#### Cell Diameter in Bacillus subtilis is Determined by the Opposing Actions of Two Distinct 1 2 Cell Wall Synthetic Systems

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#### 20 Abstract

- 21 Rod shaped bacteria grow by adding material into their cell wall via the action of two spatially distinct enzymatic systems:
- 22 The Rod system moves around the cell circumference, while the class A penicillin-binding proteins (aPBPs) are
- 23 unorganized. To understand how the combined action of these two systems defines bacterial dimensions, we examined how
- 24 each system affects the growth and width of Bacillus subtilis, as well as the mechanical anisotropy and orientation of
- 25 material within their sacculi. We find that rod diameter is not determined by MreB, rather it depends on the balance between
- 26 the systems: The Rod system reduces diameter, while aPBPs increase it. RodA/PBP2A can both thin or widen cells,
- 27 depending on its levels relative to MreBCD. Increased Rod system activity correlates with an increased density of directional
- 28 MreB filaments, and a greater fraction of directionally moving PBP2A molecules. This increased circumferential synthesis
- 29 increases the amount of oriented material within the sacculi, increasing their mechanical anisotropy and reinforcing rod
- 30 shape. Together, these experiments explain how the combined action of the two main cell wall synthetic systems build rods
- 31 of different widths, a model that appears generalizable: Escherichia coli containing Rod system mutants show the same
- 32 relationship between the density of directionally moving MreB filaments and cell width.

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#### 35 Introduction

While the length of *Bacillus subtilis* rods increases as a function of their growth rate <sup>1</sup>, their width remains constant across different growth conditions <sup>2</sup>. How bacteria define and maintain these rod shapes with such precision is not understood, but it must involve mechanisms controlling both the rate and location of glycan insertion into the peptidoglycan (PG) sacculus, the enveloping heteropolymer meshwork that holds cells in shape <sup>3</sup>. In order to understand how bacteria grow in defined shapes, we need to understand not only where these enzymes act, but how their activity affects the geometry and

41 arrangement of material within the sacculus.

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43 The PG for cell elongation is synthesized by two families of penicillin-binding proteins (PBPs). The class A penicillin-binding 44 proteins (aPBPs) both polymerize and cross-link glycans. The class B penicillin-binding proteins (bPBPs) cross-link <sup>4,5</sup> the 45 glycans polymerized by the glycosyltransferase RodA <sup>6</sup>. bPBPs and RodA are components of the "Rod complex", a group 46 of proteins essential for rod shape (Figure 1A). In B. subtilis, this contains RodA, the class B transpeptidase PBP2A, MreC, 47 MreD, RodZ, and filaments of MreB, an actin homolog <sup>7,8</sup>. MreB polymerizes together with other MreB homologs into short, 48 highly curved filaments on the membrane 9.10. To maximize their interaction with the membrane these curved filaments 49 deform to the membrane, orienting along the direction of highest inward membrane curvature, pointing around the rod width 50 <sup>11</sup>. Oriented by their association with MreB filaments, the Rod complex moves directionally around the cell circumference, 51 driven by the PG synthetic activity of RodA/PBP2A<sup>12-15</sup>. This radial motion of independent filament-enzyme complexes are 52 believed to insert PG strands around the rod width <sup>16</sup>.

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aPBPs also affect rod shape, as *B. subtilis* cells lacking them are thinner <sup>17</sup>. Single molecule and biochemical studies indicate aPBPs and the Rod system are both spatially and functionally distinct <sup>6,14</sup>: While Rod complexes move around the cell width, aPBPs have never been seen to move directionally. Rather, they diffuse within the membrane, occasionally pausing and remaining immobile for a few seconds <sup>14</sup>. Furthermore, the synthetic activity of either aPBPs or the Rod complex can be inhibited without affecting the activity or motions of the other <sup>6,14</sup>. Given that both synthetic systems are required for growth, but that Rod complex activity is spatially organized while aPBP activity is not (**Figure 1A**), it is not clear how these two PG synthetic machineries work together to create a rod-shaped sacculus of a given width.

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62 Current models of rod width have focused on MreB filaments, attributing the altered widths of Escherichia coli MreB mutants

to possible changes in MreB filament curvature, twist, angle, or localization to negative Gaussian curvature <sup>18-24</sup>. Not only do these models lack the effects of aPBP mediated synthesis, they are: A) theoretical, as these proposed changes in filament curvature or twist have not been structurally validated, or B) difficult to reconcile in *B. subtilis*, which has no detectable negative Gaussian curvature <sup>11</sup> or skew in filament angle relative to the cell body <sup>11-13,25</sup>.

67 Rather than focusing on MreB alone, we sought to develop a more thorough understanding of how the synthesis of both 68 organized and unorganized PG affects the width and growth of B. subtilis, as well as their effects on the organization and 69 mechanics of cell wall material. We find that aPBP and Rod complex-mediated PG synthesis have opposing effects on rod 70 width, and that cell diameter depends on their balance. The rate cells add surface area is largely unaffected by the level of 71 any one system, unless both become limiting. As MreBCD expression increases and rods thin, we observe a greater density 72 of directionally moving MreB filaments and a greater fraction of directionally moving enzymes. Increased Rod complex 73 activity creates a greater proportion of oriented material within the sacculus, causing the rod to stretch less across its width 74 and more along its length in response to internal turgor. Finally, we show that the different widths of E. coli Rod mutants 75 also show correlation between the density of directionally moving MreB filaments and cell width, giving a simple, 76 generalizable model that may explain the role of the Rod system in cell width.

#### 77 Results

#### 78 The Rod and aPBP systems have opposing effects on cell diameter

The width of rod-shaped bacteria has been attributed to properties encoded within MreB filaments <sup>19,21,24,26</sup>. We reasoned that if width were defined by MreB, then the *mreBCD* genes from larger bacteria should produce cells with larger diameters. To test this, we replaced the *B. subtilis mreBCD* operon with the *mreBCD* operon from *Bacillus megaterium*, a nearly 2-fold wider bacterium. Surprisingly, these cells grew as rods only slightly wider than wild type (WT) *B. subtilis,* and by further overexpressing *B. megaterium mreBCD* cells became even thinner (36 nm) (Figure 1B, S1A), suggesting that MreB filaments themselves do not encode a specific rod width.

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We next examined how width changed as we independently controlled the levels of the two main *B. subtilis* PG synthetic systems. We created strains where expression of PBP1 (the major aPBP, encoded by *ponA*) was under IPTG control. As PG synthesis by the Rod system depends upon MreB <sup>14</sup>, we created strains where the native *B. subtilis mreBCD* operon was under xylose control. We grew these strains at different induction levels, then measured their steady-state widths using

- 90 fluorescence microscopy. For a subset of inductions, we measured the relative protein abundance using proteomic mass 91 spectrometry, normalizing to the levels in WT cells grown in CH media (**Supplemental File 1**).
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- Varying *mreBCD* inductions revealed the Rod system has a thinning effect on rod shape. In the lowest inductions supporting growth, rods were ~2-fold wider than WT, with some cells losing rod shape. As we increased *mreBCD* induction, cells became thinner, reaching WT width when MreB abundance recapitulated WT levels (Figure 1C, Figure S1B). Inductions above this point resulted in cells becoming 33 nm thinner than WT, and thinner still (58 nm) in *mreBCD* merodiploids. As above, these results demonstrate MreB filaments do not *define* a given rod diameter, rather as previously hypothesized <sup>11,27</sup>, they indicate the Rod system *reduces* cell diameter.
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Different *ponA* inductions revealed that aPBPs have a widening effect: With no IPTG, cells contained 0.25 the amount of WT PBP1, and were ~23% thinner than WT, similar to  $\Delta$ *ponA* strains <sup>17</sup> (Figure 1D, Figure S1C). As we increased *ponA* induction cells became wider, reaching WT widths when PBP1 abundance recapitulated WT levels. Inductions above this point caused cells to become increasingly wider, and by expressing *ponA* under stronger promoters or in merodiploids, we could produce rods almost twice WT diameter.

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106 These results demonstrate that the aPBPs and Rod system have opposing effects: The circumferentially moving Rod 107 complex reduces cell diameter, while the spatially unorganized aPBPs increase diameter. We hypothesized that a balanced 108 expression of each system could produce WT diameter rods. We combined the IPTG-inducible ponA and xylose-inducible 109 mreBCD alleles into one "dual-inducible" strain. We found six different pairs of xylose and IPTG concentrations that 110 produced rods of WT diameter (Figure 1E, Figure S1D), even though individually, each induction resulted in perturbed 111 diameters in the singly-inducible parental strains (Figure 1C-D). Relative quantitation of protein levels revealed that, in the 112 induction pairs at or beneath WT levels, these cells contained reduced, but relatively balanced amounts of PBP1 relative to 113 MreB. To determine this balance in native conditions, we measured the widths and protein levels of WT cells grown in 114 different media that result in different growth rates (Figure 1F, Figure S1E). Together, this data suggested width depends 115 on the balance between the aPBP and the Rod systems. By plotting the ratio of the [Fold change PBP1] / [Fold change 116 MreB] for all conditions in the data sets against their width (Figure 1G), we found that B. subtilis maintains its diameter 117 within ~5% of WT only when this ratio was within a range of 0.8 to 1.5; outside of this range, cell diameter diverged. WT

cells in different media showed PBP1/MreB ratios ranging from 0.9 to 1.5. Likewise, the induction pairs yielding WT widths in the dual-inducible strain were within 0.8 to 1.0, only when their levels did not exceed WT. In contrast, the single-inducible *ponA* or *mreBCD* strains were within 5% of WT width only when the ratio was within this range; outside of it, width rapidly diverged. Together, these results indicate that, at the level of PG insertion, cell width is affected by the levels of the two opposing synthetic systems that insert material into the sacculus (Figure 1H).

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124 RodA can function outside of the Rod system to widen cells, but only when PBP2A is also in excess. Given that 125 RodA acts within the Rod complex with PBP2A (encoded by pbpA) to synthesize PG<sup>14</sup>, titrations of rodA and pbpA should 126 show the same "thinning effect" as mreBCD. However, rodA overexpression can restore WT width to ΔaPBP cells <sup>6</sup>, a 127 widening activity similar to ponA. To investigate this discrepancy, we made two sets of strains where rodA or pbpA (as the 128 only elongation-specific bPBP in the cell) were under the control of increasingly strong inducible promoters. As before <sup>28,29</sup>. 129 low rodA or pbpA yielded wide cells; and as induction increased, cells gradually thinned to WT width, suggesting both are 130 required for functional Rod complexes. However, while we expected excess RodA to create thinner cells, once rodA 131 induction exceeded the level required for normal width, it behaved like excess ponA, fattening cells far beyond WT diameter 132 <sup>6</sup> (Figure 2A). In contrast, *pbpA* inductions beyond this point had a negligible effect on diameter.

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Given that RodA and PBP2A must interact to synthesize PG <sup>30</sup>, we tested if the widening caused by RodA overexpression required excess PBP2A. We created two strains that had *rodA* under xylose control; one contained native *pbpA*, and the other had *pbpA* under IPTG control (both as the only elongation-specific bPBP in the cell). As before, when *pbpA* was under native control, depletion or overexpression of *rodA* caused diameter to increase (Figure 2B). In contrast, when *pbpA* was held at the lowest induction required for WT width and *rodA* was simultaneously overexpressed, cells remained at WT widths. However, these RodA-overproducing cells could be made increasingly wider by increasing *pbpA* induction (Figure 2B), demonstrating excess RodA requires excess PBP2A to increase width.

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These results suggested RodA/PBP2A widens cells when their levels exceed the rest of the Rod complex. If this hypothesis is correct, wide cells caused by RodA/PBP2A overexpression should become thinner if MreBCD is increased, perhaps recruiting the diffusive excess RodA/PBP2A into circumferentially-moving, "thinning" Rod complexes. Indeed, while strong induction of *rodA* made wide cells, simultaneous overexpression of a second copy of *mreBCD* reduced cell diameter

- (Figure 2C), a narrowing far greater than what we observed when overexpressing *mreBCD* in otherwise WT strains (Figure
   1C). Thus, the aPBP-like widening activity of RodA/PBP2A likely occurs once they exceed some level of MreBCD, possibly
   reflecting a saturation of binding sites within Rod complexes.
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#### 151 Growth rates are maintained across a wide range of enzyme levels, unless both systems become limiting.

We next examined how each PG synthetic system affected the rate of cell growth in our dual-induction strain using two assays: 1) single-cell measurements of the rate of surface area addition <sup>31</sup> under agar pads and 2) population measures of growth in a shaking plate reader. First, we examined the growth of the dual-inducible strain at the *mreBCD* and *ponA* inductions that produced WT widths (Figure 1E). At the lowest induction pairs, growth was greatly reduced in both assays, and increased with each increasing induction pair (Figure 2D), up to the induction pair that produced WT PBP1 and MreB

- 157 protein levels. Thus, growth can be reduced if both PG synthesis systems become limiting.
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159 Next, we assayed growth as we titrated either mreBCD or ponA while holding the other constant. Similar to E. coli MreB 160 studies <sup>22,32,33</sup>, both assays showed no difference in growth across a wide range of *mreBCD* inductions, except at the lowest 161 induction where cells frequently lost rod shape. Similarly, both assays saw no difference in growth across our ponA 162 induction range. Thus, even though these cells have different geometries, they add new surface area at the same rate. We 163 next examined how extremely low levels of aPBPs affected growth. Similar to previous observations <sup>17,34,35</sup>, both Δ*ponA* and 164 ΔaPBP (ΔpbpF, ΔpbpG, ΔpbpD, ΔponA::kan) strains showed a marked reduction in bulk growth, a defect rescued by 165 overexpression of RodA <sup>6,36</sup>. However, the single cell measurements revealed a surprise: Both Δ*ponA* and ΔaPBP cells 166 showed no defect in their single-cell growth rate, adding surface area at the same rate as WT, as did RodA overexpressing 167 ΔaPBP cells (Figure 2D). Given ΔaPBP cells display an increased rate of lysis that can be suppressed by RodA 168 overexpression <sup>6,36</sup>, it appears the population growth rate defect of aPBP-deficient cells arises not from a reduction in growth 169 rate, but from an increased frequency of death.

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Increased MreBCD increases the density of directionally moving MreB filaments and the fraction of directionally
 moving synthetic enzymes.

