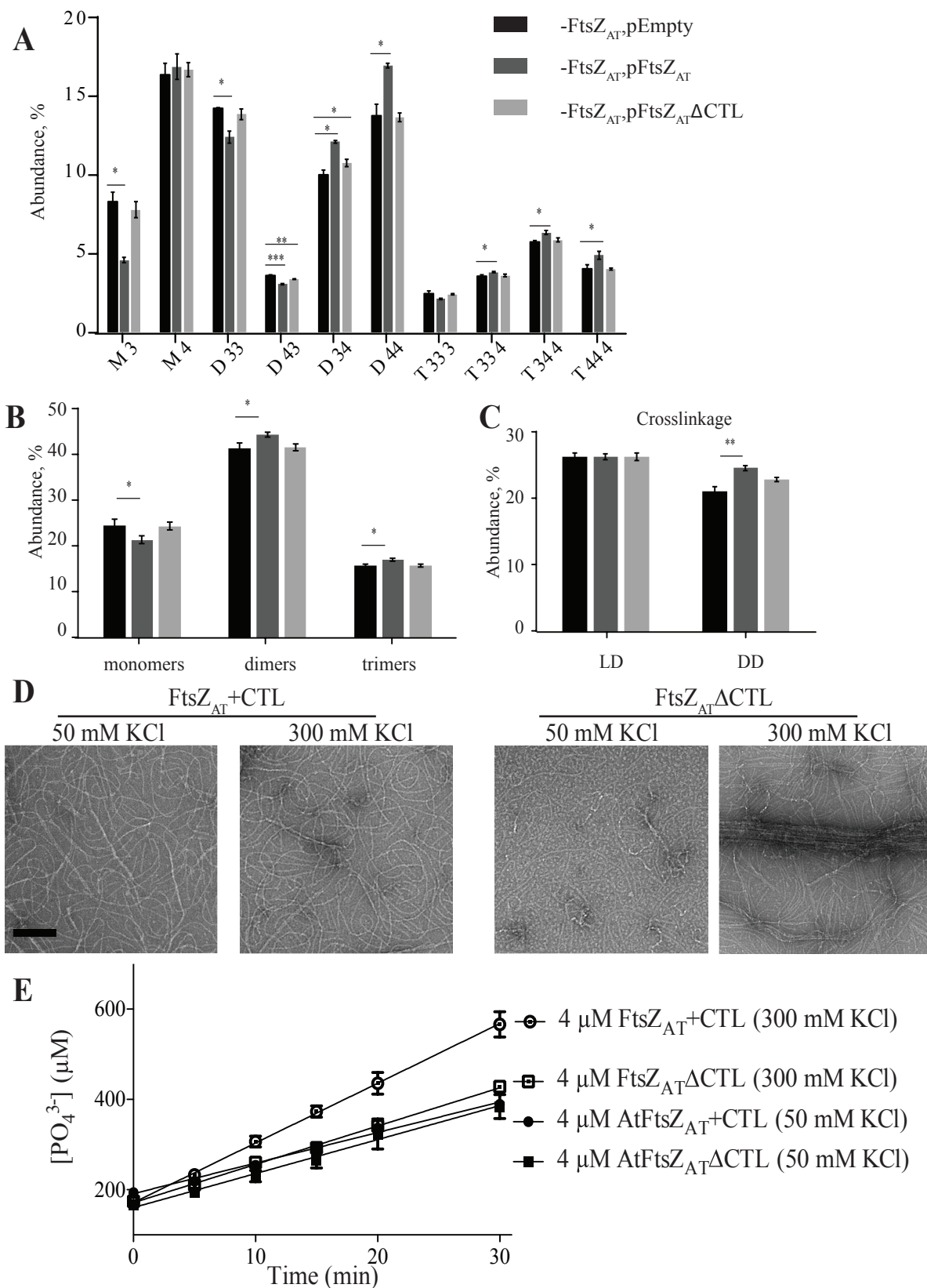


Figure 7

