Unipolar peptidoglycan synthesis in the Rhizobiales requires an essential class A penicillin-binding protein

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18 ABSTRACT

19 Members of the Rhizobiales are polarly-growing bacteria that lack homologs of the 20 canonical rod complex. To investigate the mechanisms underlying polar cell wall 21 synthesis, we systematically probed the function of cell wall synthesis enzymes in the plant-pathogen Agrobacterium tumefaciens. The development of fluorescent D-amino 22 23 acid dipeptide (FDAAD) probes, which are incorporated into peptidoglycan by penicillinbinding proteins in *A. tumefaciens*, enabled us to monitor changes in growth patterns in 24 the mutants. Use of these fluorescent cell wall probes and peptidoglycan compositional 25 analysis convincingly demonstrate that a single class A penicillin-binding protein is 26 27 essential for polar peptidoglycan synthesis. Furthermore, we find evidence of an alternative mode of cell wall synthesis that likely requires LD-transpeptidase activity. 28 Genetic analysis and cell wall targeting antibiotics reveal that the mechanism of unipolar 29 growth is conserved in Sinorhizobium and Brucella. This work provides insights into 30 31 unipolar peptidoglycan biosynthesis employed by the Rhizobiales during cell elongation.

33 INTRODUCTION

Our current understanding of peptidoglycan (PG) assembly in rod-shaped bacteria 34 35 stems largely from investigations conducted using well-known model species like Escherichia coli and Bacillus subtilis, which incorporate new cell wall material along the 36 lateral sidewalls of the cell body. Expanding our studies of cell wall synthesis to include 37 38 diverse species, with alternative modes of elongation, is an important step in unveiling the mechanisms of how and why bacteria evolve novel growth modes and generate 39 innovative morphologies. It had, for example, long been assumed that all rod-shaped 40 bacteria employed the same growth strategy; however, unipolar growth is widespread 41 42 among rod-shaped bacteria in the Alphaproteobacterial order Rhizobiales, suggesting diversification of growth strategies [1]. The Rhizobiales are comprised of diverse 43 bacteria with respect to both their cellular morphology, and their environmental niches 44 [2, 3]. This includes many species of medical and agricultural significance such as the 45 46 facultative intracellular pathogens Bartonella and Brucella, the nitrogen-fixing plant symbiont Sinorhizobium, and the causative agent of crown gall disease Agrobacterium 47 tumefaciens, [4, 5]. PG labeling experiments in A. tumefaciens, Brucella abortus, and S. 48 49 *meliloti* have confirmed that unipolar growth is the mode of elongation utilized by these rod-shaped species [1]. Remarkably, labeling experiments have revealed that polar 50 growth in A. tumefaciens results specifically in the incorporation of pentapeptides at the 51 growth tip; [6] however, the mechanisms that underlie polar PG biosynthesis remain 52 poorly understood. 53

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PG biosynthesis is an essential process that allows bacteria to grow and divide, 55 faithfully reproducing their characteristic cell shape [7]. PG assembly requires different 56 classes of synthesis enzymes including the penicillin-binding proteins (PBPs), which 57 can be further divided into two classes. Class A PBPs are bifunctional enzymes that 58 catalyze β -1,4 linkages between the N-Acetylglucosamine (NAG) and N-Acetylmuramic 59 60 acid (NAM) sugars in a process called transplycosylation, and also synthesize crosslinks between 4-3 and 4-4 peptides in a process known as transpeptidation [8]. 61 The class B PBPs are monofunctional enzymes that have only transpeptidase (TPase) 62 activity [9]. The shape, elongation, division, sporulation (SEDS) family proteins, RodA 63 and FtsW, also possess glycosyltransferase (GTase) activity [10, 11]. Current models of 64 cell wall assembly maintain that SEDS proteins, in complex with their cognate class B 65 PBP, are the primary drivers of cell wall synthesis and are required to sustain rod shape 66 [12-14]. Thus, RodA functions with PBP2 during elongation, while FtsW functions with 67 68 PBP3 (encoded by *ftsl*) during cell division. The class A PBPs are currently thought to act independently from the rod complex, functioning primarily in PG maintenance and 69 repair [15, 16]. 70

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The suite of cell wall synthesis enzymes encoded by the Rhizobiales, is distinctly different from other bacterial orders. For example, the elongation-specific rod complex of proteins including PBP2, RodA, and MreBCD are universally absent [1, 17], suggesting that RodA-PBP2 are not the primary drivers of elongation. In addition, the genomes of Rhizobiales are enriched for the presence of LD-transpeptidases (LDTs). LDTs are a class of cell wall synthesizing enzymes that carry out transpeptidation

reactions to catalyze 3-4 and 3-3 crosslinks in the cell wall [18]. The cell wall of *A*. *tumefaciens* contains a high proportion (30%) of 3-3 and 3-4 crosslinks, in contrast to
laterally growing rod-shaped bacteria, where only 1-5% of the cell wall is crosslinked by
LDTs [19]. This suggests that LDT enzymes may play an important role during polar
growth [1, 20]. Overall, these observations suggest that Rhizobiales use a noncanonical mechanism for polar elongation.

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85 Using a combination of microscopy, new probes, biochemical, and genetic analyses we have characterized the function of the six high molecular weight PBPs encoded by A. 86 tumefaciens and have identified the major cell wall synthesis enzymes required for polar 87 growth. We show that, unlike the proposed auxiliary function of PBP1a in other rod-88 shaped bacteria, in A. tumefaciens PBP1a is an essential enzyme required for polar PG 89 expansion, with depletion of PBP1a resulting in a loss of proper rod-shape. Using newly 90 developed fluorescent D-amino acid dipeptide (FDAAD) probes, we show that PBP1a is 91 92 the enzyme primarily responsible for inserting nascent PG at the pole. Additionally, 93 PBP1a depletion leads to a modified PG composition, including an increase in LDT linkages. Collectively, this suggests that the mechanism of polar growth in the 94 Rhizobiales has evolved through the expansion, diversification, and altered regulation of 95 96 the core cell wall synthesis machinery. We confirmed the essentiality of PBP1a in the closely related bacterium Sinorhizobium, suggesting that the mechanisms underlying 97 polar growth in the Rhizobiales are well conserved. Finally, we have identified the β -98 99 lactam faropenem as a specific inhibitor of polar growth in Agrobacterium, 100 Sinorhizobium, and Brucella, indicating that the target(s) of faropenem are conserved

among the Rhizobiales. These findings broaden our understanding of the role of PG
synthesis enzymes that contribute to polar growth and will inform strategies aimed at
developing novel therapeutics that target the cell wall of polar-growing bacteria in the
Rhizobiales [21, 22].

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106 **RESULTS**

107 The class A cell wall synthase PBP1a is essential for polar growth

108 To begin probing the molecular determinant(s) of polar growth in *A. tumefaciens*, we

sought to generate deletions of the predicted PG synthase enzymes. Although A.

110 *tumefaciens* lacks homologs to the predicted cell elongation synthases RodA and

111 PBP2, the genome encodes four bifunctional class A PBPs (PBP1a, PBP1b1, PBP1b2

and PBP1c), two monofunctional class B PBPs (PBP3a and PBP3b), and one

113 monofunctional GTase, MtgA (Figures 1A, 2A). To determine which of these enzyme(s)

provide the primary GTase activity in the absence of a RodA homolog, we made in-

115 frame deletions of those genes encoding predicted GTase enzymes to further explore

their contribution during cell growth or division.