174 To gain a mechanistic link between the level of the PG synthetic systems to cell width, we sought to develop microscopic 175 measures of their activity. While the fraction of stationary aPBPs would be difficult to quantify. Rod complexes exhibit a 176 more quantifiable phenotype: As their motion is driven by PG synthesis, the cellular activity of the Rod system can be 177 measured by quantitating the number of directionally moving MreB filaments <sup>37</sup>. We developed an analysis method that, 178 using total internal reflection fluorescence microscopy (TIRFM) data, determines the density of MreB filaments moving 179 directionally around the cell. Filaments undergoing directed motion are detected by taking advantage of the temporal 180 correlation occurring between adjacent pixels across the cell width as objects move through them. These objects are then 181 counted over a given time and then normalized to the total cellular surface area (Figure 3A, S3A-F). To evaluate the 182 robustness of this approach to filament number, velocity, angle, and length, we simulated TIRFM time-lapses of MreB 183 filaments in cells (Figure S3G-K). This revealed this approach was more accurate than counting filaments in the same data 184 with particle tracking. Furthermore, when applied to real TIRFM movies of MreB filaments, this approach yielded numbers 185 comparable to tracking MreB filaments inside cells imaged with TIRF - structured illumination microscopy (TIRF-SIM) 186 (Figure S3L). With this analysis tool, we examined how the density of directionally moving MreB filaments related to rod 187 width as we titrated the expression of an inducible mreB-sw-msfGFP, mreCD operon expressed as the only source at the 188 native locus. This analysis revealed a well fit, apparently linear correlation between directional filament density and width 189 (Figure 3B). Thus, the density of directionally moving MreB filaments reduces cell width, possibly by promoting directional 190 PG synthesis.

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To this end, we next examined how MreBCD levels affected PBP2A motion. Multiple studies have noted PBP2A molecules exist as a mixed population; some diffuse within the membrane, while others move directionally <sup>12-14</sup>. We titrated *mreBCD* induction as we tracked the motions of PBP2A molecules (expressed at the native locus as a HaloTag fusion, sparsely labeled with JF-549 <sup>38</sup>). At low *mreBCD*, only a small fraction of PBP2A moved directionally. As we increased *mreBCD* induction, an increasing fraction of PBP2A molecules moved directionally (Figure 3C, Movie SM1), demonstrating that *mreBCD* limits the amount of directional PBP2A.

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199 Increased MreBCD:PBP1 correlates with an increased amount of oriented material and structural anisotropy of the200 cell wall.

201 To gain more insight into how an increased density of directional rod complexes can reduce rod width, we used polarization 202 microscopy to understand how increased circumferential synthesis affected the organization of material within the cell wall. 203 Polarization microscopy reports on both the angle and extent of orientation within optically anisotropic (or birefringent) 204 materials<sup>39</sup>, and has been used to assay the orientation of various materials, including plant cell walls<sup>40-42</sup> and mitotic 205 spindles <sup>43</sup>. Polarization microscopy revealed purified WT sacculi show birefringence, indicating that some fraction of the 206 material within them is oriented. Focused at their surface, sacculi showed a predominant slow axis oriented along the rod 207 length (Figure 4A, Movie SM2). Given that amino acids have a higher refractive index compared to sugars, this suggests 208 (in agreement with previous models <sup>44</sup>) the peptide crosslinks are predominantly oriented along the rod length, and the 209 glycans are oriented around the circumference.

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211 We then examined how the balance between the Rod and aPBP systems affected the relative amount of oriented material 212 in the wall. We grew our dual-induction strain under three different induction conditions to create varied width cells: wide 213 (high ponA, low mreBCD), normal (ponA and mreBCD induced at WT levels), and skinny (low ponA, high mreBCD) (Figure 214 4B), purified their sacculi, and quantified their total retardance with polarization microscopy (Figure 4C, Movie SM3). 215 Retardance is the differential optical path length for light polarized parallel and perpendicular to the axis of molecular 216 alignment; alternatively, it is defined as birefringence ( $\Delta n$ ) times the physical path length through an anisotropic material. 217 Wide cells (high ponA, low mreBCD) had the lowest retardance, skinny cells (low ponA, high mreBCD) had the highest 218 retardance, and normal cells were in between (Figure 4D). Because retardance depends on both the thickness of material 219 (path length) and the degree of its orientation ( $\Delta n$ ), we normalized the retardance of each sample to the mean thickness of 220 the cell walls of each induction condition, obtained using transmission electron microscopy (Figure S4A). This revealed that 221 the skinny cell walls had more highly ordered material ( $\Delta n$ ) per path length (nm) of cell wall thickness, and that wide cell 222 walls had the least (Figure 4E). Thus, in agreement with recent atomic force microscopy studies showing orientated glycans 223 in E. coli require MreB<sup>16</sup>, these experiments demonstrate that as Rod system activity increases, so does the amount of 224 oriented material in the wall.

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The sacculi of *E. coli and B. subtilis* are mechanically anisotropic, stretching more along their length than across their width <sup>44-46</sup>. To test how the ratio of oriented to unoriented PG synthesis affected this property, we grew our dual-inducer strain at the three *mreBCD:ponA* inductions above, labeled their walls with Alexa-488-D-amino carboxamides <sup>4</sup>, and assayed the

dimensions of their cell walls before and after hyperosmotic shocks (**Figure 4F-G**). This revealed that increased Rod system activity correlated with an increased mechanical anisotropy of the sacculus: As we increased the expression of MreBCD relative to PBP1A, rods shrank less across their width, and more along their length (**Figure 4H, S4B-C**). Thus, the Rod system acts to reinforce rod shape against internal turgor by promoting oriented PG synthesis around the rod.

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#### 234 E. coli Rod mutants also show a correlation between cell width and the density of directionally moving filaments. 235 Previous studies have examined how MreB and PBP2 mutations affect the shape of E. coli, hypothesizing that their 236 abnormal widths arise from changes in the curvature, twist, or angle of MreB filaments. Our observations in B. subtilis 237 suggested an alternative explanation: Abnormal width might arise from simply changing the amount of Rod system activity. 238 We tested this by measuring the density of directional GFP-MreB filaments in these same mutant E. coli strains. As a 239 benchmark, we first assayed the width of *E.coli* as we titrated the expression of mreB-sw-msfGFP using CRISPRi against 240 msfGFP <sup>47</sup> <sup>28</sup>. As in *B. subtilis*, this yielded an inverse relationship between directional MreB filament density and cell width 241 (Figure 5A). Examining each group of mutants showed the same result: 1) An identical trend was observed for the 242 mutations hypothesized to change filament twist <sup>19</sup>, 2) as well as in the mutations designed to change filament curvature <sup>24</sup>. 243 And 3) notably, the same correlation between directional filament density and cell width was observed in strains where E. 244 coli mrdA (PBP2) was replaced with mrdA genes from other species <sup>22</sup>. TIRF-SIM imaging of these MreB mutants revealed 245 some insight into these effects (Movie SM4): Some of the wide mutants showed either A) longer but fewer filaments, or B) a 246 large fraction of immobile filaments. Conversely, some thinner mutants appeared to have more, but shorter filaments. Finally 247 4), while Colavin et al. hypothesized that RodZ reduces width by changing MreB filament curvature <sup>24</sup>, we found that 248 increased RodZ induction increased MreB filament density as cells thinned (Figure 5B, Movie SM5). This suggests that the 249 decrease in cell width most likely occurs from to RodZ's ability to nucleate MreB filaments 48 rather than changing their 250 curvature. Thus, the density of directionally moving Rod complexes correlates with cell width in both B. subtilis and E. coli 251 across multiple genetic perturbations.

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#### 253 Discussion

The shape of bacteria is defined by their cell walls; these experiments demonstrate that the two systems that insert PG into it have opposing roles on its shape. Due to the intrinsic orienting of MreB filaments around the rod width <sup>11</sup>, the Rod system inserts circumferentially oriented material around the rod circumference, reducing its diameter. As the number of MreB

filaments increases, so does the fraction of directional enzymes and the amount of oriented material in the wall. In contrast, the aPBPs do not move circumferentially, inserting material that isotropically enlarges the sacculus. Our data indicates the macroscopic shape of the sacculus arises from the nature of the material inserted into it: The more it is oriented around the rod circumference by the Rod complex, the less the rod stretches across its width, and the more it stretches along its length (Figure 5C).

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263 If the balance between the two PG synthetic systems is perturbed, the shape of the sacculus becomes altered, though its 264 rate of expansion remains constant. As both systems utilize the same pool of lipid II <sup>49</sup>, the flux through each may depend on 265 their relative levels; if one is reduced, the flux through the other may increase. This would explain why disrupting the Rod 266 system causes cells to swell <sup>27,29,50-52</sup>: In the absence of Rod-mediated thinning, aPBPs add more material uniformly over the 267 cell surface. Likewise, in the absence of aPBP-mediated widening, increased flux through the Rod system would explain 268 why cells become extremely thin <sup>17,53</sup>. However, if both systems are equivalently reduced, cells grow with normal widths, but 269 at slower rates; as long as the activities are balanced - identical shape arises from the balanced levels of enzymes, but 270 growth is reduced due to their combined activity becoming limiting. This would explain why ponA mutations rescue mreB 271 deletions <sup>54</sup>; equally crippling both systems may rebalance the activities such that the cell retains normal shape and viability.

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#### 273 Implications for the role of MreB in rod width determination.

274 Given that 1) mreBCD from the wide bacterium B. megaterium creates close to normal diameter B. subtilis rods, and 2) B. 275 subtilis diameter depends on mreBCD levels, we find it unlikely that any property of MreB filaments defines a given cell 276 diameter. Rather, MreB appears to be one component of a rod-thinning system, working in opposition to aPBP-mediated 277 widening. Indeed, in vitro studies have revealed MreB filaments are extremely curved (> 200 nm), allowing them to deform 278 to, and orient around the width of bacteria of any larger width 9,11. While it remains possible that given MreB mutations or 279 interactions with RodZ could indeed alter filament curvature or twist <sup>19,21,24,26</sup>, these experiments suggest a more 280 parsimonious explanation for their effect on Rod activity: These mutations alter MreB's polymerization dynamics or cellular 281 distribution. Reducing the amount or number of active MreB polymers would reduce Rod activity, causing cells to widen. 282 Likewise, mutations altering MreB's polymer length distribution or tendency for filaments to bundle would cause the 283 distribution of Rod activity to become non-uniform, causing some parts of the cell to thin while others would widen, as 284 previously observed for certain MreB mutants <sup>12,27</sup>.

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286	Additionally, our experiments reinforce observations that aPBPs and RodA (when in excess to MreBCD) serve reparative,
287	anti-lytic roles: aPBP-deficient cells grow at the same rate as WT, yet have an increased frequency of death, lysing as thin
288	rods without losing shape 6,36. Likewise, aPBP-mediated synthesis increases upon endopeptidase overexpression in E. coli
289	<sup>55</sup> . Given the active state of aPBPs correlates with single aPBP molecules displaying periods of transient immobility <sup>14,56</sup> ,
290	their synthesis may be localized to small (<50 nm) regions. Combined, these observations support a model where aPBPs
291	synthesize material to fill gaps in the PG meshwork <sup>36,14</sup> . Gaps could arise via mechanical damage, hydrolases <sup>55</sup> , or
292	between the imperfectly oriented strands built by the Rod complex <sup>11</sup> . If this model is correct, the different spatial activities of
293	these two systems might allow the sacculus to maintain integrity at any Rod/aPBP ratio: the fewer the Rod complexes, the
294	larger the gaps filled by aPBPs.
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296	While many different activities affect the shape of the sacculus, such as its cleavage by hydrolases or rigidification by wall
297	teichoic acids <sup>11</sup> , the first process defining its geometry is the spatial coordination controlling where glycans are inserted into

it. While these experiments give a coarse-grained description into how each synthetic system affects cell shape, a fine

increased Rod activity make cells thinner? Understanding the physical mechanisms causing these changes will require not

downstream of glycan insertion affect the shape and mechanics of sacculi as they subsequently modify, remodel, and break

only a better understanding of the molecular architecture of the sacculus, but also an investigation into how the enzymes

scale, mechanistic understanding remains to be determined - How does aPBP activity make cells wider? How does

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down the nascently-polymerized glycan architecture.

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#### 314 Author Contributions.

- 315 B. subtilis strains were cloned by MD, YS, and MK. All width and bulk growth measurements of B. subtilis were done by MD,
- 316 and E. coli widths by YS. E. coli CRISPRi strains were cloned by AV, who was supervised by SVT. Single cell growth rates
- 317 were done by YS and MK. TIRFM and tracking of PBP2A was done by MK. All TIRFM of MreB was done by YS. TIRF-SIM
- 318 of MreB was done by YS, EG, and JR. Purified sacculi and proteomic mass spec sample preps were done by MD. YS wrote
- 319 the code for analysis of single cell growth rate, filament density, and simulations of data. Polarization microscopy and
- 320 analysis was conducted by JR, RO, and EG. SW did the FDAA synthesis, osmotic shocks, and TEM. The paper was written
- 321 by EG, MK, MD, and YS.
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#### 429 Figure 1 – Rod width depends on the relative levels of the widening aPBPs to the thinning 430 Rod system. B

Rod complex

PBP

PBP2A

(pbpA)

aPBF

PBP1

(ponA)

A

431 432 Error bars are SD of the mean.

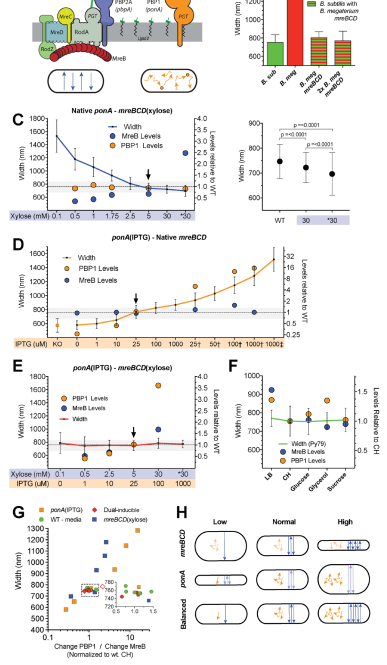
433 434 A. Schematic of the two PG synthetic systems 435 responsible for growth. The Rod complex 436 (depicted on the left) polymerizes glycans via the 437 peptidoglycan glycosyltransferase (PGT) activity of 438 RodA. These glycans are crosslinked into the 439 sacculus by the transpeptidase (TP) activity of 440 PBP2A. aPBPs are depicted on the right, 441 containing both PGT and TP activity. Below -442 Schematic of each system's in vivo motions. Rod 443 complexes move circumferentially around the rod 444 as they synthesize PG. In contrast, aPBP synthesis 445 does not appear organized, as they diffuse along 446 the membrane, occasionally halting and remaining 447 immobile for 1-3 seconds. Protein names are 448 capitalized, gene names italicized. 449

450 B. B. megaterium mreBCD in B. subtilis forms 451 rods close to B. subtilis width. "B. sub" indicates 452 WT B. subtilis. "B. meg" indicates B. megaterium. 453 Checkered boxes are bMD465 (amyE::Pxyl-454 mreBCD. minCD [B. megal::erm. mreBCD. 455 minCD::spec [B. mega]), a B. subtilis strain where 456 the native mreBCD locus is replaced with B. 457 megaterium mreBCD, and an additional B. 458 megaterium mreBCD is under xylose control at an 459 ectopic locus. "B. meg mreBCD" indicates bMD465 460 grown with 1% glucose to repress expression from 461 the additional copy. "2x B. meg mreBCD" indicates 462 bM465 growth with 30 mM xylose to overexpress 463 B. megaterium mreBCD from the additional copy. 464 See Figure S1A for a zoomed view. 465

#### 466 C-F. Titrations of ponA and mreBCD vs. cell

467 width. Each strain was grown in the inducer 468 concentrations shown below the graph. Mean width 469 is plotted on the left axis. PBP1 and MreB relative 470 protein abundances (determined by mass 471 spectrometry and normalized to the levels in WT 472 Py79 grown in CH) are plotted on the right axis.