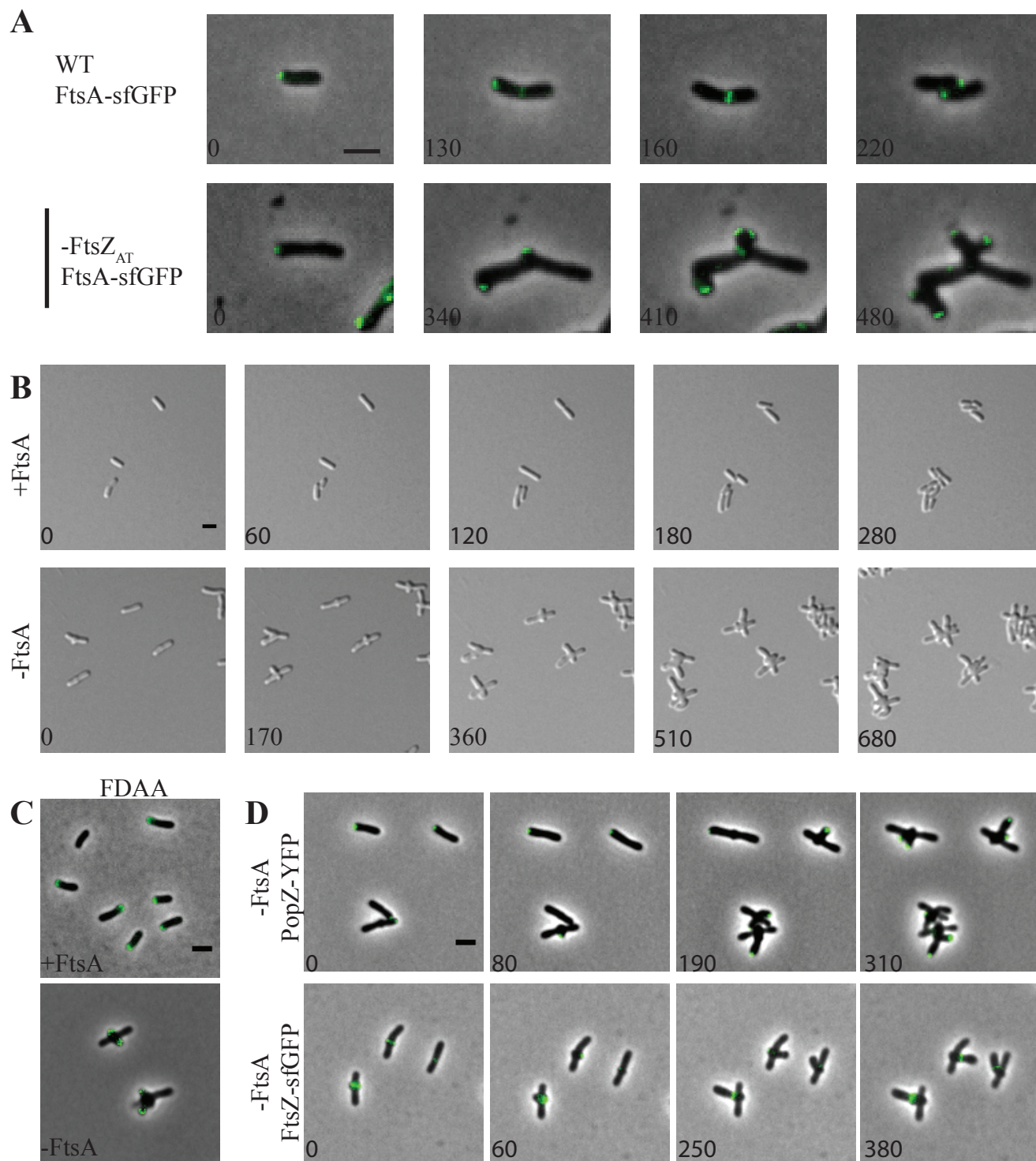


Figure 8