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118 Consistent with a previous transposon screen [26] it was not possible to obtain a PBP1a 119 deletion. We therefore constructed a PBP1a depletion strain by introducing a copy of 120 the PBP1a-encoding gene, under an IPTG-inducible promoter, at a heterologous site on 121 the chromosome and subsequently succeeded in creating an in-frame deletion of the 122 native gene encoding PBP1a in the presence of the IPTG inducer [27]. The PBP1a

depletion strain grown in the presence of IPTG is referred to as + PBP1a and the 123 depletion strain grown in the absence of IPTG is referred to as - PBP1a. We confirmed 124 depletion of PBP1a in the absence of IPTG using Bocillin-FL, a fluorescent penicillin 125 derivative. Two bands were observed that could correspond to the predicted molecular 126 weight of PBP1a (~88 kDa), but only the second band was absent in PBP1a-depleted 127 128 cells and likely represents PBP1a protein (Supplementary Figure 1A). Strikingly, cells depleted of PBP1a for 16 hours lost their rod shape, becoming shorter (Figure 1A, B) 129 and wider (Supplementary Figure 1B) and had a severe viability defect, as measured by 130 spotting serial dilutions, compared to the same strain when PBP1a is induced 131 (Supplementary Figure 1C). The addition of IPTG to wild-type A. tumefaciens led to a 132 slight increase in the median cell length (Figure 1B, 1C) compared to wildtype alone but 133 had no effect on cell viability or rod shape (Supplementary Figure 1B, 1C). 134

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Of the remaining bifunctional PBPs, single deletions of the genes encoding PBP1b1, 136 PBP1b2, and PBP1c had no effect on cell length (Figure 1B, C) or cell viability 137 138 (Supplementary Figure 1D). Similarly, a double mutant of PBP1b1 and PBP1b2 or a triple mutant of PBP1b1, PBP1b2 and PBP1c (referred to as ∆3pbp) had no obvious 139 140 mutant phenotype with respect to cell length (Figure 1B, C) or cell viability (Supplementary Figure 1D). Thus, despite all of the predicted bifunctional PBPs being 141 produced during exponential growth of A. tumefaciens (Supplementary Figure 1E), our 142 data indicate that only PBP1a makes a major contribution to cell growth under standard 143 laboratory conditions. Similarly, deletion of the monofunctional GTase encoding mtgA 144 produced cells of normal length (Figure 1B, C). The lack of a readily observed 145

- phenotype in the $\Delta mtgA$ strain is consistent with findings in *E. coli* and *Hyphomonas*
- *neptunium* [28-30]. Together, these data suggest that the bifunctional enzyme PBP1a,
- 148 which likely has both GTase and TPase activities, fulfils the role of RodA and PBP2, as
- the primary PG synthase required for polar elongation in *A. tumefaciens*.



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Figure 1. Functional characterization of PG synthases in A. tumefaciens. (A) Domain structure of the 152 putative PG biosynthesis enzymes showing the transmembrane (orange), glycosyltransferase (GT, 153 PF00912), transpeptidase (TP, PF00905), OB-like (PF17092), and biPBP_C (PF06832) domains. The 154 155 regions of intrinsic disorder (grey) as predicted by MoBiDB are also shown [23]. The scale indicates the length in amino acids (aa). The corresponding ATU numbers are listed in parentheses beside each gene 156 157 name. (B) Phase microscopy images showing the phenotypes of PG synthase mutants. Each strain was 158 grown to exponential phase, spotted on an ATGN agar pad (ATGN is a minimal medium with glucose and 159 (NH4)2SO4), and imaged by phase microscopy. Scale bar: 2 µm. (C) Cell length distributions of PG 160 synthase mutants. The indicated strains were grown as in B and subjected to cell lengths measurements using MicrobeJ [24]. The data are represented as box and whisker plots in the style of Tukey [25], and 161 visualizes five summary statistics (the center line is the median, the two hinges correspond to the first 162 and third guartiles (the 25th and 75th percentiles), the two whiskers (representing the smallest and 163 largest values no further than 1.5 times the interguartile range), and all "outlying" points are plotted 164 165 individually. The PBP1a depletion strain grown in the presence of IPTG is referred to as + PBP1a and the 166 depletion strain grown in the absence of IPTG is referred to as - PBP1a. Distributions of cells significantly different from wildtype (WT) are indicated (***; One-Way ANOVA with Bonferroni correction, p > 2*10^16). 167 168 n = > 800 cells per strain.

169 Class B synthases PBP3a and PBP3b are required for cell division

Incorporation of PG at the septum prior to cell division typically requires synthesis 170 171 enzymes that are distinct from the cell elongation machinery. Although A. tumefaciens 172 lacks the cognate SEDS-PBP pair that is typically required for elongation, the SEDS protein FtsW and PBP3, which are required for cell division, are conserved. While most 173 174 bacteria possess a single, essential *ftsl* gene that encodes PBP3, some Rhizobiales, including A. tumefaciens, encode two Ftsl homologs (PBP3a and PBP3b) (Figure 2A). 175 pbp3a resides in the mra operon of cell division and cell envelope biogenesis genes 176 similar to most *ftsl*-encoding homologs [31], while PBP3b is encoded as a monocistronic 177 gene elsewhere in the genome. This raises the possibility that the second 178 monofunctional transpeptidase (PBP3b) may serve the role of a PBP2 homolog that 179 functions in polar elongation. Thus, we sought to determine if either of the two class B 180 PBP homologs were required for cell division or polar elongation in A. tumefaciens. 181 182 Saturating transposon mutagenesis in LB medium indicated that PBP3a was likely essential [26]; however, it was possible to make a deletion of *pbp3a* when cells were 183 grown in minimal medium. These observations suggest that the PBP3a-encoding gene 184 was conditionally essential. 185

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In minimal medium, deletion of *pbp3a* caused a severe cell viability defect
(Supplementary Figure 1F). In addition, cells were longer (Figure 2B, C), and were
frequently observed to adopt branching or bulging morphologies; these features are a
hallmark of cell division defects in *A. tumefaciens* [32, 33]. In contrast, deleting *pbp3b*had no effect on cell viability (Supplementary Figure 1F) or cell length (Figure 2B, C).

Together, these results indicate that PBP3a is the major class B PBP contributing to cell 192 division in *A. tumefaciens*. Since the deletion of the gene encoding PBP3a did not fully 193 inhibit cell division, we hypothesized that PBP3b may be able to partially compensate 194 for the loss of PBP3a. To address this possibility, we first created a PBP3a depletion 195 strain; when this strain was grown in the absence of IPTG for 24 hours, it phenocopied 196 197 the cell viability and cell length defects of the *pbp3a* deletion mutant (Figure 2B, C). We then depleted PBP3a in a $\Delta pbp3b$ mutant background, and found that cells not only 198 failed to divide, but also swelled at the mid-cell before lysing (Figure 2C, Supplemental 199 200 movie 1), indicating that PBP3a and PBP3b both contribute to septal PG biosynthesis during cell division. The phenotype observed in the absence of both PBP3a and PBP3b 201 was remarkably similar to the phenotype observed during depletion of FtsW in A. 202 tumefaciens [32]. FtsW possesses GTase activity, and is a major synthase required for 203 cell division [12, 34]. Consistent with current models of cell division, we hypothesize that 204 205 FtsW provides the GTase activity while PBP3a and PBP3b provide the DDtranspeptidase (TPase) activity necessary for proper septal PG biosynthesis during cell 206 division in A. tumefaciens. In all, our findings support a model in which PBP3a can 207 208 sustain proper cell division in the absence of PBP3b, and that, while PBP3b contributes to septal PG synthesis, it cannot fully compensate for the loss of PBP3a. Thus, both 209 210 class B PBPs contribute primarily to septal PG biosynthesis.