- 473 Arrowhead indicates inductions producing WT
- 474 widths and cellular levels of induced protein.
- 475 Dashed line depicts mean steady state diameter of
- 476 Py79 in CH, with SD shown as gray shaded area.
- 477



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🔲 B. subtilis

B. megateriun

B. subtilis with
 B. megateriun

478 C. Cell diameter decreases with mreBCD induction. All data are from strain bMD545 (amyE::Pxyl-mreBCD::erm

479 AmreBCD::spec), except for the induction marked \* which are bMK355 (amyE::Pxyl-mreBCD::erm) containing a xylose-inducible 480 mreBCD in addition to the native mreBCD. Right - Zoomed view of the right two bars compared to WT. P-value calculated with

- 481 Mann-Whitney.
- 482

483 **D. Cell diameter increases with ponA induction**. All data are from strain bMD598 (yhdG::Pspank-ponA::cat \DeltaponA::kan), except 484 for the inductions marked + and + which are under stronger promoters than bMD598. + is bMD586 (vhdG:Phyperspank-ponA::cat 485 Δ*ponA::kan*), and ‡ is bMD554 (*yhdG::Phyperspank-ponA::cat*) which has an inducible *ponA* in addition to the native copy. 486

#### 487 E. A balanced expression of both PG synthetic systems yields normal width rods across a large range.

488 Strain bMD620 (amyE::PxyI-mreBCD::erm \DeltamreBCD::spec yhdH::Pspank-ponA::cat \DeltaponA::kan) grown in combinations of 489 inducers shown along the bottom. \* indicates bMD622 (amyE::PxyI-mreBCD::erm yhdH::Pspank-ponA::cat  $\Delta ponA::kan$ ) containing 490 a xylose-inducible mreBCD in addition to the native mreBCD. 491

492 F. B. Subtilis maintains a constant width in different media. Mean and SD of cell diameters of B. subtilis strain Py79

493 (exponential phase) grown in different growth media (media where a carbon source is indicated is S750). Width is plotted on the 494 left axis. PBP1 and MreB relative protein abundances (determined by mass spectrometry, normalized to the levels in CH media) 495 are plotted on the right axis. 496

497 G. WT width is maintained within a narrow range of relative PBP1/MreB ratios. Plotted are the widths of all cells in Figures 498 1C-F, against the ratio of the fold change in PBP1 to that of MreB. Inset is a zoomed view of the dotted box, showing the widths of 499 cells within a 0.5-1.5 PBP1/MreB range. Green indicates ratios in different media. Orange squares indicate ratios in ponA inducible 500 strains. Blue squares indicate ratios in mreBCD inducible strains. Red diamonds indicate ratios in the dual-induction strain, with the 501 open red diamond indicating the ratio when protein levels exceeded WT.

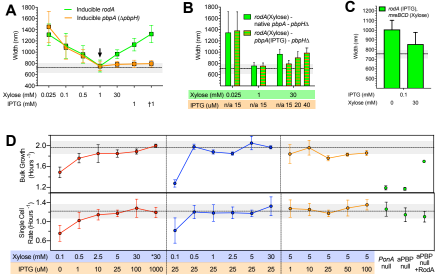
503 H. Model for how the two PG synthesis systems affect rod width. Top – As the amount of circumferentially organized 504 synthesis increases (blue straight lines), cell diameter decreases. Center - In contrast, as the level of unorganized synthesis 505 increases (orange, non-directed lines), so too does cell diameter. Bottom - As long as the activities of the unorganized (orange) 506 and circumferentially organized (blue) syntheses are balanced, cell width can remain constant, even across a range of protein 507 levels.

#### 509 Figure 2 – Effects of RodA/PBP2 on cell width, and how each PG synthetic system affects 510 growth.

511 В 512 1800 1800 A. As rodA or pbpA induction is Α Inducible rodA 1600 Inducible pbpA (ApppH) 1600 513 increased, the mean diameter of cells 1400 1400 (mu 514 Ĩ decreases up to a point, beyond which 1200 1200 515 ₽ 1000 Nidth 1000 it increases with rising rodA induction, 516 800 800 but not for *pbpA*. Green line represents 600 600 517 diameters of bMD592 (Pxyl-rodA::erm) at 400 518 Xvlose (mM) 0.025 0.1 five different xylose concentrations, save 0.5 1 30 IPTG (mM) †1 519 bMD580 (yhdG::Phyperspank-rodA::cat 520 ∆rodA::kan) and bMD556 521 522 523 524 (yhdG::Phyperspank-rodA::cat - labeled †) D 2.0 which were induced with IPTG. Orange Bulk Growth sinoH) line represents the diameters of bMD597 (Pxyl-pbpA::erm ∆pbpH::spec) at five 1.2 525 different xylose concentrations, save 1.4 526 -0.1 Hours bMD574 (yhdG::Phyperspank-pbpA::cat Single Cell 527 528 Δ*pbpA::erm* Δ*pbpH::spec*) and bMD573 (vhdG::Phyperspank-pbpA::cat 529 △*pbpH::spec* - labeled †) which were 2.5 0.5 0.5 5 30 \*30 0.1 Xvlose (mM) 0.1 1 530 induced with IPTG. IPTG (uM) 0 10 25 100 1000 25 531 532 533 534

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B. Overexpression of rodA increases cell diameter, but only when pbpA expression is also sufficiently high. Light green bars represent the steady state diameter of bMD627 (Pxyl-rodA::erm \DeltappH::spec) induced at the specified concentrations of xylose. Green + orange striped bars represent the steady state diameter of bMD631 (Pxyl-rodA::erm yhdG::Pspank-pbpA::phleo 535 *Apph::spec*) induced at the specified concentrations of xylose and IPTG. 536

#### 537 C. The increase in cell diameter caused by overexpression of rodA is reduced by simultaneous overexpression of

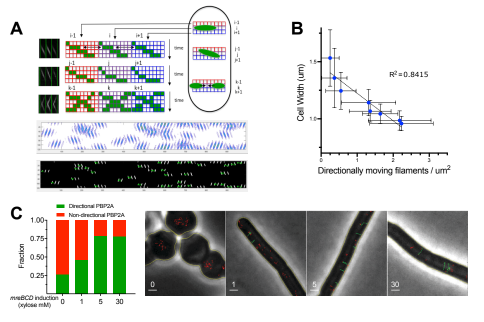
*mreBCD*. bMD557 (*yhdG*::Phyperspank-*rodA*::*cat amyE*::Pxyl-*mreBCD*::*erm*) was induced at the indicated levels of IPTG and xylose.

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541 **D. Rates of cell growth measured at the population (***Top***) and single cell (***Bottom***) level. Mean rates of cell growth were
542 measured either by (***Top***) optical density at 600 nm in a shaking plate reader in CH, or (***Bottom***) by microscopically assaying the
543 rate single cells (grown under an agar CH pad) added surface area. All measures are from bMD620 grown in the inducer
544 concentrations indicated along the bottom, with the exception of: "***ponA* **null", which is bMK005 (***AponA::cat***), "aPBP null", which is
545 bAM268 (***ApbpF, ApbpG, ApbpD, AponA::kan***), and "aPBP Null + RodA", which is bAM288 (***ApbpF ApbpG ApbpD AponA:kan***546** *amyE::***Phyperspank-***rodA***-***His10::spec***), where RodA is induced with 0.05 mM IPTG. Dotted horizontal lines represent the mean
547 growth rate of Py79 in CH medium, with the shaded area representing the SD.** 

# Figure 3 – Increased *mreBCD* increases directional MreB filament density and the fraction of directional PBP2A molecules.

A. Schematic of the method to quantitate directionally moving MreB filaments. Data shown is from simulated TIRFM time-

553 lapses, with a 1 sec exposure time and 554 65 nm/pixel magnification. First, a 555 kymograph is generated for each row 556 of pixels along the midline of the cell. 557 These kymographs are then lined up 558 side by side to generate a single 2D 559 image, where each column of the 560 image contains a kymograph of each 561 sequential row of pixels in the cell. This 562 image is adaptively thresholded, then 563 segmented with contour analysis to 564 extract all fluorescent objects (middle). 565 These objects are used to get velocity 566 (slope), time (centroid), and position 567 (row) for each particle. As each particle 568 will show similar intensities in adjacent 569 rows, or sometimes move at an angle, 570 objects up to two rows apart are 571 grouped based on time, position, and 572 velocity. This yields the final particle 573 count (bottom). See Figure S3 for 574 further details and tests. 575



**B.** The decreased rod width caused by increasing *mreBCD* induction correlates with an increasing density of directionally **moving MreB filaments.** bYS981 (*amyE*::PxyI-*mreB-SW-msfGFP, mreCD::erm ΔmreBCD::spec*) was grown in different amounts of xylose. Plotted are the steady state mean cell widths against the density of directionally moving filaments (determined as in **A**, above). All error bars are SD of the mean.

581 C. The fraction of directionally moving Halo-PBP2A molecules increases with mreBCD expression. mreBCD was induced at 582 the indicated levels in bMK385 (amyE::Pxyl-mreBCD::erm ΔmreBCD::spec HaloTag-11aa-pbpA::cat), and the single molecule 583 motions of JF-549-labeled Halo-PBP2A were imaged by TIRFM using 300 msec continuous exposures. Plotted are the fractions of 584 labeled PBP2A trajectories over 7 frames in length that moved directionally. *Right* - Representative montage of Halo-PBP2A 585 trajectories at different levels of mreBCD inductions overlaid on phase images. Directionally moving tracks are labeled green; all 586 other tracks are labeled red. Scale bar is 1 μm. See also corresponding Movie S3.

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#### Figure 4 - Increased Rod activity increases both the amount of oriented material within 589 590 sacculi and their mechanical anisotropy.

592 All error bars are SD of the mean.

#### 593 594 A. Polarization microscopy

#### 595 reveals oriented material within

596 the cell wall. Left - LC-PolScope 597 images of purified WT Py79 sacculi. 598 Focused at the surface, the cell wall 599 is seen to be birefringent. Color 600 indicates the orientation of the slow 601 axis, with intensity corresponding to 602 retardance in that direction. 603 Orientation key is the color wheel at 604 upper left. *Right* - Polarization 605 orientation view of the sacculi

- 606 surface, with lines pointing in the 607 predominant orientation of the slow
- 608 axis. See also Movie S2.

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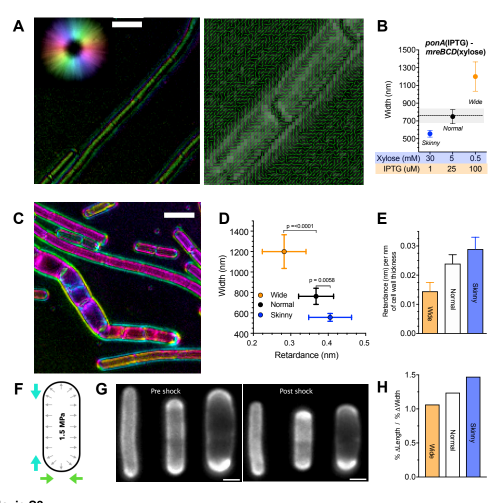
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#### 610 B. Induction conditions used to 611 assay sacculi properties. bMD620

612 (amyE::PxyI-mreBCD::erm

613 ΔmreBCD::spec yhdG::Pspank-ponA::cat 614 △ponA::kan) was induced to grow at 3 615 different widths: Skinny (30 mM 616 xylose, 1 µM IPTG), Normal (5 mM 617 xylose, 25 µM IPTG), and Wide (0.5 618 mM xylose, 100 µM IPTG). 619

620 C. Example LC-PolScope image of bMD620 sacculi induced at different 621 622 623 widths. For each experiment, pairs of purified sacculi ("Wide" and 624 "Normal", or "Wide" and "Skinny") 625 were combined, and Z-stacks were 626 collected in 100 nm steps. See also Movie S3.



D. The amount of oriented material in the cell wall increases with mreBCD induction, and inversely correlates with rod width. Plotted is the mean retardance vs. width along the length of projected Z-stacks of at least 90 different cells for each induction condition.

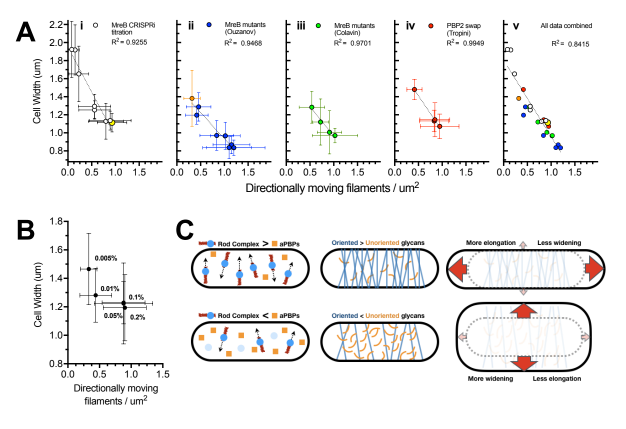
E. Mean retardance for each sample normalized to the thickness of the cell wall, as measured by TEM.

F. Schematic of osmotic shock assay of anisotropy. B. subtilis sacculi are normally under tension due to the high internal turgor (black arrows). Hyperosmotic shock negates this pressure, allowing observation of how sacculi shrink in length and width (colored arrows).