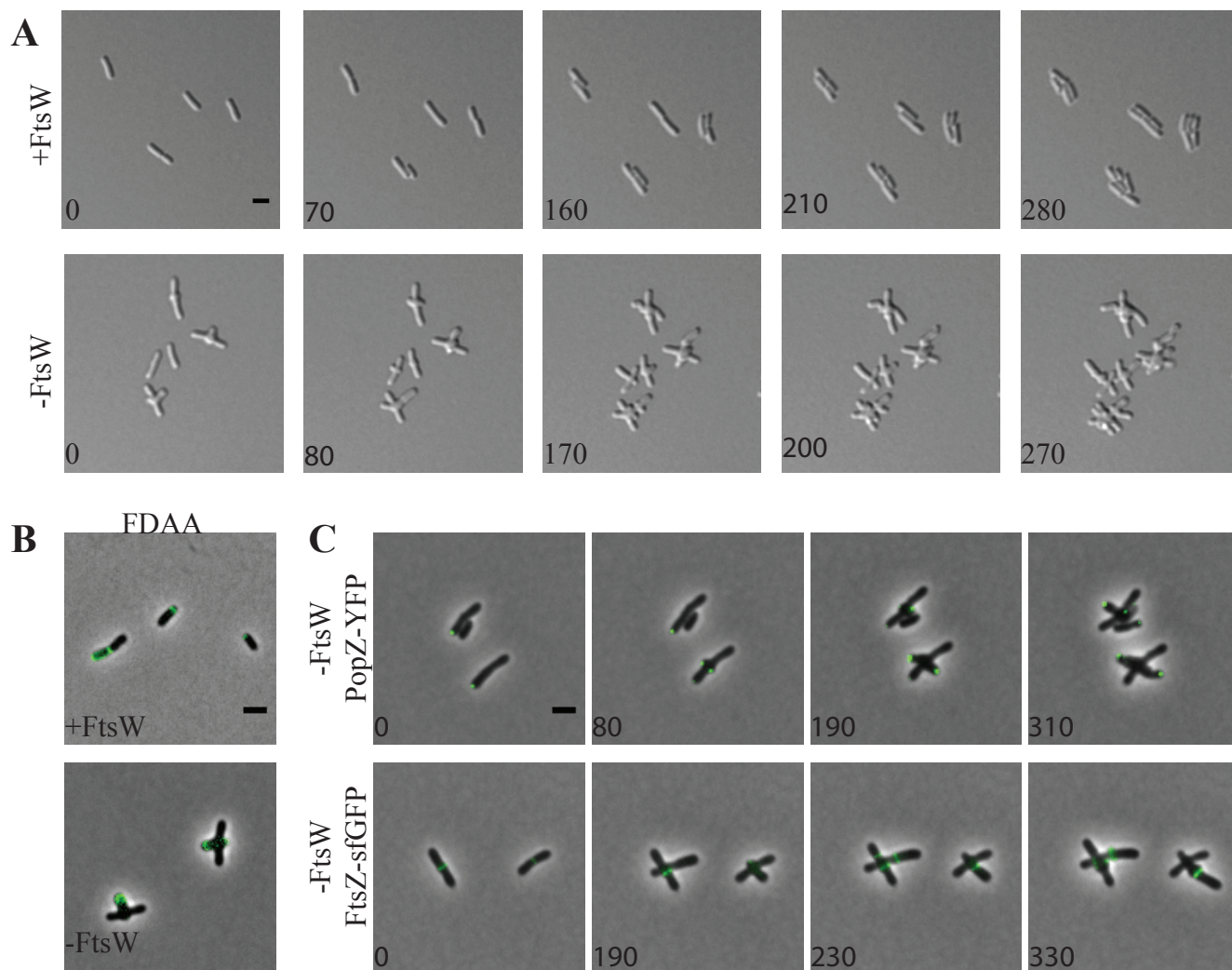


Figure 9