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Figure 2. Functional characterization of monofunctional synthases PBP3a and PBP3b. (A) Domain 212 structure of the putative PG biosynthesis enzymes showing the transmembrane (orange), transpeptidase 213 214 (TP, PF00905), and PBP dimer (PF03717) domain. The scale indicates the length in amino acids (aa). 215 The corresponding ATU numbers are listed in parentheses beside each gene name. (B) Phase 216 microscopy images showing the phenotypes of PG synthase mutants. Each strain was grown to exponential phase, spotted on an ATGN agar pad, and imaged by phase microscopy. Scale bar: 2 µm. 217 (C) Cell length distributions of PG synthase mutants. The indicated strains were grown to exponential 218 219 phase, spotted on an agar pad, imaged by phase microscopy, and subjected to cell lengths 220 measurements using MicrobeJ [24]. The data are shown as box and whisker plots in the style of Tukey 221 [25]. Distributions of cells significantly different from wildtype (WT) are indicated (***; One-Way ANOVA 222 with Bonferroni correction, p< $2*10^{16}$). n = > 500 per strain. 223

224 Development of new fluorescent cell wall probes to monitor PBP activity

225 Traditional fluorescent D-amino acid (FDAA) probes are an exceptionally useful tool for 226 investigating the patterning of cell wall synthesis in diverse microbes [35]. However, 227 FDAAs report on the activity of extracellular/periplasmic DD and LD transpeptidases, and as a result can be incorporated into the PG in a growth-independent mechanism [6]. 228 229 Here, we have developed fluorescent D-amino acid dipeptide (FDAADs) probes to observe nascent sites of PG crosslinking in living cells, thus eliminating the need for 230 click chemistry that is required when using traditional D-amino acid dipeptide probes 231 (DAADs) [6, 36]. DAADs are incorporated into the cell wall precursors by the 232 233 cytoplasmic MurF ligase and probe incorporation reports specifically on nascent PG synthesis [36-38]. The resulting lipid II-linked modified precursor is most likely covalently 234 crosslinked to an existing glycan strand through the activity of bifunctional PBPs (Figure 235 3A). The FDAAD probe HADA—DA successfully labeled the PG of several bacteria. 236 237 including B. subtilis, E. coli, Streptomyces venezuelae and A. tumefaciens (Figure 3B, C), and we demonstrated and evaluated PG labeling with four additional FDAADs of 238 different sizes and molecular weights in diverse species (Supplemental figure 2A-D). In 239 240 addition, S. venezuelae, a polar growing Actinobacteria, was short pulsed with three different FDAADs to illustrate that these probes report on the newest PG synthesis 241 242 activity (Figure 3B), similar to FDAAs [39, 40]. Given their cytoplasmic mechanism of incorporation, complementary to FDAAs (which are incorporated by transpeptidases 243 periplasmically) [6], these probes will be particularly useful to distinguish between 244 growth-dependent and growth-independent PG crosslinking in species with a higher 245 proportion of extracellular LDT activity, such as polar-growing bacterial species. 246





248 Figure 3. Fluorescent D-amino acid dipeptide (FDAAD) labeling is absent from the growth pole in 249 the PBP1 depletion. (A) Schematic of the incorporation pathway of fluorescent D-amino acid dipeptides 250 (FDAADs), which are incorporated into the muropeptide precursor molecule in the cytoplasm by MurF ligase. The modified muropeptide precursor is flipped across the membrane and the activity of a 251 252 bifunctional PBP crosslinks the new PG monomer into the existing PG sacculus. (B) Top row: structure of 253 the FDAAD HADA—DA and merged phase and fluorescent channels of labeling patterns of HADA—DA 254 in E.coli and B. subtilis. Bottom row: short pulse labeling of S. venezuelae sequentially labeled first with BADA—DA, followed by HADA—DA and Atto610DA—DA. (C) Demographs of wild-type (WT) cells depict 255 incorporation of FDAADs at a population level. Median profiles of the fluorescence channel are stacked 256 257 and ordered by cell length n = 513. (D) Demographs depict incorporation of FDAAD in the PBP1a 258 depletion strain grown in the presence or absence of IPTG labeled + PBP1a and - PBP1a, respectively. 259 Merged phase and fluorescent channels of cells with representative polar and septum labeling of FDAAD are shown to the right of each demograph. Scale bars: $2 \mu m$. n = > 1000 per strain 260

262 **PBP1a is the major synthase incorporating PG at the pole**

A. *tumefaciens* was labeled for ~5% of the cell cycle with HADA—DA. As expected, cells exhibit labeling at the growth pole during elongation, and at the septum in cells undergoing cell division (Figure 3C). This pole-to-septum labeling pattern is consistent with other cell wall labeling methods including fluorescent D-amino acids (FDAAs) [41] and D-cysteine labeling [1]. Similar to wild-type cells, a pole-to-septum labeling pattern was observed in the Δ 3pbp mutant, consistent with a limited role for these class A PBPs in polar growth under the conditions tested (Supplementary Figure 3).

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271 In stark contrast to wild-type (Figure 3C) or PBP1a replete cells (Figure 3D), pole-272 specific labeling with HADA—DA was almost completely absent from cells depleted of PBP1a for 20 hours (Figure 3D). Similar results were observed with NADA—DA and 273 BADA—DA (Supplemental Figure 4A, B). Consistent with these observations, we 274 tracked the growth of PBP1a-depleted cells for seven generations using a microfluidic 275 276 device and found that the reduction in cell length occurred first for the new pole 277 daughter cell and likely resulted from the loss of polar PG insertion by PBP1a (Supplemental Figure 3A, Supplemental movie 2). We thus concluded that PBP1a is the 278 major PG synthase required for polar PG incorporation. Notably, PBP1a-depleted cells 279 280 have more robust FDAAD labeling at the septum (Figure 3D), suggesting that additional 281 glycosyltransferase enzymes remain active at the site of cell division.