638 G. Example FDAA-labeled cells from each condition (left to right: Skinny, Normal, Wide) before and after osmotic shock. Scale bar 639 is 1 μm. 640

641 H. As the relative amount of Rod system activity increases, so does the mechanical anisotropy of the sacculus. Anisotropy 642 (% change in length / % change in width) for each induction condition following osmotic shock. Raw data and reduction for each 643 dimension (both absolute and percent) are in Figure S4.

# Figure 5 – Directional MreB filament density also correlates with cell width of *E. coli* Rod mutants, suggesting a generalizable model for the effects of Rod complex on cell width.



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### A. Cell width vs. density of directionally moving MreB filaments in different *E. coli* strains. (i) AV88 (186::Ptet-

dCas9, mreB::msfGFP-mreBsw), allows the tunable expression of mreB-msfGFPsw by expressing various sgRNAs with different matches against msfGFP. Yellow indicates WT expression. (ii) mreB-SW-msfGFP mutant strains from Ouzounov et al., 2016.
 Orange indicates RM478 (ΔrodZ, mreB(S14A)-msfGFPsw) from Morgenstein et al., 2015. (iii) mreB-msfGFPsw mutants believed to change filament curvature from Colavin et al., 2018. (iv) msfGFP-mreB strains from Tropini et al., 2015, where mrdA is replaced with mrdA from other species. (v) All data from (i)-(iv) combined. See also Movie SM4. All values are detailed in Table S1.

### B. Decreased cell width caused by increased RodZ expression correlates with an increased density of directionally

6 moving MreB filaments. KC717 (*csrD::kan, mreB::msfGFP-mreB, ProdZ* (*frt araC PBAD*)) was grown at different arabinose 7 concentrations (indicated on the graph), and filament density was calculated as in Figure 3B. See also Movie SM5.

659 **C. Model for how the balance between aPBPs and the Rod system affects cell width.** *Left* – When Rod complex activity (perhaps 660 set by the number of MreB filaments) is high relative to that of aPBPs, sacculi have more circumferentially oriented material (*Center*) 661 compared to when aPBP activity is greater. *Right* – As the amount of oriented material increases, the less elastic sacculi are more 662 rigid across their width, but less rigid along their length. Being stretched by the internal turgor pressure, sacculi with greater Rod 663 activity are better able to maintain their width, while stretching more along their length. In contrast, cells with reduced Rod activity 664 have less circumferentially oriented glycans to reinforce their width, and thus expand more along their width.

#### Supplemental Material for: 666

667

#### Cell Diameter in Bacillus subtilis is Determined by the Opposing Actions of Two Distinct 668 **Cell Wall Synthetic Systems** 669

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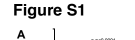
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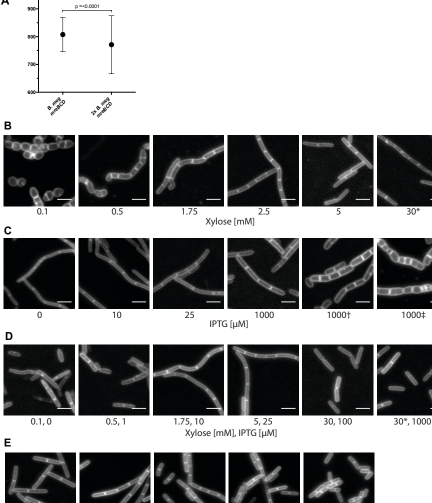
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A. Zoomed view of the two rightmost distributions of widths in Figure 1B. Error bars are SD of the mean.

CH

S750

glucose

**B-D.** Cells were grown in CH media with the indicated concentrations of inducers during steady state growth, then stained with 0.5  $\mu$ g/mL FM 5-95. Scale bars are 4  $\mu$ m.

S750

sucrose

**B.** Images are of bMD545 (*amyE::PxyI-mreBCD::erm*, Δ*mreBCD::spec*), save the induction annotated \* which is bMK355 (*amyE::PxyI-mreBCD::erm*), containing a xylose inducible *mreBCD* in addition to the native *mreBCD*.

S750

glycerol

**C.** Images are of bMD598 (*yhdG::Pspank-ponA::cat, ΔponA::kan*), induced with the indicated amounts of IPTG, save the inductions marked † and ‡ which are under stronger promoters than bMD598; † is bMD586 (*yhdG*::Phyperspank-*ponA*::*cat*, *ΔponA::kan*), and ‡ is bMD554 (*yhdG*::Phyperspank-*ponA*::*cat*), containing an inducible *ponA* in addition to the native copy.

D. Images of bMD620 (*amyE*::PxyI-*mreBCD*::*erm*, Δ*mreBCD*::*spec*, *yhdH*::Pspank-*ponA*::*cat*, Δ*ponA*::*kan*) grown in CH, and induced with the indicated amounts of IPTG and xylose. The image annotated \* indicates bMD622 (*amyE*::*PxyI-mreBCD*::*erm yhdH*::*Pspank-ponA*::*cat* Δ*ponA*::*kan*) containing a xylose-inducible *mreBCD* in addition to the native *mreBCD*.

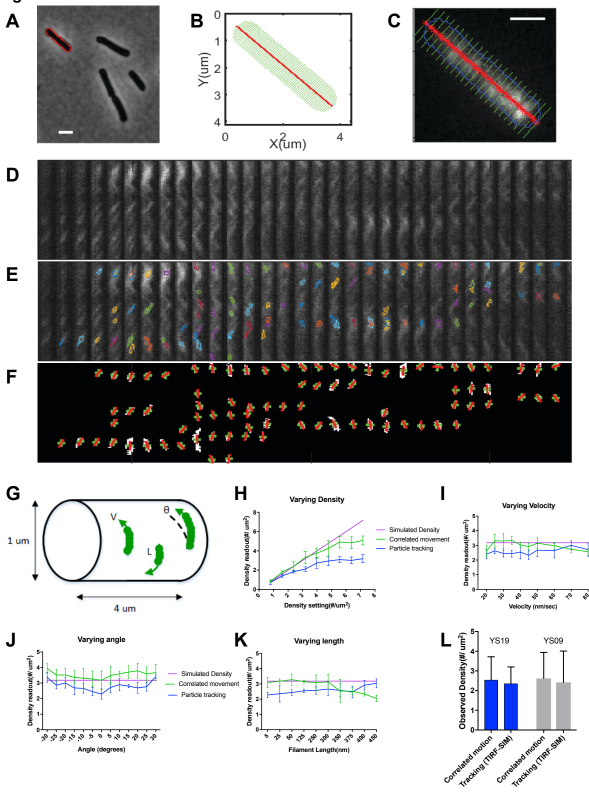
 $694 \\ 695 \\ 696 \\ 697 \\ 698 \\ 699 \\ 700 \\ 701 \\ 702 \\ 703 \\ 704 \\ 705 \\ 706 \\ 707 \\ 708 \\ 709$ 

LB

710 E. Representative images of Py79 grown in indicated media, then stained with 0.5 μg/mL FM 5-95. Scale bars are 4 μm.

711

## 712 Figure S3



714 A-F. Expanded description of the method used to determine directional filament density. Cells expressing fluorescent MreB 715 fusions were imaged with TIRF for 120 seconds, followed by a single image in phase-contrast. From the phase-contrast images  $\begin{array}{c} 716\\ 717\\ 718\\ 719\\ 720\\ 721\\ 722\\ 723\\ 724\\ 725\\ 726\\ 727\\ 728\\ 729\\ 730\\ 731\\ 732\\ 733\\ 734 \end{array}$ ((A) - scale bar is 2 um), cells were segmented to determine their dimensions and midline (B). The resulting segmentation mask was used to create kymographs out of the fluorescence time-lapse data. (C) Shows the mask overlaid on a maximum intensity projection of a TIRF time-lapse (red line indicates midline, green lines are the axis along which kymographs are drawn, scale bar is 2 um). Next, kymographs were created for each sequential row of pixels along the midline, and the kymographs from each row were displayed side by side as in (D). Next, objects within these kymographs were identified and segmented based on contour analysis (E), using adaptive thresholding. For each identified object, a set of properties [X, t, v, I] were measured. (X, t) are the centroid of the object in the binary image (green crosses in (F)). X is the position of the filament along the cell width, and t is time, corresponding to rows in the kymographs. Velocity (v) is calculated based on the diagonal slope of the object in the binary image (red lines in (E)), and can be positive or negative depending on the direction of motion. Intensity (I) is the sum of the intensities of all pixels in the object. Because each kymograph represents one row of pixels in the cell, and the signal from each particle can span more than one pixel (65 nm) in width, the same filament can appear in several sequential kymographs (or rows in the original image). In order to link the objects occurring in multiple rows together, each object in a given row [Xi] can be linked to corresponding objects in the next row  $[X_{i+1}]$  or previous row  $[X_{i+1}]$  based on their time and velocity. Objects are only linked if they have a velocity v + -20 nm/sec of each other, and their peak time t is within 2 seconds; these objects are then counted as a single filament. This also allows us to capture filaments moving at angles of up to +/- 30 degrees to the midline. Finally, the number of counts occurring during the imaging interval is normalized based on the cell width and length (or total surface area  $\pi^*$  width\*length).

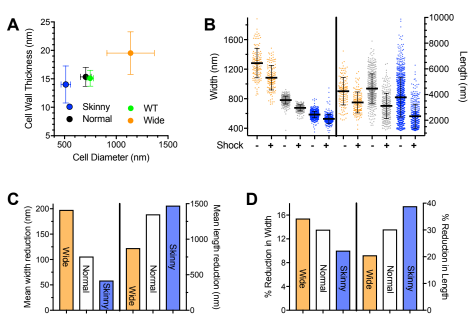
#### G-J. Evaluation of filament density quantitation tested using simulations.

(G) Schematic of the MreB data simulations. Simulations were run using custom MATLAB code, with the following parameters: 735 736 737 738 739 740 Maximum fluorescence intensity was 300 counts and noise was 10 (approximating the signal to noise conditions in our imaging). The simulated cell had a width of 1 µm and a length of 4 µm. We tested the robustness of our tracking by varying one parameter while the others were fixed at the following values: Filament density was 3.2/µm<sup>2</sup> (corresponding to 40 MreB filaments per cell), filament velocity was 30 nm/sec, filament orientation was 0 degrees, and filament length was 250 nm. This data was tracked using the correlated motion method detailed above. For comparison, we tracked the same simulated data with the linear motion LAP tracker in Trackmate, with a low bar for calling a trace a directional motion (tracks longer than 6 frames with a displacement greater 741 742 than 250 nm). (H) Filament detection performance with respect to different filament density settings (from 0.8 to 7.2/um<sup>2</sup>, corresponding to 10 - 90 MreB filaments per cell). (I) Filament detection with respect to filament angles (from -30 to 30 degrees). 743 744 (J) Filament detection with respect to different filament velocities (from 20 to 90 nm/sec). (K) Filament detection with respect to different filament lengths (5 nm to 450 nm). 745

#### 746 L. Correlated motion analysis yields a similar density of directional filaments as tracking of MreB filaments imaged by

747 TIRF-SIM. For this comparison, two different strains with natively expressed MreB fusions, YS09 (mreB-mNeonGreen) and YS19 748 (mreB-SW-msfGFP), were grown under the same conditions, and were analysed using two different imaging/analysis pipelines: 1) 749 As above, standard TIRFM imaging followed by correlated motion analysis, and 2) TIRF-SIM imaging, followed by tracking 750 filaments with particle tracking (using the same settings as in (G) above). 751

### 754 Figure S4



**A.** Cells, induced under identical conditions to **Figure 4B**, were fixed and stained with uranyl acetate, the block was cut in ultrathin sections, then imaged with transmission electron microscopy (TEM). The thickness of the cell wall was measured and plotted against cell diameter (also determined via TEM).

**B.** All data of the width (left) and length (right) of cells before and after osmotic shocks (orange = Wide, grey = Normal, blue = Skinny). Error bars are SD of the mean.

C Absolute and D percent reduction in width and length of labeled sacculi under each induction condition after osmotic shock.

# Supplemental Movie Legends.

**Movie S1** – Example single molecule movies of PBP2A-HaloTag (labeled with JF549<sup>1</sup>) at different *mreBCD* induction levels in bMK385 (*amyE::Pxyl-mreBCD*::*erm*, Δ*mreBCD*, HaloTag-11aa-pbpA::cat). Xylose concentrations are indicated in each panel. Frames are 300 msec apart.

**Movie S2** – Z-stack of LC-PolScope images of sacculi purified from wild type Py79. (Left) Z-stack of the retardance of each plane. (Right) Color map of birefringence at each plane, with color wheel at upper left giving orientation reference. Z-steps were taken in 100 nm increments.

Movie S3 – Z-stack color map of birefringence of sacculi purified from strain bMD620 (*amyE::Pxyl-mreBCD::erm, ΔmreBCD, yhdG::Pspank-ponA::cat, ΔponA::kan*), where cells were induced to "Wide" (0.5 mM xylose, 0.1 mM IPTG) and
 "Normal" widths (5 mM xylose, 0.025 mM IPTG). Color wheel at upper left indicates orientation direction. Z-steps were taken
 in 100 nm increments.

Movie S4 – TIRF-SIM movies of different mreB-SW-msfGFP mutants. Cells were grown in LB at 37°C, then placed under an LB agar and imaged at 37°C. Frames are 1 sec apart. Scale bars are 1 um.

Movie S5 – TIRF-SIM movies of strain KC717(*mreB::msfGFP-mreB, ProdZ* (*frt araC PBAD*)). Cells were grown for 5
 hours in LB with the indicated arabinose concentrations, were placed under an LB agar pad and imaged at 37°C. Frames
 are 1 sec apart. Scale bars are 1 um.

# 788 Materials and Methods.

### 789 Media and culture conditions

For all experiments, unless otherwise noted, *B. subtilis* and *B. megaterium* were grown in casein hydrolysate (CH) medium (where indicated, xylose and/or isopropyl thiogalactoside (IPTG) was added), and *E. coli* strains were grown in lysogeny broth (LB) medium (where indicated, arabinose or anhydrotetracycline (ATc) was added), at 37°C with rotation. When pre-cultures reached exponential phase the next day, after overnight growth at 25°C, they were further diluted into fresh growth medium (and where indicated, with the specified concentrations of inducer) and were grown for at least 3 hours at 37°C with rotation, to an OD<sub>600</sub> of ~0.3 to 0.7.