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LD-transpeptidases contribute to growth-independent and growth-dependent PG modification

285 In contrast to FDAAD incorporation, which is linked to the PG precursors in the cytoplasm, the more conventional fluorescent D-amino acid (FDAA) labeling occurs in 286 the periplasm through either DD-transpeptidase reactions carried out by PBPs or LD-287 transpeptidase reactions carried out by LDTs [41, 42]. As expected, wild-type and 288 Δ 3pbp cells labeled with a short pulse of the FDAA probe HADA show the characteristic 289 pole-to-septum labeling pattern that is typical of A. tumefaciens growth (Figure 4A. 290 Supplementary Figure 3). Notably, FDAADs label discrete regions at the pole and 291 292 midcell, while FDAA labeling is more prominent in the sidewalls, particularly of the new pole compartment prior to cell division, while the old pole remains unlabeled. (Figure 4A 293 white arrow). This labeling pattern is consistent with the reported growth-independent 294 incorporation of FDAAs by LDTs [6]. Interestingly, we observed that a majority of 295 growth-independent labeling occurred preferentially in the new pole compartment. To 296 further support this idea, we labeled A. tumefaciens for 60 minutes with either HADA or 297 HADA—DA and compared the labelling pattens (Supplemental figure 5). After a 60-298 minute incubation, HADA—DA labeling was primarily found in distinct regions at the 299 300 pole and midcell, with little sidewall labeling, similar to a short pulse and consistent with areas of new synthesis. In contrast, the HADA labelling was much brighter, and fully 301 labeled the sidewalls. HADA labeling was particularly enriched along the sidewalls of 302 the new-pole compartment. Our observations during short and long-pulse labeling 303 experiments suggests that, in addition to growth-dependent pole and midcell labeling, 304

LDTs contribute to spatially distinct crosslinking of the cell wall, particularly along the sidewalls of the new pole daughter cell.

308	We next sought to test whether FDAA labeling at the growth pole was absent in the
309	PBP1a depletion strain. In contrast to the absence of polar incorporation that was seen
310	following FDAAD labeling, cells depleted of PBP1a labeled robustly at the growth pole
311	with HADA (Figure 4A). Since PBP-mediated incorporation of FDAADs is absent from
312	the growth pole in the PBP1a depletion (Figure 3D, Supplemental Figure 4A, B), the
313	major enzymes incorporating HADA at the pole are most likely LDTs, consistent with
314	recent findings for incorporation of FDAAs in E. coli [42]. Additionally, the activity of
315	LDTs has been shown to be functionally linked to the activity of class A PBPs [42, 43].
316	Therefore, our findings indicate that LD-transpeptidation contributes to PG crosslinking
317	at the growth pole.



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319 Figure 4. Fluorescent D-amino acid (FDAA) and PG composition analysis illustrates a role for 320 LDTs in polar growth. (A) Demographs of wildtype (WT) and the PBP1a depletion strain grown in the 321 presence or absence of IPTG labeled + PBP1a and - PBP1a, respectively. Demographs depict 322 incorporation of FDAA at a population level. Median profiles of the fluorescence channel are stacked and ordered by cell length wildtype n = > 600 per strain. Merged phase and fluorescent channels of cells with 323 representative polar and septum labeling by FDAAs are shown below each demograph. Scale bar: 2 µm. 324 325 (B) Bar graphs depicting the average abundance of muropeptides obtained by UPLC analysis from 326 wildtype, and the PBP1a depletion strain grown in the presence or absence of IPTG for 16 hours. Major 327 muropeptides are labeled M, monomers; D, dimers; and T, trimers. Numbers indicate the length of the 328 muropeptide stems and the position of cross-links in dimers and trimers. Data shown are averages taken 329 from analysis of three independent biological samples. Samples that are statistically significant are indicated (One-Way ANOVA with Tukey's multiple comparison test, ** p<0.005, *** p<0.0005). p-value 330 331 between + PBP1a and - PBP1a for D34 was 0.0588 and between WT and - PBP1a for D44 was 0.072.

333 Depletion of PBP1a or loss of PBP1b2 leads to increased levels of LD-crosslinks 334 in PG

335 Since depletion of PBP1a led to dramatic reduction in the polar incorporation of 336 FDAADs, while maintaining polar FDAA incorporation we hypothesized that a decrease in the activity of PBPs relative to LDTs may lead to altered PG composition. We grew 337 338 the PBP1a depletion strain in the presence or absence of IPTG for 16 hours, collected the cell wall fraction, and analyzed muropeptides by ultra-performance liquid 339 chromatography (UPLC). The major muropeptides found in wild-type A. tumefaciens 340 PG, included monomeric (M), dimeric (D) and trimeric (T) muropeptides (Figure 4B). 341 PG from PBP1a-depleted cells had significantly reduced levels of muropeptides with DD-342 crosslinks, as seen by the ~3% reduction in D44 abundance (Figure 4B). As expected, 343 this observation confirmed that depleting PBP1a leads to decreased DD-transpeptidase 344 activity. Depletion of PBP1a also resulted in an increase in muropeptides containing LD-345 crosslinks, as indicated by the ~4% increase in D34 abundance (Figure 4B). A similar 346 decrease in D44 abundance and increase in D34 abundance was also observed in the 347 348 $\Delta pbp1b2$ (Supplemental Figure 4A), but not in the single deletions of pbp1b1, pbp1c, pbp3a or pbp3b (Supplemental Figure 4B). Therefore, the compositional changes in the 349 double and triple mutant strain could likely be attributed to the deletion of *pbp1b2*. 350 351 These results implicate PBP1b2 as an important DD-transpeptidase enzyme in A. 352 tumefaciens and suggest that increased LDT activity may be a general response to decreased DD-transpeptidase levels, and not necessarily a specific response to 353 depletion of PBP1a. Furthermore, since we saw an increase in muropeptide D34 but not 354 D33 (Figure 4B), we hypothesize that decreased DD-transpeptidase levels activate only 355

a subset of LDTs, and that another group of LDTs may function along with PBP1a
during polar growth, consistent with the observation of HADA labeling at the growth
pole.

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360 Faropenem treatment inhibits polar growth in the Rhizobiales

361 Since species in the Rhizobiales have a large number of LD-transpeptidase enzymes (A. *tumefaceins* encodes 14 LDTs), deleting them all is a significant undertaking. β-lactam 362 antibiotics are one of the most widely used classes of antibiotics that target cell wall 363 synthesis enzymes, and these have been primarily studied for their ability to target 364 PBPs [44]. A subclass of β -lactam antibiotics known as the carbapenems, including the 365 penem antibiotic faropenem, are, however, useful for probing the activity of LDTs [45, 366 46]. Therefore, to characterize the global contribution of LDT activity during A. 367 tumefaciens growth, we investigated the effect of five carbapenem antibiotics and one 368 369 penem antibiotic on cell growth and morphology. In A. tumefaciens, we find that treatment with meropenem or faropenem leads to an overall decrease in PG 370 crosslinkage, including both LD- and DD-crosslinks (Figure 5A, Supplemental Figure 5), 371 372 suggesting that the targets of these drugs impact cell wall biosynthesis.



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374 Figure 5. Phenotypic characterization of carbapenem and penem antibiotic treatments. (A) 375 Abundance of total LD- and DD-crosslinkages in PG isolated from wild-type cells, wild-type cells grown in 376 the presence of 1.5 µg/mL meropenem for 4 hours, and wild-type cells grown in the presence of 1.5 377 µg/mL of faropenem for 6 hours. Timepoints were chosen based on the onset of phenotypic changes. 378 Data shown are the total abundance of the muropeptides containing LD- or DD-crosslinks from analysis of 379 two independent samples. (B) Representative images of wild-type cells grown in the presence of 1.5 µg/mL meropenem and faropenem. Cells were incubated with antibiotics for 24 hours, then spotted on a 380 381 1% agarose pad and imaged using DIC microscopy. Scale bar: 2 µm (C) Time-lapse microscopy of wild-382 type cells spotted on a 1% ATGN agarose pad supplemented with 1.5 µg/mL of faropenem; images were 383 acquired every 10 minutes. Indicated time in hours is shown. Scale bar: 2 um (D) Representative phase microscopy images of A. tumefaciens, S. meliloti, or B. abortus grown overnight in ATGN, Tryptone-Yeast 384 (TY) Extract or Brucella Broth, respectively, to an OD₆₀₀ of 0.6 and spotted on a 1% ATGN agarose pad 385 386 with or without 1.5 µg/mL faropenem. Cells were imaged after 16 hours of growth. Scale bar: 2 µm.

To better understand the impacts of these antibiotics, we observed morphological 388 changes induced by drug treatment. Treatment with sub-minimum inhibitory 389 390 concentrations (MIC) treatments with any of five carbapenem antibiotics: meropenem, imipenem, doripenem, ertapenem, or tebipenem for 24 hours induced mid-cell swelling 391 (Figure 5B, Supplementary Figure 6). These data indicated that these carbapenem 392 393 antibiotics may target an enzyme(s) with a specific role at the septum during cell division. Interestingly, faropenem-treated cells became wider and rounder after 24-hour 394 exposure (Figure 5A), which pointed to the cellular target of faropenem as being 395 396 important for the maintenance of rod shape during polar growth. In agreement, time-397 lapse microscopy of faropenem-treated cells revealed a loss of rod shape preferentially in the daughter compartment. Remarkably, following cell division the daughter cell 398 which inherits the growth pole is large and round whereas the daughter cell inheriting 399 the old growth pole retains its rod shape (Figure 5B, Supplemental movie 3). This 400 401 phenotype was distinct from that associated with the treatment of the other carbapenem antibiotics and indicated that the cellular target(s) of faropenem was important to 402 maintain proper PG integrity in the growth pole compartment. 403

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The phenotype of faropenem-treated cells, along with the growth-independent labelling of HADA along the sidewalls of the new pole (Supplemental figure 6) suggested that the old and new cell compartments may have a distinct repertoire of PG synthesis enzymes that help to confer polar identity between the daughter cells. To determine if the cellular target(s) of faropenem were conserved in other Rhizobiales we treated the closely related plant symbiont *S. meliloti* and the obligate intracellular pathogen *B. abortus* with

sub-lethal concentrations of faropenem. We observed swelling of the growth pole in *A. tumefaciens, S. meliloti,* and *B. abortus* (Figure 5D). Thus, we have identified the β lactam antibiotic faropenem as a specific antibiotic inhibitor of polar growth among the Rhizobiales. Altogether, these data suggest that cell wall enzymes that are important for polar growth have a conserved role in agriculturally and medially important species of Rhizobiales.

417

418 The essentiality of PBP1a is conserved among the Rhizobiales

Since faropenem targeting of the growth pole machinery is conserved, we sought to 419 determine if the essential role of PBP1a is also conserved in other Rhizobiales. We found 420 that treatment of *B. abortus* with the PBP1a-specific GTase inhibitor flavomycin 421 (moenomycin) [47] led to growth arrest and the formation of large, round cells (Figure 422 6B). Consistent with our findings that treatment of *B. abortus* with the PBP1a inhibitor 423 flavomycin causes cells to lose their rod-shape, a transposon mutagenesis screen of B. 424 abortus predicted that out of the bifunctional PBPs, only PBP1a may be essential for 425 426 growth [48]. In line with this, Bandara and colleagues were unable to obtain a PBP1a mutant in Brucella melitensis, indicating that PBP1a is likely essential for viability [49]. 427



428

429 Figure 6. Mechanisms of polar growth are conserved in the Rhizobiales *Brucella* and

430 Sinorhizobium.(A) Representative phase microscopy images of B. abortus grown overnight without antibiotic, with 20 µg/mL of flavomycin, or with 1.5 µg/mL of faropenem, after 16 hours cells were spotted 431 432 on a 1% Brucella Broth agarose pad and imaged. (B) Top panel: phase microscopy images showing the 433 phenotypes of S. meliloti wildtype (WT) and PBP mutants. Each strain was grown to exponential phase, 434 spotted on a 1% TY agarose pad, and imaged by phase microscopy. Bottom panel: cell length 435 distributions of PG synthase mutants. The indicated strains were grown to exponential phase, spotted on 436 an agarose pad, imaged by phase microscopy, and subjected to cell lengths measurements using 437 MicrobeJ [24]. The data are shown as box plots in the style of Tukey [25]. Distributions of cells

438 significantly different from wildtype are indicated (***; One-Way ANOVA with Bonferroni correction, p<

439 2*10^16). n = >400 per strain.

Next, we explored the function of PBPs in the closely related plant symbiont S. meliloti. 441 S. meliloti encodes six bifunctional PBP homologs, including one PBP1a homolog, four 442 PBP1b homologs, and one PBP1c homolog. We constructed a Δ 5pbp mutant, which is 443 lacking all four PBP1b homologs and the PBP1c homolog. The ∆5pbp mutant had a 444 median cell length similar to that of wild-type S. meliloti, but with a slightly broader 445 distribution of cell lengths, with ~88% of cells falling between 1.5 and 4 µm compared to 446 ~97% of WT cells falling between these cell lengths. Interestingly, the Δ 5pbp mutant 447 retained its rod shape, similar to the \triangle 3pbp mutant of *A. tumefaciens*, suggesting that 448 449 these enzymes contribute minimally to sustaining proper rod shape under standard growth conditions. 450

451

Similar to our findings for *A. tumefaciens*, we were unable to make a deletion of *pbp1a*,
so we constructed a PBP1a depletion strain. Depletion of PBP1a led to a severe
viability defect (Supplementary Figure 7A) and cells became shorter (Figure 6B) and
wider (Supplementary Figure 7B). Thus, PBP1a is essential for polar growth and
maintenance of proper rod shape in both *S. meliloti* and *A. tumefaciens*. Taken
together, our data from *B. abortus*, *S. meliloti* and *A. tumefaciens* support the notion that
PBP1a plays an essential and conserved role in polar growth among the Rhizobiales.

459

460 **DISCUSSION**

Bacteria employ widely diverse growth strategies. Unipolar growth, or incorporation of
new cell wall material at a single pole, is a shared mode of growth among the