## 797 Sacculi purification

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798 20 mL cultures were grown in 250 mL baffled flasks at 37°C with vigorous shaking, to an OD<sub>600</sub> of ~0.5, were harvested and were centrifuged at 5,000 × g for 5 min at 4°C. Cell pellets were resuspended in 1 mL of ice-cold 799 800 phosphate buffered saline (PBS), were centrifuged at 6,000  $\times$  g for 30 sec, were resuspended in 500  $\mu$ L of PBS, and 801 were killed by boiling in a water bath for 10 min. Cells were centrifuged at  $6,000 \times q$  for 2 min, were resuspended in 802 500 µL of PBS containing 5% sodium dodecyl sulfate (SDS), and were boiled in a water bath for 25 min; this was 803 repeated once, except boiling was for 15 min. To remove the SDS, the samples were centrifuged at  $6,000 \times q$  for 2 804 min, and were resuspended in 500  $\mu$ L of PBS; this was repeated 5 times. The cells were centrifuged at 6,000 × g for 2 805 min, were resuspended in 1 mL of 50 mM Tris-HCl, pH 7.5, containing 10 mM NaCl, and 2 mg of pronase from 806 Streptomyces griseus (MilliporeSigma, MA), and were incubated at 60°C for 90 min with gentle shaking. To remove 807 the pronase, the samples were boiled twice in PBS/5% SDS, followed by 6 rounds of washes in PBS, exactly as 808 described in the steps above that precede the pronase treatment. The samples were centrifuged at 6,000  $\times$  g for 2 809 min, were resuspended in 1 mL of 25 mM Tris-HCl, pH 8.5, containing 0.5 M NaCl, 5 mM MgCl<sub>2</sub>, and 100 U of salt-810 active nuclease (SAN) (ArticZymes, Norway), and were incubated at 4°C overnight with gentle mixing. To remove the 811 SAN, the samples were boiled twice in PBS/5% SDS, followed by 6 rounds of washes in PBS, exactly as described in 812 the steps above that precede the pronase treatment. The sacculi were centrifuged at  $6,000 \times q$  for 2 min, the 813 supernatant was removed, and the pellets were stored at -80°C.

### 815 **Polarization microscopy**

816 Purified sacculi were resuspended in PBS and placed on ethanol-cleaned No. 1.5 coverslips under a cleaned glass

817 slide. Polarization images were acquired on an inverted LC-PolScope mounted on a Nikon Ti-E equipped with a

60x/1.4NA Plan Apo oil immersion objective and oil condenser with matching NA, and a Hamamatsu Photonics

Flash4 camera. Z-stacks in 100 nm steps were taken of each sample using 50 msec acquisitions of 546/12 nm light. All image acquisition, processing, and display, including colored display and line maps were prepared using the

821 OpenPolScope Hardware Kit and plugins for ImageJ/Micro-Manager from OpenPolScope.org.

# 822823 Calculation of retardance

Z-stacks of the computed total retardance for each slice were exported from OpenPolScope. These stacks were cropped at one frame above and beneath the focal planes of the top and the bottom of the cells, then projected into a single plane using ImageJ. To avoid getting high retardance values from the edge effects arising from the sides of the cell or the septa, we selected the retardance at the middle of the cell, using the average of line scans (5 pixels long and 3 wide) that were drawn down the center of cells, taking care to avoid edges and septa.

## 830 Transmission electron microscopy of cell wall thickness

831 Overnight cultures were diluted into fresh medium and grown to an OD<sub>600</sub> of ~0.3 to 0.5. Cells were pelleted by

- centrifugation at 5,000 x g and fixed by resuspending in 100 mM MOPS pH 7 containing 2% paraformaldayde, 2.5%
- gluteraldehyde, and 1% DMSO overnight at 4°C. Cells were centrifuged at 5,000 x g and were washed 3 times with
- 100 mM MOPS pH 7. The pellet was stained with 2% osmium tetroxide in 100 mM MOPS for 1 hr, washed twice with
- deionized water, and stained overnight with 2% uranyl acetate. The pellet was washed twice with deionized water and dehydrated by washing once with 50% ethanol, once with 70% ethanol, once with 95% ethanol, and then three times
- with 100% (v/v) ethanol. Samples were prepared for resin infiltration by washing once with 50% ethanol, 50%
- propylene oxide (v/v) and then once with 100% propylene oxide. All wash steps were for 5 min. Infiltration of resin was

achieved by incubation with 50% Embed 812 (EMS, PA)/50% propylene oxide for 1 hr, followed by incubation with

67% Embed 812/33% propylene oxide for 1hr, and incubation with 80% Embed 812/20% propylene oxide for 1 hr.
Samples were then incubated twice with Embed 812 for 1 hr, followed by an overnight incubation in molds. The molds

- 842 were baked at 65°C for 18 hr before sectioning.
- 843

Serial ultrathin sections (80 nm) were cut with a Diatome diamond knife (EMS, PA) on a Leica Ultracut UCT (Leica
 Microsystems, Germany) and collected on 200-mesh thin-bar formvar carbon grids. Sections were imaged on an FEI
 Tecnai transmission electron microscope.

847

848 Cell wall thickness measurements were performed using a custom-built MATLAB (Mathworks, MA) script. Image

intensity profiles extracted from lines were drawn perpendicular to a user-input line defining the middle of the cell wall.

The distance between the two lowest points below a threshold within 40 nm of the middle of the cell wall was

measured as the cell wall thickness at ~30 points in each cell.

# 853 Measurements of cell diameter

Cultures grown to an OD<sub>600</sub> of ~0.3 to 0.7 were stained with 0.5  $\mu$ g/mL FM 5-95 (Thermo Fisher, MA) for 1 min, and were concentrated by centrifugation at 6,000 x *g* for 30 sec. The cell pellet was resuspended in ~1/20 volume of growth medium, and 3  $\mu$ L was applied to ethanol-cleaned No. 1.5 coverslips under a 3% agarose pad containing growth medium. Fluorescent cells were imaged with the top surface of the agarose pad exposed to air, in a chamber heated to 37°C. Epifluorescence microscopy was performed using a Nikon Eclipse Ti equipped with a Nikon Plan Apo  $\lambda 100 \times /1.4$ NA objective and an Andor camera. Cell contours and dimensions were calculated using the

860 Morphometrics software package <sup>2</sup>. 861

## 862 Measurements of single-cell growth rate

Cultures grown to an  $OD_{600}$  below 0.3 were concentrated by centrifugation at 6,000 x *g* for 30 sec. The cell pellet was resuspended in growth medium, and applied to No. 1.5 glass-bottomed dishes (MatTek Corp., MA). All cells were imaged under a 2% agarose pad containing growth medium, with the top surface exposed to air, in a chamber heated to 37°C. Phase-contrast microscopy was performed using a Nikon Eclipse Ti equipped with a Nikon Plan Apo  $\lambda$ 100×/1.4NA objective and an Andor camera. We used a custom-built package in MATLAB to perform segmentation on phase-contrast time-lapse movies, then calculated the growth rate of the surface area of single *B. subtilis* chains.

## 870 Measurements of bulk growth rate

For cell culture measurements of growth rate, overnight cultures were diluted in fresh medium and grown for ~3 hr at 37°C to an OD<sub>600</sub> of ~0.3 to 0.7. The cultures were diluted back to a calculated OD<sub>600</sub> of 0.07 in 100-well microtiter plates, and their growth rates were measured in a Bioscreen-C Automated Growth Curve Analysis System (Growth Curves USA, NJ) plate reader, at 37°C with continuous shaking. Growth rates were calculated from OD<sub>600</sub> measurements that were recorded every 5 min for at least 6 hr.

# 876877 TIRF microscopy of Halo-PBP2A

Cultures grown to an  $OD_{600}$  below 0.3 were labelled with 10 nM JF549<sup>3</sup>, and were concentrated by centrifugation at 6,000 x g for 30 sec. The cell pellet was resuspended in growth medium and applied to ethanol-cleaned No. 1.5 coverslips. All cells were imaged under a 2% agarose pad containing growth medium with the top surface exposed to air, in a chamber heated to 37°C. TIRFM and phase-contrast microscopy were performed using a Nikon Eclipse Ti equipped with a Nikon Plan Apo  $\lambda$  100×/1.45NA objective and a Hamamatsu ORCA-Flash4.0 V2 sCMOS camera. Fluorescence time-lapse images were collected by continuous acquisition with 300 msec exposures.

884

## 885 Analysis of the density of MreB

886 Phase images of bacteria were segmented using Morphometrics <sup>4</sup>, and the width and length of each cell was

- calculated. Next, the fluorescence time-lapses were analyzed based on the segmentation mask of the phase image
- 888 (Figure S3C). Filament counting was performed in several steps (Figure S3). First, kymographs were generated for
- each row of pixels along the midline of the cell. Next, the kymographs for each row were placed side by side,
- converting the TIRF time lapse data into a single 2D image (**Figure S3D**). To identify filaments in the kymograph,
- 891 closed contours were generated in the 2D image (Figure S3E). We only selected contours within a given size range

892  $(0.04 \ \mu m^2 to 0.17 \ \mu m^2)$ . For these contours, we calculated the total intensity (the sum of the intensities of the pixels in 893 the contour), the centroid, the velocity (calculated from the slope of the major axis line of the contour) (Figure S3F), 894 and time (from the centroid). Next, to identify cases where the same MreB filament appears in multiple sequential 895 kymographs, each object in a given kymograph is linked to a corresponding object in the next and previous 896 kymographs based on the above properties of the object (see Figure S3F for details). Finally, the counting is verified 897 manually by numbering each filament on the 2D image (Figure 3A). To test the performance of the filament counting, 898 we analyzed simulated data with different filament density, velocity, and orientation settings (Figure S3). All of the 899 image analyses were performed using MATLAB.

900

## 901 Simulation of directionally moving MreB

902 The Image Correlation Spectroscopy <sup>5</sup> MATLAB package was used for the simulation of MreB moving around the cell. 903 The following parameters were set for the MreB simulations: velocity, orientation, filament number and filament length. 904 The default velocity setting is 30 nm/sec and the default orientation is 0, which means MreB moves perpendicular to 905 the central axis. The default filament length is set to 250 nm and each MreB monomer is assumed to be 5 nm. The 906 cell width is set to 1 um and the cell length is set to 4 µm. The pixel size is 65 nm and the time interval is 1 sec, which 907 is the same as the TIRF imaging obtained with our Nikon Eclipse Ti equipped with a Hamamatsu ORCA-Flash4.0 V2 908 sCMOS camera. The particles are randomly distributed on the surface of the cell. Each simulation data point was 909 repeated 5 times for the counting analysis. To compare the correlated motion approach against particle tracking, we 910 counted the number of tracks observed after tracking the simulated data using the Linear Motion LAP tracker in FIJI 6 911 with TrackMate v3.8<sup>7</sup>. The threshold for spot size was 0.195 µm and the intensity threshold was 10 counts. The 912 search radius is 0.085 µm, the link radius is 0.085 µm, and the gap size is 1. All traces longer than 6 frames that had 913 moved more than 250 nm were considered directional motions.

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## 915 TIRF-SIM imaging of *E. coli* strains

Cells were prepared as described in "Media and culture conditions". Cells were placed under an LB agarose pad, on a cleaned No.1.5 coverslip, and imaged at 37°C. Live-cell SIM data were acquired as described previously <sup>8</sup> on a Zeiss Axio Observer.Z1 inverted microscope outfitted for structured illumination. An Olympus 100×/1.49NA objective was used instead of the Zeiss 1.45NA objective because the slightly larger NA of the Olympus objective gives higher tolerance for placing the excitation beams inside the TIRF annulus. Data was acquired at 1 sec frame rates, with 20 msec exposures from a 488 nm laser for each rotation. TIRF-SIM images were reconstructed as described previously <sup>8</sup>.

## 924 TIRF-SIM imaging of *B. subtilis* strains

Cells were prepared as described in "Media and culture conditions". Cells were placed under a CH agarose pad in a No. 1.5 glass-bottomed dish (MatTek Corp., MA) for imaging. Images were collected on a DeltaVision OMX SR Blaze system in TIRF mode, using an Edge 5.5 sCMOS camera (PCO AG, Germany) and a 60x objective. 75 msec exposures from a 488 nm diode laser were used for each rotation. Spherical aberration was minimized using immersion oil matching. Raw images were reconstructed using SoftWoRx (GE Healthcare, MA) software.

### 931 Particle tracking of JF549-Halo-PBP2A

Particle tracking was performed using the software package FIJI <sup>6</sup> and the TrackMate v3.8 <sup>7</sup> plugin. For calculation of particle velocity, the scaling exponent  $\alpha$ , and track orientations relative to the midline of the cell, only tracks persisting for 7 frames or longer were used. Particle velocity for each track was calculated from nonlinear least squares fitting using the equation MSD(t) = 4Dt + (vt)<sup>2</sup>, where MSD is mean squared displacement, t is time interval, D is the diffusion coefficient, and v is speed. The maximum time interval used was 80% of the track length. To filter for directionally moving tracks, we discarded those with a velocity lower than 0.01 nm/sec. Tracks were also excluded if the R<sup>2</sup> for log MSD versus log t was less than 0.95, indicating a poor ability to fit the MSD curve.