Rhizobiales, but the mechanisms that drive polar PG insertion remain poorly 463 understood. Most bacteria have multiple class A PBPs with semi-redundant functions 464 and growth is supported by the presence of any one of the PBPs [50] In C. crescentus, 465 four class A PBPs can support growth when expressed alone. [51] In B. subtilis, all 4 466 class A PBPs are dispensable and the SEDS protein RodA has likely taken over the 467 468 essential function of the class A PBPs during elongation [52] In contrast, here we identified the class A PBP1a homolog as an essential protein needed to synthesize PG 469 at the growth pole in A. tumefaciens and related genera, despite the presence of other 470 class A PBPs. Current findings support the proposal that in laterally growing bacterial 471 species, PBP1a homologs act as autonomous entities involved in PG remodeling or 472 repair and do not function as a part of the core elongasome [15, 16]. Instead, the 473 monofunctional glycosyltransferase RodA and the monofunctional transpeptidase PBP2 474 are required to maintain rod-shape [10, 53]. In contrast, we find that in the Rhizobiales, 475 476 which lack MreB, RodA and PBP2 homologs, the bifunctional enzyme PBP1a is an essential core component of the elongasome of polar growing Rhizobiales. Loss of 477 class A PBPs in *E. coli* or *B. subtilis* led to a decrease in cell width [54, 55]. Conversely, 478 479 depletion of PBP1a in A. tumefaciens and S. meliloti led to a significant decrease in cell length and an increase in cell width (Figure 1B, C, Supplementary Figure 1B). This 480 481 indicates that cells lacking PBP1a have a shorter period of cell elongation, and as a 482 result, likely spend more time synthesizing the septum (Figure 3C, Supplementary 483 Figure 3A, B). Thus, the regulation of PG synthesis that governs cell length and cell width in the Rhizobiales utilizes a novel mechanism compared to well-studied model 484 485 bacteria. Perhaps because class A PBPs function independently of cytoskeletal

complexes [10], PBP1a was freely available to assume the role of the primary enzyme 486 driving polar PG synthesis prior to the loss of the mre operon. Notably, the other 487 488 bifunctional PBPs minimally contributed to the maintenance of proper rod shape under the conditions tested in A. tumefaciens and S. meliloti. However, it is likely that the 489 additional class A PBPs may make dedicated contributions under specific growth 490 491 conditions. For example, in *E. coli*, PBP1a homologs are required for optimal growth in alkaline pH, while PBP1b homologs are required under acidic conditions [56]. Since the 492 plant rhizosphere is an acidic environment [57], it is possible that the remaining 493 494 bifunctional PBPs have specialized functions to maintain growth when bacteria are associated with a plant host. Additionally, duplication of the monofunctional PBP3 495 homolog among the Rhizobiales is restricted to a few species that interact with plants, 496 suggesting this is not a broad solution to the loss of PBP2, but that PBP3b homologs 497 may contribute more significantly to bacterial growth in plantae. In agreement with this 498 499 idea, we found that of the two monofunctional PBP homologs (PBP3a and PBP3b), PBP3a - a clear homolog of the division-specific PBP3 - contributes to cell division, 500 while deletion of *pbp3b* plays a minimal role in cell growth or cell shape under standard 501 502 laboratory conditions but forms a synthetic lethal pair with PBP3a that functions in cell division. 503

504

In laterally growing bacterial species, the role of scaffolding PG synthase enzymes
during elongation is fulfilled in part by the actin homolog MreB [58]. Homologs of
MreBCD are absent in the Rhizobiales; thus, how PBP1a is recruited to the growth pole
and how its activity is regulated remains unexplored. Recently, GPR (for Growth Pole

509	Ring), a large (~226 kDa) apolipoprotein with similarity to the polar organizing protein
510	TipN from <i>C. crescentus</i> , was reported to form a ring at the growth pole in <i>A.</i>
511	tumefaciens, with depletion of this protein leading to rounded cells [59]. This phenotype
512	implicates GPR as a possible candidate to scaffold PG enzymes during elongation. In
513	addition, PG synthesis by PBP1a also requires hydrolysis of the existing sacculus to
514	allow for insertion of new muropeptides. A DD-endopeptidase (RgsM) that is predicted to
515	have hydrolysis activity was recently shown to be essential for polar growth in S. meliloti
516	[60], and thus represents an interesting candidate for polar PG hydrolysis.
517	
518	Several lines of evidence suggest that FDAAs are primarily incorporated into the cell

wall through remodeling by LDTs [41, 42, 61, 62]. Cells depleted of PBP1a label 519 robustly at the growth pole with FDAAs, independently of PBP-mediated FDAAD 520 labeling. We also found that FDAAs label WT cells along the sidewalls of the new pole 521 much more brightly than the old pole. This points to a role for LDTs in not only polar 522 growth, but also sidewall remodeling of the new pole daughter cell (Figure 7). 523 524 Collectively, our results indicate that at the pole in *A. tumefaciens* FDAAD probes report on nascent PG synthesis by PBP1a, whereas FDAAs are incorporated by the localized 525 activity of LD-transpeptidases on the bacterial cell surface. These findings demonstrate 526 527 the usefulness of FDAAD probes for distinguishing between growth-dependent, nascent cell wall synthesis and growth-independent cell wall remodeling. In particular, the 528 529 growth pattern of species for which a high proportion of growth-independent remodeling 530 occurs can be obscured by FDAAs. Therefore, FDAADs are more useful in 531 distinguishing nascent PG synthesis mediated by PBPs.



532

533 Figure 7. Model of cell wall synthesis in the Rhizobiales. Unipolar elongation: A. tumefaceins 534 elongates from a single pole using the action of an essential PBP1a homolog. As the cell gets longer, the sidewalls of the new pole begin to be remodeled in a growth-independent manner, through the activity of 535 536 LD-transpeptidases. Transition: prior to cell division growth at the pole is terminated and growth at the 537 midcell is initiated, while remodeling of the new pole compartment continues. Cell division: during cell 538 division new PG is added at the midcell through the transpeptidase activity of two class B PBPs (with 539 PBP3a being the primary synthase) and the glycosyltransferase FtsW. Cell separation: after cell separation, the new poles resume polar elongation from what was the site of cell division. Continuous 540 remodeling of the new pole daughter cell occurs throughout the cell cycle and is inherited into the new 541 542 pole daughter cell.

543

Since LDTs are resistant to most classes of β -lactam antibiotics [63], crosslinking the 545 cell wall via LDTs may contribute to the high antibiotic resistance to β -lactam antibiotics 546 547 in the Rhizobiales. Here, we show that meropenem and faropenem inhibit LD- and DDtranspeptidation in *A. tumefaciens*, indicating that they may target LDTs and/or PBPs. 548 Perhaps the loss of LDT activity disrupts the activity of high molecular weight PBPs if 549 550 they function together in a growth pole complex. A combination of LDT and PBPtargeting antibiotics acted synergistically in killing *M. tuberculosis* [52], and a similar 551 approach may also be effective against species in the Rhizobiales. Since faropenem 552 553 causes swelling of the growth pole in A. tumefaciens, S. meliloti and B. abortus it is likely that the target(s) of this drug are conserved components of the growth machinery. 554 Identification of the specific cellular target(s) of faropenem will provide candidate 555 proteins for further characterization. 556

557

Expanding our understanding of the mechanism of polar growth in the Rhizobiales will 558 help to shed light on the different strategies that can be employed by bacteria during cell 559 elongation. Indeed, our findings highlight intriguing parallels with other, distantly related, 560 polarly growing bacteria including the Actinobacteria. For example, PBP1a is essential 561 in Mycobacterium smegmatis [64], an Actinobacteria that grows by bipolar elongation. In 562 563 addition, PBP1a localizes to growth poles and is important for maintenance of rod shape in other Actinobacteria [65, 66]. Therefore, the reliance on PBP1a for synthesis of 564 PG at the pole may be a key feature of polar-growing bacteria that arose independently 565 566 in the Rhizobiales and Actinobacteria, indicating convergent evolution. Notably, the cell 567 wall of polar-growing bacteria in both clades also contain a high proportion (~30-80%) of

LDT-crosslinked PG [1, 67, 68], suggesting that LD-crosslinks may provide structural 568 integrity. Furthermore, in *Mycobacterium* deletion of LDT-encoding genes leads to a 569 loss of rod-shape [63], and LDTs also contribute to active PG synthesis of the sidewalls 570 [37]. Finally, carbapenem antibiotics are routinely used to treat *M. tuberculosis* 571 infections [20] hinting that the target of these drugs may be important in polar growing 572 573 bacteria. Future work directed at characterizing the role of LDTs during polar growth in Rhizobiales and Actinobacteria is needed to determine if the high degree of LD-574 crosslinking is an innovation which allows for polar elongation to be adopted as the 575 primary mode of growth. Overall, the possibility that there may be governing principles 576 which allow for polar growth to emerge as a successful growth strategy is a fascinating 577 concept which merits further study. 578

579

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596	
597	MATERIALS AND METHODS
598	Bacterial strains, plasmids, and growth conditions. A list of all bacterial strains used
599	in this study is provided in Supplemental Table 1. Agrobacterium tumefaciens C58 and
600	derived strains were grown in ATGN minimal media [69] without exogenous iron at 28°C
601	with shaking. When appropriate, kanamycin (KAN) was used at the working
602	concentration of 300 μ g/ml. When indicated, isopropyl β -D-1-thio-galactopyranoside
603	(IPTG) was used as an inducer at a concentration of 1 mM. Sinorhizobium meliloti
604	strains were grown in Tryptone-Yeast (TY) medium. When appropriate, KAN was used
605	at the working concentration of 100 $\mu\text{g/ml},$ Gentamicin (GM) was used as 20 $\mu\text{g/ml},$ and
606	IPTG was used at a concentration of 500 μ g/ml. Brucella abortus strain S19 was grown
607	in brucella broth. E. coli strains were grown in Luria-Bertani medium at 37°C. For E. coli
608	DH5 α and S17-1 λ <i>pir</i> , when appropriate 50 µg/ml or 30 µg/ml of KM, respectively, was
609	added.

610

611 Construction of strains and plasmids. A list of all primers used in this study is
612 provided in Supplemental Table 2. For amplification of target genes, primer names

indicate the primer orientation and added restriction sites. All expression vectors were
verified by sequencing. All vectors were introduced into *A. tumefaciens* strains utilizing
standard electroporation protocols [70] with the addition of IPTG in the media when
introducing plasmids into depletion backgrounds.

617

Construction of deletion/depletion plasmids and strains. Vectors for gene deletion 618 by allelic exchange were constructed using recommended methods for A. tumefaciens 619 620 [70]. Briefly, 500-bp fragments upstream and 500 bp downstream of the target gene were amplified using primer pairs P1/P2 and P3/4 respectively. Amplicons were spliced 621 together by SOEing using primer pair P1/P4. The amplicon was digested and ligated 622 into pNTPS139. The deletion plasmids were introduced into A. tumefaciens by mating 623 using an *E. coli* S17 conjugation strain to create KM resistant, sucrose sensitive primary 624 integrants. Primary integrants were grown overnight in media with no selection. 625 Secondary recombinants were screened by patching for sucrose resistance and KM 626 sensitivity. Colony PCR with primers P5/P6 for the respective gene target was used to 627 628 confirm deletion. PCR products from P5/P6 primer sets were sequenced to further confirm deletions. 629

For depletion strain construction, target genes (*pbp1a* or *pbp3a*) were amplified, digested and ligated into pUC18-mini-Tn7T-GM-P_{lac}. The mini-Tn7 vector, along with the pTNS3 helper plasmid, were introduced into C58 Δ *tetRA*::a-*att*Tn7 as described previously [27]. Transformants were selected for GM resistance and insertion of the target gene into the a-*att* site was verified by colony PCR using the tet forward and Tn7R109 primer. PCR products were sequenced to confirm insertion of the correct

gene. Next, the target gene was deleted from the native locus as described above in thepresence of 1 mM IPTG to drive expression of the target gene from the engineered site.

638 To generate the S. meliloti strain lacking the five non-essential PBPs, the corresponding genes were consecutively deleted from the Rm2011 rgsP-egfp genome 639 using the sucrose selection method. [71] To generate the S. meliloti MrcA1 depletion 640 strain, first plasmid pK18mobsac-mrcA1del was integrated into the Rm2011 rgsP-egfp 641 genome, then an ectopic mrcA1 copy was introduced on plasmid pGCH14-mrcA1 642 followed by sucrose selection of mutant clones with deletion of the native mrcA1 allele. 643 The curable plasmid pGCH14, which is maintained in single copy in S. meliloti due to 644 the replication operon repABCpMlb lacO, prone to repression by LacI was used as 645 vector to conditionally establish the ectopic copy of mrcA1 under control of its native 646 promoter. pSRKKm, carrying *lacl*, was introduced into the strain with chromosomal 647 deletion of *mrcA1*, carrying pGCH14-mrcA1, and the resulting strain Rm2011 rgsP-eafp 648 mrcA1^{dpl} was grown in presence of 500 µm IPTG. Growth in the absence of IPTG 649 induced MrcA1 depletion due to loss of the ectopic mrcA1 copy. 650

651

Phase and Fluorescence microscopy. A small volume (~1 µl) of cells in exponential
phase (OD₆₀₀ = 0.2 - 0.4) was applied to a 1% ATGN agarose pad as described
previously [72]. DIC, Phase contrast and epifluorescence microscopy were performed
with an inverted Nikon Eclipse TiE and a QImaging Rolera em-c2 123 1K EMCCD
camera with Nikon Elements Imaging Software. For time-lapse microscopy, images
were collected every ten minutes, unless otherwise stated.

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A T O N

659	Quantification of cell length distributions. Cells were grown overnight in ATGN.
660	Cells were diluted in ATGN to an $OD_{600} = 0.2$ and allowed to grow until reaching an
661	$OD_{600} = 0.4 - 0.6$. Live cells were imaged using phase contrast microscopy and cell
662	length distributions of the indicated number of cells per strain were determined using the
663	longest medial axis as measured by MicrobeJ software [24].

664

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Quantification of cell morphologies, FDAA and FDAAD labeling patterns. For A. 665 666 tumefaciens, cells were grown overnight in ATGN media and diluted in the same conditions to an $OD_{600} = 0.20$ and allowed to grow until reaching an $OD_{600} = 0.4 - 0.6$. At 667 this point cells were labeled with 1mM of the fluorescent-D-amino acid (FDAA) HCC-668 amino-D-alanine (HADA) or the fluorescent dipeptide (FDAAD) NBD-amino-D-alanine-D-669 alanine (HADA—DA) as previously described [36, 38, 41]. Immediately following a five-670 minute incubation, cells were ethanol fixed to prevent further growth. Phase contrast 671 and epifluorescence microscopy was performed on the reported number of cells. For E. 672 coli, cells were grown overnight at 37°C in M9 + 0.2% glucose minimal medium and 673 diluted in the same conditions to an $OD_{600} = 0.1$ and allowed to grow until reaching an 674 OD₆₀₀ = 0.4 - 0.6. At this point cells were labeled with 1 mM of HADA—DA. After an 675 incubation of 90 minutes cells were ethanol fixed and washed 3 times in 1 mL PBS 676 677 before imaging. For *B. subtilis 3610*, cells were grown overnight at 37°C in S750 + 1% glucose defined minimal medium and diluted in the same conditions to an $OD_{600} = 0.1$ 678 and allowed to grow until reaching an $OD_{600} = 0.4 - 0.6$. At this point cells were labeled 679 680 with 5 mM of HADA—DA. After an incubation of 120 minutes cells were ethanol fixed 681 and washed 3 times in 1 mL PBS before imaging. For S. venezuelae, cells were grown