939

### 940 Osmotic shock experiments

- 941 Overnight, exponentially growing cultures (as described in "Media and culture conditions") were diluted into fresh CH
- medium, grown at 37°C to an OD<sub>600</sub> of 0.1 to 0.2, then stained by growing in 100  $\mu$ M Alexa Fluor 488-D-Lysine-NH2
- for 1 hr. Without washing, cells were then loaded into a CellASIC microfluidic flow cell (MilliporeSigma, MA) pre-
- onditioned with media and washed in the chamber via channels 6 and 5. Media in channel 6 was replaced with 5 M

- NaCl, and the flow cell was resealed and imaged immediately. After collecting images across the whole chip pre-
- 946 shock, 5 M NaCl was flowed into the chip via channel 6 and imaged immediately.
- 947

## 948 Alexa Fluor 488-D-Lysine-NH2

- Alexa Fluor 488-D-Lysine-NH2 was synthesized as in Lebar et al., 2014. Briefly, Boc-D-Lys(Cbz)-OH (Bachem,
- 950 Switzerland) was reacted with carbonyldiimidazole (CDI) (MilliporeSigma, MA) in dimethylformamide (DMF) for 1.5 hr,
- then aqueous ammonia was added and stirred for 6 hr to form the carboxamide Boc-D-Lys(Cbz)-NH2. The Cbz
- 952 protecting group was removed by catalytic hydrogenation (20% Pd(OH)2/C) in methanol. The product, Boc-D-Lys-
- 953 NH2, was reacted with CDI in DMF for 1.5 hr, then Alexa Fluor 488 carboxylic acid in DMF was added and reacted for
- 6 hr to yield Boc-D-Lys(Alexa Fluor 488)-NH2. The Boc protecting group was removed by stirring in neat
   trifluoroacetic acid (TFA) for 30 min. The reaction was stopped by dropwise addition of TFA solution to ice-cold ether.
- 955 The precipitate was then HPLC-purified to obtain Alexa Fluor 488-D-Lysine-NH2.
- 950 957

## 958 **CRISPRi titration of MreB expression**

- 959 We used complementarity-based CRISPR knockdown to titrate the MreB expression level in *E. coli*. The degree of
- 960 MreB-SW-msfGFP repression is controlled by introducing mismatches between the guide RNA and the target DNA <sup>9</sup>.
- 961 The repression strength can be tuned by modulating spacer complementarity to msfGFP using different numbers of
- mismatches. To repress msfGFP using CRISPR knockdown, we placed the dCas9 cassette under the control of a
- 963 Ptet promoter and different plasmids to target msfGFP. We induced dCas9 at a constant high level with 100 ng/ml of
- ATc and changed the degree of guide RNA complementarity with different plasmids. For pcrRNA plasmid we use four
- different guide RNAs with 10, 11, 14, and 20 bp of complementarity. For pAV20 plasmid we use four different guide
- RNAs with 5, 10, 14, and 20 bp of complementarity. The cells were grown and imaged in LB containing 50 ug/ml
   kanamycin.

# 969 Protein extraction and labelling

Cell cultures were grown to an  $OD_{600}$  of ~0.4 to 0.6 (cell amounts were normalized by harvesting the equivalent of 3 mL of culture at an  $OD_{600}$  of 0.5). Cells were centrifuged at 6,000 × *g* for 30 sec, washed once in 1 mL of ice-cold 20 mM Tris-HCl, pH 7.5, and 10 mM EDTA (TE), were resuspended in 100 µL of TE, and were killed by boiling in a water bath for 10 min. All samples were frozen at -80°C overnight (or up to 1 week). Thawed samples were digested with 50 µg of lysozyme (Thermo Fisher, MA) in the presence of 1 mM PMSF, at 37°C for 15 min.

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Protein extraction was achieved utilizing a Covaris S220 ultrasonicator (Covaris, MA), under denaturing conditions
upon the addition of urea-based Protein Extraction Buffer DF (Covaris, MA), followed by ice-cold methanol/chloroform
precipitation. Proteins were digested with trypsin (Promega, WI). Each resulting peptide mixture was labeled with one
of a set of up to eleven isotopic tandem mass tags (TMTs) (Thermo Fisher, MA).

## 981 Peptide fractionation and mass spectrometry

Equal amounts of each TMT-labelled sample were combined and fractionated by electrostatic repulsion-hydrophobic
 interaction chromatography, on an Agilent (Santa Clara, CA) 1200 HPLC system using a PolyWAX LP 200 x 2.1 mm,
 5 µm, 300Â column (PolyLC, MD). Peptides were separated across a 70 min gradient from 0% of "buffer A" (90%)

- 984 5 µm, 300A column (PolyLC, MD). Peptides were separated across a 70 min gradient from 0% of buller A (90%)
   985 acetonitrile, 0.1% acetic acid) to 75% of "buffer B" (30% acetonitrile, 0.1% formic acid), with 20 fractions collected over
   986 time. Each fraction was dried in a SpeedVac (Eppendorf, Germany) and resuspended in 0.1% formic acid before
- 987 injection to a mass spectrometer.
- 988

LC-MS/MS was performed on a Thermo Orbitrap Elite (Thermo Fisher, MA) mass spectrometer equipped with a
Waters nanoACQUITY HPLC pump (Waters Corp., MA). Peptides were separated on a 150 μm inner diameter
microcapillary trapping column packed with ~3 cm of C18 Reprosil 5 μm, 100 Å resin (Dr. Maisch GmbH, Germany),
followed by an analytical column packed with ~20 cm of Reprosil 1.8 μm, 200 Å resin. Separation was achieved by
applying a gradient of 5–27% acetonitrile in 0.1% formic acid, over 90 min at 200 nl/min. Electrospray ionization was
achieved by applying a voltage of 2 kV using a home-made electrode junction at the end of the microcapillary column
and spraved from fused-silica PicoTips (New Objective, MA). The Orbitrap instrument was operated in data-

- dependent mode for the mass spectrometry methods. The mass spectrometry survey scan was performed in the
- 997 Orbitrap in the range of 410 1,800 m/z at a resolution of  $12 \times 10^4$ , followed by the selection of the twenty most

intense ions for HCD-MS2 fragmentation using a precursor isolation width window of 2 m/z, AGC setting of 50,000,
and a maximum ion accumulation of 200 msec. Singly-charged ion species were not subjected to HCD fragmentation.
Normalized collision energy was set to 37 V and an activation time of 1 msec. Ions in a 10 ppm m/z window around
ions selected for MS2 were excluded from further selection for fragmentation for 60 sec.

1002

1003 Mass spectrometry analysis: Raw data were submitted for analysis in Proteome Discoverer 2.1.0.81 (Thermo 1004 Scientific) software. Assignment of MS/MS spectra was performed using the Sequest HT algorithm by searching the 1005 data against a protein sequence database that including all entries from B. subtilis (UniProt proteome ID 1006 UP000018540) and other known contaminants such as human keratins and common lab contaminants. Sequest HT 1007 searches were performed using a 15 ppm precursor ion tolerance and required each peptide's N- and C-termini to 1008 adhere with trypsin protease specificity, while allowing up to two missed cleavages. TMT tags on peptide N-termini 1009 and lysine residues (+229.162932 Da) were set as static modifications while methionine oxidation (+15.99492 Da) 1010 was set as a variable modification. An MS2 spectra assignment false discovery rate (FDR) of 1% on protein level was 1011 achieved by applying the target-decoy database search. Filtering was performed using Percolator (64-bit version <sup>10</sup>). 1012 For guantification, a 0.02 m/z window centered on the theoretical m/z value of each of the TMT reporter ions and the 1013 intensity of the signal closest to the theoretical m/z value was recorded. Reporter ion intensities were exported in a 1014 results file of the Proteome Discoverer 2.1 search engine in Microsoft Excel format. The total signal intensity across all 1015 peptides quantified was summed for each TMT channel, and all intensity values were adjusted to account for 1016 potentially uneven TMT-labelling and/or sample handling variance for each labelled channel. For our final relative 1017 protein guantitation analysis, all contaminants from the database search were removed from the results, and only the 1018 remaining *B. subtilis* proteins were used to re-normalize all protein abundances. 1019

# 1021 Strain construction

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1023 bMD277 containing amyE::Pxyl-mreBCD, minCD (B. megaterium)::erm was generated upon transformation of Py79 1024 with a five-piece Gibson assembly reaction<sup>11</sup>, that contained the following PCR products. (1) A 1228 bp fragment 1025 containing sequence upstream of the amyE locus (amplified from Py79 genomic DNA using primers oMD191 and 1026 oMD108); (2) the 1673 bp erythromycin-resistance cassette loxP-erm-loxP (amplified from pWX467 [gift of D. Rudner] 1027 using primers oJM028 and oJM029); (3) a 1532 bp fragment containing the xyIR gene, and the PxyIA promoter with 1028 an optimized ribosomal binding sequence (amplified from pDR150 [gift of D. Rudner] using primers oMD73 and 1029 oMD226); (4) a 4106 bp fragment containing the mreB, mreC, mreD, minC and minD alleles of the B. megaterium 1030 mreB operon (amplified from strain QMB 1551 (ATCC 12872) genomic DNA using primers oMD227 and oMD228); 1031 and (5) a 1216 bp fragment containing the amyE terminator, and sequence downstream of the amyE locus (amplified 1032 from Py79 genomic DNA using primers oMD196 and oMD197).

1033 1034 bMD465 harboring amyE::Pxyl-mreBCD, minCD (B. megaterium)::erm, mreBCD, minC,D::spec (B. megaterium) was 1035 generated upon transformation of bMD277 with a four-piece Gibson assembly reaction, that contained the following 1036 PCR products. (1) A 1275 bp fragment containing sequence upstream of the mreB gene (amplified from Py79 1037 genomic DNA using primers oMD96 and oMD308); (2) a 4121 bp fragment containing the mreB, mreC, mreD, minC 1038 and minD alleles of the B. megaterium mreB operon (amplified from strain QMB 1551 (ATCC 12872) genomic DNA 1039 using primers oMD313 and oMD314); (3) the 1274 bp spectinomycin-resistance cassette loxP-spec-loxP (amplified 1040 from pWX466 [gift of D. Rudner] using primers oJM028 and oJM029); and (4) a 1152 bp fragment containing 1041 sequence downstream of the minD gene (amplified from Py79 genomic DNA using primers oMD300 and oMD315).

1043 bMK355 harboring *amyE::PxyI-mreBCD::erm* was built identical to bMD277, except that PCR product (4) was instead
 1044 a 2493 bp fragment containing the *mreB*, *mreC*, and *mreD* alleles of the *B. subtilis mreB* operon (amplified from Py79
 1045 genomic DNA using primers oMD334 and oMK221).

bMD543 containing amyE::Pxyl-mreBCD::erm, ΔmreBCD, ΔminCD::cat was generated upon transformation of
 bMK355 with a three-piece Gibson assembly reaction, that contained the following PCR products. (1) A 1305 bp
 fragment containing sequence upstream of the mreB gene (amplified from Py79 genomic DNA using primers oMD96
 and oMD298); (2) the 1139 bp chloramphenicol-resistance cassette loxP-cat-loxP (amplified from pWX465 [gift of D.
 Rudner] using primers oJM028 and oJM029); and (3) a 1215 bp fragment containing sequence downstream of the
 minD gene (amplified from Py79 genomic DNA using primers oMD299 and oMD300).

1054 **bMD545** containing *amyE::PxyI-mreBCD::erm*, *AmreBCD*, *PmreB-minC*, *D::spec* was generated upon transformation 1055 of bMD543 with a four-piece Gibson assembly reaction, that contained the following PCR products. (1) A 1305 bp 1056 fragment containing sequence upstream of the mreB gene (amplified from Py79 genomic DNA using primers oMD96 1057 and oMD379); (2) a 1582 bp fragment containing the minC and minD genes, and the minD terminator of the mreB 1058 operon (amplified from Py79 genomic DNA using primers oMD380 and oMD381); (3) the 1274 bp spectinomycin-1059 resistance cassette loxP-spec-loxP (amplified from pWX466 using primers oJM028 and oJM029); and (4) a 1215 bp 1060 fragment containing sequence downstream of the minD gene (amplified from Py79 genomic DNA using primers 1061 oMD300 and oMD382).

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bMD599 harboring ΔponA::kan was generated upon transformation of Py79 with a three-piece Gibson assembly
reaction, that contained the following PCR products. (1) A 1329 bp fragment containing sequence upstream of the
ponA gene (amplified from Py79 genomic DNA using primers oMK001 and oMK002); (2) the 1577 bp kanamycinresistance cassette loxP-kan-loxP (amplified from pWX470 [gift of D. Rudner] using primers oJM028 and oJM029);
and (3) a 1313 bp fragment containing sequence downstream of the ponA gene (amplified from Py79 genomic DNA
using primers oMK005 and oMK006).

1070 bMK005 containing ΔponA::cat was built identical to bMD599, except that PCR product (2) was instead the 1139 bp
 1071 chloramphenicol-resistance cassette loxP-cat-loxP (amplified from pWX465 using primers oJM028 and oJM029).
 1072

1073 **bMD554** containing *yhdG::Phyperspank-ponA::cat* was generated upon transformation of Py79 with a five-piece 1074 Gibson assembly reaction, that contained the following PCR products. (1) A 1219 bp fragment containing sequence 1075 upstream of the yhdH gene (amplified from Py79 genomic DNA using primers oMD328 and oMD367), (2) the 1139 bp 1076 chloramphenicol-resistance cassette loxP-cat-loxP (amplified from pWX465 using primers oJM028 and oJM029); (3) a 1077 1895 bp fragment containing the lacl gene, and the Phyperspank promoter with an optimized ribosomal binding 1078 sequence (amplified from pDR111 [gift of D. Rudner] using primers oMD232 and oMD234); (4) a 2799 bp fragment 1079 containing the ponA coding region (amplified from Py79 genomic DNA using primers oMD365 and oMK370); and (5) a 1080 1252 bp fragment containing the *yhdG* terminator, and sequence downstream of the *yhdG* gene (amplified from Py79 1081 genomic DNA using primers oMD371 and oMD372). 1082

- 1083 bMD586 harboring *yhdG::Phyperspank-ponA::cat*, Δ*ponA::kan* was generated upon transformation of bMD554 with
   1084 genomic DNA from bMD599.
   1085
- bMD594 harboring *yhdG::Pspank-ponA::cat* was built identical to bMD554, except that PCR product (1) was instead a
   1287 bp fragment made with primers oMD328 and oMD329; and PCR product (3) containing the *Pspank* promoter
   was instead amplified from pDR110 (gift of D. Rudner).
- 1090 **bMD598** containing *yhdG::Pspank-ponA::cat*, *ΔponA::kan* was generated upon transformation of bMD594 with genomic DNA from bMD599.
- bMD619 containing amyE::Pxyl-mreBCD::erm, ΔmreBCD, PmreB-minCD::spec, yhdG::Pspank-ponA::cat was
   generated upon transformation of bMD545 with genomic DNA from bMD594.
- bMD620 containing *amyE::Pxyl-mreBCD::erm*, Δ*mreBCD*, PmreB-minCD::spec, yhdG::Pspank-ponA::cat, ΔponA::kan
   was generated upon transformation of bMD619 with genomic DNA from bMD599.
- 1099 bMD622 harboring *amyE::PxyI-mreBCD::erm*, *yhdG::Pspank-ponA::cat*, *ΔponA::kan* was generated upon
   transformation of bMD598 with genomic DNA from bMK355.
- bMD556 harboring *yhdG::Phyperspank-rodA::cat* was built identical to bMD554, except that PCR product (4) was
   instead a 1236 bp fragment containing the *rodA* coding region (amplified from Py79 genomic DNA using primers
   oMD364 and oMK369).
- bMD580 containing *yhdG::Phyperspank-rodA::cat*, Δ*rodA::kan* was generated upon transformation of bMD556 with a three-piece Gibson assembly reaction, that contained the following PCR products. (1) A 1265 bp fragment containing sequence upstream of the *rodA* gene (amplified from Py79 genomic DNA using primers oMD388 and oMD389); (2) the 1577 bp kanamycin-resistance cassette *loxP-kan-loxP* (amplified from pWX470 using primers oJM028 and oJM029); and (3) a 1258 bp fragment containing sequence downstream of the *rodA* gene (amplified from Py79 genomic DNA using primers oMD386 and oMD387).
- **bMD592** harboring *Pxyl-rodA::erm* was generated upon transformation of Py79 with a four-piece Gibson assembly reaction, that contained the following PCR products. (1) A 1265 bp fragment containing sequence upstream of the *rodA* gene (amplified from Py79 genomic DNA using primers oMD388 and oMD389); (2) the 1673 bp erythromycinresistance cassette *loxP-erm-loxP* (amplified from pWX467 using primers oJM028 and oJM029); (3) a 1532 bp fragment containing the *xylR* gene, and the *PxylA* promoter with an optimized ribosomal binding sequence (amplified from pDR150 using primers oMD73 and oMD226); and (4) a 1271 bp fragment containing the *rodA* coding region (amplified from Py79 genomic DNA using primers oMD394 and oMD395).
- bMD627 containing *Pxyl-rodA::erm*, Δ*pbpH::spec* was generated upon transformation of bMD592 with genomic DNA
   from bDR2487.
- 1123

bMD563 containing *yhdG::Phyperspank-pbpA::cat* was built identical to bMD554, except that PCR product (4) was
 instead a 2204 bp fragment containing the *pbpA* coding region (amplified from Py79 genomic DNA using primers
 oMD316 and oMK368).