682	overnight at 30 $^{\circ}$ C in LB medium and diluted in the same conditions to an OD ₆₀₀ = 0.1
683	and allowed to grow until reaching an $OD_{600} = 0.4 - 0.6$. At this point cells were labeled
684	with 1 mM BODIPY FL-amino-d-alaninyl-D-alanine (BADA—DA). After an incubation of
685	15 minutes, cells were washed once with 1 mL LB and resuspended in 500 uL LB
686	containing 2 mM HADA—DA. After an incubation of 15 minutes, cells were washed
687	once with 1 mL LB and resuspended in 500 uL LB containing 1 mM ATTO 610-amino-d-
688	alaninyl-D-alanine (Atto610ADA—DA). After an incubation of 15 minutes cells were
689	ethanol fixed and washed 3 times in 1 mL PBS before imaging.
690	

Demographs were constructed using MicrobeJ. For demographs, cells were arranged from top to bottom according to their cell lengths and each cell was oriented such that the new pole (defined as the cell pole with the higher fluorescence intensity as determined by FDAA or FDAAD labeling or the smaller pole diameter in cells without label) was oriented to the right.

696

697 Synthesis of HADA—DA.

698



A solution of 7-hydroxycoumarin-3-carboxylic acid (HCC) (105 mg, 0.51 mmol) and 701 carbonyldiimidazole (83 mg, 0.51 mmol) in anhydrous DMF (5 mL) was stirred at room 702 703 temperature (RT) for 2 h under an atmosphere of argon. Boc-D-2,3-diaminopropionic acid (104 mg, 0.51 mmol) was then added and the reaction mixture was allowed to stir 704 at RT for 18 h. DMF was removed in vacuo. The residue was diluted with EtOAc (35 705 mL), washed with 1N HCI (20 mL) and water (30 mL). The combined aqueous layers 706 707 are back-extracted with EtOAc (20 mL). The combined organic layers were washed with 708 brine (20 mL), dried over sodium sulfate, filtered, and concentrated to dryness. The 709 crude product was dissolved in anhydrous DMF. D-alanine tert-butyl ester hydrochloride (111 mg, 0.612 mmol), HOAt (83.5 mg, 0.612 mmol) and NaHCO₃ (94 mg, 1.12 mmol) 710 were added successively, and the reaction mixture was cooled to 0 °C. EDCI (117 mg, 711 0.612 mmol) was then added, and the reaction mixture was stirred for 12 h. The solvent 712 713 was removed in *vacuo* and the product was diluted with EtOAc, washed with 1N HCl, water, brine, dried over sodium sulfate, filtered and concentrated. The crude product 714 715 was dissolved in DCM/TFA (1:1, 4 mL) mixture, stirred for 1 h and evaporated to 716 dryness. The yellow color solid was dissolved in acetonitrile/water and purified by 717 reverse phase HPLC and lyophilized to yield the desired product as a pale yellow solid (63 mg, 26 %). 1H NMR (500 MHz, DMSO-d6) 11.37 (br s, 1H), 8.91 (m, 1H), 8.90 (m, 718 1H), 8.81 (s, 1H), 8.42 (br s, 3H), 7.83 (m, 1H), 5.94 (m, 1H), 5.89 (s, 1H), 4.24 (m, 1H), 719 4.04 (br s, 1H), 3.77 (m, 1H), 3.68 (m, 1H), 1.32 (s, 1H); HRMS-ESI-TOF m/z calc 720 C16H18N3O7 (M+H) 364.1145, Found 364.1161. 721 722

723 **PG compositional analysis**.

For PG analysis, three cultures of each of strain were grown overnight in 3 ml culture 724 tubes of ATGN minimal media at 28°C with shaking; the + PBP1a strain was 725 726 supplemented with 1mM IPTG. The 3 ml cultures were then added to 50 ml flasks of fresh ATGN and allowed to grow under the same conditions until reaching an 727 exponential phase OD_{600} of 0.5-0.6. Cells were then pelleted by centrifugation at 4000 x 728 729 q for 10 minutes. Cell pellets were washed three times with ATGN by centrifugation and resuspension to remove IPTG. After the final wash, the 3 cell pellets from the + PBP1a 730 strain were split and resuspended in 50 ml ATGN with or without IPTG. Each culture 731 732 was grown for 16 hours (hr). to an OD₆₀₀ of 0.6 After 146 h of growth, 50 ml of the exponential cultures were collected and pelleted by centrifugation at 4000 x g for 20 733 minutes. Cell pellets were resuspended in 3 mL of ATGN and 6 mL of 6% SDS and 734 stirred with magnets while boiling for 4 h. Next, samples were removed from heat but 735 continued to stir overnight. Samples were then shipped to Cava laboratory for 736 737 purification and analysis. Upon arrival, cells were boiled and simultaneously stirred by magnets for 2 h. After 2 h, boiling was stopped, and samples were stirred overnight. PG 738 was pelleted by centrifugation for 13 minutes (min) at 60,000 rpm (TLA100.3 Beckman 739 740 rotor, Optima Max-TL ultracentrifuge; Beckman), and the pellets were washed 3 to 4 times by repeated cycles of centrifugation and resuspension in water. The pellet from 741 742 the final wash was resuspended in 50 µl of 50 mM sodium phosphate buffer, pH 4.9, 743 and digested overnight with 100 µg/ml of muramidase at 37°C. Muramidase digestion 744 was stopped by boiling for 4 min. Coagulated protein was removed by centrifugation for 15 min at 15,000 rpm in a desktop microcentrifuge. The muropeptides were mixed with 745 746 15 µl 0.5 M sodium borate and subjected to reduction of muramic acid residues into

747	muramitol by sodium borohydride (10 mg/ml final concentration, 20 min at room
748	temperature) treatment. Samples were adjusted to pH 3 to 4 with orthophosphoric acid
749	and filtered (0.2- μ m filters). Analysis of muropeptides was performed on an ACQUITY
750	Ultra Performance Liquid Chromatography (UPLC) BEH C18 column, 130Å, 1.7 μ m, 2.1
751	mm x 150 mm (Water, USA) and detected at Abs. 204 nm with ACQUITY UPLC UV-
752	visible detector. For data shown in Figure 4B and Supplementary Figure 7B,
753	muropeptides were separated with organic buffers at 45°C using a linear gradient from
754	buffer A (formic acid 0.1% (v/v) in water) to buffer B (formic acid 0.1% (v/v) in
755	acetonitrile) in a 18 minutes run with a 0.25 ml/min flow. For data shown in Figure 5A
756	and Supplementary Figure 5A, muropeptides were separated using a linear gradient
757	from buffer A (sodium phosphate buffer 50 mM pH 4.35) to buffer B (sodium phosphate
758	buffer 50 mM pH 4.95 methanol 15% (v/v)) with a flow of 0.25 mL/min in a 20 min run.
759	Individual muropeptides were quantified from their integrated areas using samples of
760	known concentration as standards. Muropeptide abundance was statistically compared
761	using a one-way ANOVA with Tukey's multiple comparisons test.

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