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bMD573 harboring *yhdG::Phyperspank-pbpA::cat*, Δ*pbpH::spec* was generated upon transformation of bMD563 with
 genomic DNA from bRB776.

bMD574 harboring *yhdG::Phyperspank-pbpA::cat, ΔpbpA::erm, ΔpbpH::spec* was generated upon transformation of
 bMD573 with genomic DNA from bRB776.

bMD590 containing *Pxyl-pbpA::erm* was generated upon transformation of Py79 with a four-piece Gibson assembly reaction, that contained the following PCR products. (1) A 1233 bp fragment containing sequence upstream of the *pbpA* gene (amplified from Py79 genomic DNA using primers oMD75 and oMD125); (2) the 1673 bp erythromycin-resistance cassette *loxP-erm-loxP* (amplified from pWX467 using primers oJM028 and oJM029); (3) a 1537 bp fragment containing the *xylR* gene, and the *PxylA* promoter containing the *pbpA* ribosomal binding sequence (amplified from pDR150 using primers oMD73 and oMD72); and (4) a 1230 bp fragment containing the *pbpA* coding region (amplified from Py79 genomic DNA using primers oMD393 and oMD68).

bMD597 harboring *Pxyl-pbpA::erm*, Δ*pbpH::spec* was generated upon transformation of bMD590 with genomic DNA
 from bDR2487.

bMD557 containing *amyE::Pxyl-mreBCD::erm*, *yhdG::Phyperspank-rodA::cat* was generated upon transformation of
 bMK355 with genomic DNA from bMD556.

bMD571 harboring *amyE::Pxyl-mreBCD::erm*, *yhdG::Phyperspank-pbpA::cat* was generated upon transformation of
 bMK355 with genomic DNA from bMD563.

bMD631 containing *yhdG::Pspank-pbpA::phleo, ΔpbpH::spec, ΔpbpA::cat, Pxyl-rodA::erm* was generated upon
 transformation of bRB773 with genomic DNA from bMD592.

1154 **bYS201** harboring HaloTag-11aa-pbpA::cat was generated upon transformation of Py79 with a four-piece Gibson 1155 assembly reaction, that contained the following PCR products. (1) A 1233 bp fragment containing sequence upstream 1156 of the pbpA gene (amplified from Py79 genomic DNA using primers oMD75 and oMD125); (2) the 1139 bp 1157 chloramphenicol-resistance cassette loxP-cat-loxP (amplified from pWX465 using primers oJM028 and oJM029); (3) a 1158 962 bp fragment containing the native *pbpA* promoter, and *B. subtilis*-optimized coding sequence of the HaloTag 1159 protein (amplified from double-stranded DNA custom-ordered from DNA 2.0/ATUM [Newark, CA] using primers 1160 oYS598 and oYS599); and (4) a 1229 bp fragment containing the pbpA coding region (amplified from Py79 genomic 1161 DNA using primers oMD68 and oMD69).

1163 **bMK385** harboring *amyE::Pxyl-mreBCD::erm*, Δ*mreBCD*, PmreB-minCD::spec, HALO-11aa-pbpA::cat was generated upon transformation of bMD545 with genomic DNA from bYS201.

1165 1166 **bYS19** containing *mreB-SW-msfGFP* was generated upon transformation of bMD88<sup>12</sup> with a three-piece Gibson 1167 assembly reaction that contained the following PCR products. (1) A 898 bp PCR fragment containing sequence 1168 upstream of the *mreB* gene (amplified from Py79 genomic DNA using primers oMD134 and oYS34); (2) a 774 bp 1169 fragment containing B. subtilis-optimized coding sequence of the msfGFP fluorescent protein (amplified from plasmid 1170 DNA custom-ordered from DNA 2.0/ATUM [Newark, CA] using primers oYS35 and oYS36); and (3) a 1629 bp 1171 fragment containing sequence downstream of the mreB gene (amplified from Py79 genomic DNA using primers 1172 oYS37 and oMD116). This strain is markerless, and selection was in the presence of 0.5% glucose for colonies that 1173 no longer required xylose for viability. 1174

- bYS977 harboring *amyE::Pxyl-mreB-SW-msfGFP, mreCD::erm,* was built identical to bMD277, except that PCR
   product (4) was instead a 3234 bp fragment containing *mreB-SW-msfGFP, mreC*, and *mreD*, amplified from bYS19
   genomic DNA using primers oMD334 and oMK221).
- bYS979 containing *amyE::Pxyl-mreB-SW-msfGFP, mreCD::erm, ΔmreBCD, ΔminCD::cat* was generated upon transformation of bYS977 with genomic DNA from bMD543.
- bYS981 harboring *amyE::Pxyl-mreB-SW-msfGFP, mreCD::erm, ΔmreBCD, PmreB-minC,D::spec* was generated
   upon transformation of bYS979 with genomic DNA from bMD545.
- 1184 1185 AV88 containing 186::Ptet-dCas9, mreB-SW-msfGFP was obtained by transduction of strain LC69 with a P1 phage 1186 lysate obtained from strain NO50, and subsequent excision of the kanamycin resistance cassette by transient 1187 expression of a flippase. To make the plasmids expressing single-guide RNAs, the psgRNA plasmid was amplified 1188 three times with primers pairs V162/V165, V163/V166 and V164/V167. The three generated fragments were 1189 assembled together by Gibson assembly to obtain the pAV20 vector. To insert the anti-GFP single-guide RNAs into 1190 this vector, we annealed and phosphorylated the following pairs of oligonucleotides: V275/V283 for G5; V274/V282 1191 for G10; V273/V281 for G14; and V272/V280 for G20. These were then inserted in pAV20 by restriction cloning with 1192 Bsal. As pAV20 carries two sgRNA insertion sites, a non-targeting oligonucleotide pair (V279/V287) was inserted into 1193 a second site during the same reaction. 1194
- 1195

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117.

# 1196 Statistics.

All P-values are reported in figures. P-values were calculated in GraphPad Prism using the Mann-Whitney test with a two-tailed P value. Means, SD, and N for all data points are reported in table S1.

# 1200 Table S1 – Data summary and statistics.

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## 1202 **A - Figure 1.**

Panel	Strain condition.		parameter	Value	SD	Ν
В	Py79		width(nm)	754.70	81.19	466
В	B. megaterium		width(nm)	1377.88	98.09	337
В	bMD465	1% glucose	width(nm)	807.52	61.24	267
В	bMD465	30 mM xylose	width(nm)	771.20	104.12	135
С	bMD545	0.1 mM xylose	width(nm)	1533.00	247.40	501
С	bMD545	0.5 mM xylose	width(nm)	1179.00	165.50	466
С	bMD545	1 mM xylose	width(nm)	1059.00	145.70	395
С	bMD545	1.75 mM xylose	width(nm)	930.80	83.53	282
С	bMD545	2.5 mM xylose	width(nm)	807.80	90.69	461
С	bMD545	5 mM xylose	width(nm)	735.00	80.10	269
С	bMD545	30 mM xylose	width(nm)	722.10	60.64	619
С	bMK355	30 mM xylose	width(nm)	696.50	85.40	276
D	bMD599		width(nm)	575.90	97.00	174
D	bMD598	0 M IPTG	width(nm)	580.60	60.57	148
D	bMD598	1 μM IPTG	width(nm)	601.00	68.93	141
D	bMD598	10 μM IPTG	width(nm)	651.40	74.63	742
D	bMD598	25 μM IPTG	width(nm)	767.80	91.60	309
D	bMD598	100 μM IPTG	width(nm)	825.00	78.83	361
D	bMD598	1000 μM IPTG	width(nm)	868.70	78.53	654
D	bMD586	25 μM IPTG	width(nm)	935.30	106.00	332
D	bMD586	50 μM IPTG	width(nm)	1031.30	97.14	381
D	bMD586	100 µM IPTG	width(nm)	1149.45	119.40	133
D	bMD586	1000 µM IPTG	width(nm)	1283	137.8	619
D	bMD554	1000 μM IPTG	width(nm)	1519	169.4	276
E	bMD620	0.1 mM xylose, 0 M IPTG	width(nm)	786.00	159.30	452
E	bMD620	0.5 mM xylose, 1 μM IPTG	width(nm)	744.50	137.90	265
E	bMD620	2.5 mM xylose, 10 µM IPTG	width(nm)	754.40	76.76	522
Е	bMD620	5 mM xylose, 25 µM IPTG	width(nm)	749.00	73.91	288
E	bMD620	30 mM xylose, 100 µM IPTG	width(nm)	784.80	76.63	589
E	bMD622	30 mM xylose, 1000 µM IPTG	width(nm)	773.30	58.09	665
F	Py79	LB	width(nm)	770.50	44.09	590
F	Py79	CH	width(nm)	754.70	81.19	466
F	Py79	S750 glucose	width(nm)	767.49	78.52	389
F	Py79	S750 Glycerol	width(nm)	756.90	70.21	596
F	Py79	S750 Sucrose	width(nm)	761.51	63.79	360

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## 1205 1206 **E**

### B - Figure 2

<u>Б-Г</u>	iguie	2				
Panel	Strain	Conditions	parameter	Value	<sup>SD</sup> 12	0⋫
А	bMD592	0.1mM xyl	width(nm)	1113	221.8	_ 387
А	bMD592	0.5mM xyl	width(nm)	959.8	156.6 2	089
А	bMD592	1mM xyl	width(nm)	753.2	106.6	370
А	bMD592	30mM xyl	width(nm)	964.2	77.79	466
А	bMD580	1mM IPTG	width(nm)	1133	110.8	555
А	bMD556	1mM IPTG	width(nm)	1323	155.5	788
А	bMD598	0.1mM xyl	width(nm)	1081	206.5	455
А	bMD599	0.5mM xyl	width(nm)	927.6	117.8	442
А	bMD600	1mM xyl	width(nm)	749.3	77.61	135
А	bMD601	30mM xyl	width(nm)	782	52.83	91
А	bMD602	1mM IPTG	width(nm)	790.1	66.96	288
А	bMD603	1mM IPTG	width(nm)	795.1	59.96	443
В	bMD627	0.025mM xyl	width(nm)	1381	334.7	92
В	bMD627	1mM xyl	width(nm)	756.4	52.33	983
В	bMD627	30mM xyl	width(nm)	793.4	62.71	414
В	bMD631	025µM xyl, 15µM IPTG	width(nm)	1346	371.4	79
В	bMD631	1mM xyl, 15µM IPTG	width(nm)	757.9	55.22	441
В	bMD631	30mM xyl 15µM IPTG	width(nm)	966.1	80.25	503
В	bMD631	30mM xyl 20µM IPTG	width(nm)	902	106.4	262
С	bMD557	30mM xyl, 40µM IPTG	width(nm)	987.1	87.58	230
С	bMD557	30mM xyl 100µM IPTG	width(nm)	856.5	121.3	266
D	bMD620	0.1m M xyl 0 IPTG	Surface growth	0.755	0.1681	183
D	bMD620	0.5mM xyl 1µM IPTG	Surface growth	1.025	0.171	63
D	bMD620	2.5mM xyl 10µM IPTG	Surface growth	1.146	0.1108	57
D	bMD620	5mM xyl 25µM IPTG	Surface growth	1.174	0.0692	78
D	bMD620	30mM xyl 100µM IPTG	Surface growth	1.282	0.1895	46
D	bMD622	30mM xyl 1mM IPTG	Surface growth	1.194	0.1063	54
D	bMD620	0.1mM xyl 25µM IPTG	Surface growth	0.816	0.2653	17
D	bMD620	0.5mM xyl 25µM IPTG	Surface growth	1.203	0.138	111
D	bMD620	1mM xyl 25µM IPTG	Surface growth	1.182	0.174	59
D	bMD620	2.5mM xyl 25µM IPTG	Surface growth	1.188	0.1229	121
D	bMD620	5mM xyl 25µM IPTG	Surface growth	1.174	0.0691	78
D	bMD620	30mM xyl 25µM IPTG	Surface growth	1.326	0.1969	50
D	bMD620	5mM xyl 1µM IPTG	Surface growth	1.274	0.182	78
D	bMD620	5mM xyl 10µM IPTG	Surface growth	1.256	0.1302	140
D	bMD620	5mM xyl 25µM IPTG	Surface growth	1.174	0.0692	78
D D	bMD620	5mM xyl 50µM IPTG	Surface growth	1.288	0.1677	486 123
D	bMD620 bMD599	5mM xyl 100µM IPTG	Surface growth Surface growth	1.353 1.263	0.1076 0.1487	84
D	bAM268		•	1.147	0.1487	04 177
D	bAM288	25µM IPTG	Surface growth Surface growth	1.147	0.1432	51
D	bMD631	1mM xyl 15µM IPTG	Surface growth	1.398	0.1488	187
D	bMD631	30mM xyl 15µM IPTG	Surface growth	1.338	0.0789	49
D	bMD631	30mM xyl 40µM IPTG	Surface growth	1.173	0.1377	49
D	Py79	оопшилугторийн то	Surface growth	1.173	0.1527	84
D	bMD620	0.1mM xyl 0 IPTG	Bulk growth	1.49	0.1527	04 5
D	bMD620	0.5mM xyl 1µM IPTG	Bulk growth	1.49	0.09	10
D	bMD620	2.5mM xyl 10µM IPTG	Bulk growth	1.85	0.03	9
D	bMD620	5mM xyl 25µM IPTG	Bulk growth	1.85	0.07	5
D	bMD620	30mM xyl 100µM IPTG	Bulk growth	1.89	0.05	5
D	bMD622	30mM xyl 1mM IPTG	Bulk growth	2	0.03	5
D	bMD620	0.1mM xyl 25µM IPTG	Bulk growth	1.277	0.07	5
D	bMD620	0.5mM xyl 25µM IPTG	Bulk growth	1.98	0.05	5
D	bMD620	1mM xyl 25µM IPTG	Bulk growth	1.88	0.04	5
D	bMD620	2.5mM xyl 25µM IPTG	Bulk growth	1.85	0.03	5
D	bMD620	5mM xyl 25µM IPTG	Bulk growth	2.05	0.13	10
D	bMD620	30mM xyl 25µM IPTG	Bulk growth	1.97	0.03	5
D	bMD620	5mM xyl 1µM IPTG	Bulk growth	1.84	0.18	10
D	bMD620	5mM xyl 10µM IPTG	Bulk growth	1.96	0.14	9
D	bMD620	5mM xyl 25µM IPTG	Bulk growth	1.76	0.05	5
D	bMD620	5mM xyl 50µM IPTG	Bulk growth	1.83	0.03	5
D	bMD620	5mM xyl 100µM IPTG	Bulk growth	1.86	0.05	5
D	bMD599		Bulk growth	1.19	0.04	10
D	bAM268		Bulk growth	1.17	0.03	5
D	bAM288	25µM IPTG	Bulk growth	1.7	0.02	5
					2.02	

## C - Figure 3

Panel	Strain	Conditions	Width	Width	(N)	Filament	Density
			(um)	SD	()	density	SD
В	YS981	0.1mM Xyl	1.3826	0.2	336	0.29	0.28
В	YS981	0.25mM Xyl	1.2808	0.2894	214	0.37	0.35
в	YS981	0.5mM Xyl	1.2557	0.1674	130	0.55	0.42
В	YS981	1mM Xyl	1.0093	0.1128	113	1.49	0.85
В	YS981	1.75mM Xyl	1.0661	0.0969	116	1.36	0.58
В	YS981	2.5mM Xyl	1.0469	0.1315	107	1.64	0.48
В	YS981	5mM Xyl	1.0051	0.0985	92	2.14	0.85
В	YS981	30mM Xyl	1.0124	0.0731	190	2.12	0.83

**D - Figure S3 – Simulation** 

Panel	Parameter	Input value	Assay	Filament density	\$ <b>P</b> 210	N
A	density	0.7958	Correlated motion	0.8435	0.1207	5
А	density	1.5915	Correlated motion	1.7985	0.1207	5
А	density	2.3873	Correlated motion	2.3873	0.4674	5
A	density	3.1831	Correlated motion	3.3263	0.62	5
A	density	3.9789	Correlated motion	3.7561	0.5261	5
A	density	4.7746	Correlated motion	4.2494	0.795	5
А	density	5.5704	Correlated motion	4.902	0.5997	5
А	density	6.3662	Correlated motion	4.8383	0.6149	5
A	density	7.162	Correlated motion	5.077	0.462	5
A	density	0.7958	Tracking	0.7321	0.2478	5
A	density	1.5915	Tracking	1.4006	0.1332	5
A	density	2.3873	Tracking	1.8462	0.1424	5
A	density	3.1831	Tracking	2.1008	0.2293	5
A	density	3.9789	Tracking	2.7375	0.5559	5
A	density	4.7746	Tracking	2.9444	0.497	5
A	density	5.5704	Tracking	3.0876	0.2999	5
A	density	6.3662	Tracking	3.008	0.3487	5
A	density	7.162	Tracking	3.1831	0.4135	5
В	velocity	20	Correlated motion	2.6101	0.4034	5
В	velocity	25	Correlated motion	3.3263	0.43	5
В	velocity	30	Correlated motion	3.2945	0.5114	5
В	velocity	35	Correlated motion	3.3423	0.2699	5
в	velocity	40	Correlated motion	3.0399	0.1815	5
В	velocity	45	Correlated motion	2.9125	0.2361	5
В	velocity	50	Correlated motion	3.1513	0.3585	5
В	velocity	60	Correlated motion	2.9762	0.2361	5
В	velocity	70	Correlated motion	2.7375	0.215	5
В	velocity	80	Correlated motion	2.5624	0.2414	5
В	velocity	90	Correlated motion	2.9125	0.3716	5
В	velocity	20	Tracking	2.4032	0.2945	5
В	velocity	25	Tracking	2.6261	0.1866	5
в	velocity	30	Tracking	2.451	0.2414	5
В	velocity	35	Tracking	2.4351	0.2293	5
В	velocity	40	Tracking	2.5465	0.4708	5
В	velocity	45	Tracking	2.3237	0.3051	5
В	velocity	50	Tracking	2.6738	0.438	5
В	velocity	60	Tracking	2.642	0.495	5
В	velocity	70	Tracking	3.008	0.3051	5
В	velocity	80	Tracking	2.7056	0.4606	5
В	velocity	90	Tracking	2.2759	0.165	5

Panel	Parameter	Input value	Assay	Filament density	SD	Ν
С	number	-30	Correlated motion	3.947	0.3009	5
С	number	-25	Correlated motion	3.5173	0.3749	5
С	number	-20	Correlated motion	3.5173	0.4112	5
С	number	-15	Correlated motion	3.3741	0.4271	5
С	number	-10	Correlated motion	3.3104	0.6381	5
С	number	-5	Correlated motion	3.2627	0.5033	5
С	number	0	Correlated motion	3.1831	0.5955	5
С	number	5	Correlated motion	3.4855	0.5912	5
С	number	10	Correlated motion	3.5969	0.1631	5
С	number	15	Correlated motion	3.7083	0.412	5
С	number	20	Correlated motion	3.7879	0.4761	5
С	number	25	Correlated motion	3.581	0.464	5
С	number	30	Tracking	3.6765	0.5261	5
С	angle	-30	Tracking	3.3741	0.3262	5
С	angle	-25	Tracking	2.8648	0.232	5
С	angle	-20	Tracking	3.008	0.3252	5
С	angle	-15	Tracking	2.6897	0.43	5
С	angle	-10	Tracking	2.6579	0.3061	5
С	angle	-5	Tracking	2.451	0.2722	5
С	angle	0	Tracking	2.2918	0.3348	5
С	angle	5	Tracking	2.7215	0.3252	5
С	angle	10	Tracking	2.8966	0.3923	5
С	angle	15	Tracking	2.8011	0.1038	5
С	angle	20	Tracking	2.6897	0.3441	5
С	angle	25	Tracking	2.7693	0.2836	5
С	angle	30	Tracking	3.4218	0.1125	5
D	length	5	Correlated motion	3.0876	0.3348	5
D	length	25	Correlated motion	3.1831	0.1489	5
D	length	50	Correlated motion	3.2786	0.3664	5
D	length	125	Correlated motion	3.1513	0.0907	5
D	length	250	Correlated motion	3.0717	0.5114	5
D	length	300	Correlated motion	3.1354	0.4893	5
D	length	350	Correlated motion	2.5306	0.76	5
D	length	375	Correlated motion	2.5146	0.2293	5
D	length	400	Correlated motion	2.3396	0.1332	5
D	length	450	Correlated motion	2.0531	0.206	5
D	length	500	Correlated motion	1.8303	0.4285	5
D	length	5	Tracking	2.2759	0.1743	5
D	length	25	Tracking	2.3555	0.5415	5
D	length	50	Tracking	2.4351	0.1832	5
D	length	125	Tracking	2.5465	0.1125	5
D	length	250	Tracking	2.5783	0.2615	5
D	length	300	Tracking	2.6579	0.438	5
D	length	350	Tracking	2.5783	0.4344	5
D	length	375	Tracking	2.4669	0.3858	5
D	length	400	Tracking	2.9125	0.3163	5
D	length	450	Tracking	3.0399	0.2945	5
D	length	500	Tracking	2.8011	0.4886	5

# 1211 E - Figure 4, S4

Panel	Strain	Conditions	Parameter	Value	SD	(N)
B, C	YS981	30/0.001	Width	1.3826	0.2	93
B, C	YS981	5/0.025	Width	1.2808	0.2894	214
B, C	YS981	0.5/100	Width	1.2557	0.1674	130
B, E	YS981	30/0.001	Retardance			93
B, E	YS981	5/0.025	Retardance			91
B,E	YS981	0.5/100	Retardance			137
S4A	bMD620	30/0.001	Cell width (TEM)	512.38	45.5616	16
S4A	bMD620	5/0.025	Cell width (TEM)	705.71	51.6678	35
S4A	bMD620	0.5/100	Cell width (TEM)	1138.8	229.043	58
S4A	Py79		Cell width (TEM)	748.08	31.5936	32
S4A	bMD620	30/0.001	Cell wall width (TEM)	14	3.2609	16
S4A	bMD620	5/0.025	Cell wall width (TEM)	15.34	1.6439	35
S4A	bMD620	0.5/100	Cell wall width (TEM)	19.51	3.7475	58
S4A	Py79		Cell wall width (TEM)	15.11	1.3955	32
H, S4B, S4C	bMD620	5/0.025 preshock	Sacculus length	4470	1138	446
H, S4B, S4C	bMD620	5/0.025 preshock	Sacculus Width	782.6	53.03	446
H, S4B, S4C	bMD620	5/0.025 shocked	Sacculus length	3120	973.5	309
H, S4B, S4C	bMD620	5/0.025 shocked	Sacculus Width	676.6	41.38	309
H, S4B, S4C	bMD620	0.5/100 preshock	Sacculus length	4261	1063	145
H, S4B, S4C	bMD620	0.5/100 preshock	Sacculus Width	1283	199.2	145
H, S4B, S4C	bMD620	0.5/100 shocked	Sacculus length	3386	812.6	146
H, S4B, S4C	bMD620	0.5/100 shocked	Sacculus Width	1085	164.9	146
H, S4B, S4C	bMD620	30/0.001 preshock	Sacculus length	3789	1535	1112
H, S4B, S4C	bMD620	30/0.001 preshock	Sacculus Width	586.4	43.09	1112
H, S4B, S4C	bMD620	30/0.001 shocked	Sacculus length	2315	919.6	408
H, S4B, S4C	bMD620	30/0.001 shocked	Sacculus Width	527.6	59.77	408

# **F - Figure 5**.

Panel	Strain	Mutation	Conditions	Width (um)	Width SD	(N)	Filament density	Density SD	Source.
А	NO50 - A20S	A20S		0.8671	0.0969	474	1.15	0.39	13
А	NO50 - L322Q	L322Q		0.8387	0.084	661	1.19	0.65	13
А	NO50 - S14A	S14A		0.975	0.1879	559	1.01	0.31	13
А	NO50 - R74C	R74C		0.91	0.1645	517	1.01	0.43	13
А	NO50	WT		1.1343	0.1243	656	1.04	0.24	13
А	NO50 - A53∆	A53∆		1.2579	0.1261	610	0.40	0.23	13
А	NO50 - S185F	S185F		0.9296	0.154	715	0.88	0.37	13
А	NO50 - F84V	F84V		1.2876	0.2494	946	0.39	0.22	13
А	RM478	S14A ∆RodZ		1.3285	0.2075	388	0.33	0.18	14
A	eKC508	WT		1.18	0.11	323	0.41	0.41	15
A	eKC507	E276D		0.97	0.08	220	1.03	0.47	16
A	eKC967	D83E		1.12	0.255	384	0.72	0.17	15
A	eKC968	R124C		1.0061	0.2395	197	0.91	0.26	15
А	eKC969	A174T		1.2832	0.1777	279	0.54	0.27	15
А	eKC407	EC - mrdA		1.1325	0.1199	249	0.83	0.31	16
А	eKC408	ST - mrdA		1.0715	0.1358	131	0.95	0.41	16
А	eKC409	VC - mrdA		1.4799	0.1121	773	0.42	0.16	16
A	eKC410	YP - mrdA		1.1447	0.1887	457	0.85	0.31	16
А	eKC717		0.005% Arab.	1.4657	0.2501	775	0.33	0.13	15
А	eKC717		0.01% Arab.	1.2809	0.1892	561	0.44	0.25	15
А	eKC717		0.05% Arab.	1.229	0.149	794	0.88	0.34	15
А	eKC717		0.1% Arab.	1.2239	0.2842	575	0.88	0.46	15
А	eKC717		0.2% Arab.	1.194	0.2329	239	0.90	0.34	15
В	AV88 pAV20 G5	WT – no repression	100 ng/ml ATc	1.1132	0.1044	812	0.93	0.31	This work.
В	AV88 pAV20 G10		100 ng/ml ATc	1.1284	0.1981	691	0.80	0.37	This work.
В	AV88 pAV20 G14		100 ng/ml ATc	1.6528	0.308	531	0.23	0.21	This work.
В	AV88 pAV20 G20		100 ng/ml ATc	1.9188	0.278	451	0.15	0.09	This work.
В	AV88 pcrRNA G10		100 ng/ml ATc	1.1355	0.1052	1103	0.89	0.44	This work.
В	AV88 pcrRNA G11		100 ng/ml ATc	1.2541	0.1008	878	0.56	0.33	This work.
В	AV88 pcrRNA G14		100 ng/ml ATc	1.2938	0.1323	342	0.56	0.34	This work.
В	AV88 pcrRNA G20		100 ng/ml ATc	1.9238	0.3106	409	0.08	0.07	This work.

# 1216 Table S2 – *B. subtilis* strains used in this study

Strain	Relevant genotype	Source
bMD277	amyE::PxyI-mreBCD, minCD (B. megaterium)::erm	This work
bMD465	amyE::PxyI-mreBCD, minCD (B. megaterium)::erm, mreBCD, minC,D::spec (B. megaterium)	This work
bMK355	amyE::Pxyl-mreBCD::erm	This work
bMD543	amyE::Pxyl-mreBCD::erm ΔmreBCD, ΔminCD::cat	This work
bMD545	amyE::PxyI-mreBCD::erm ΔmreBCD, PmreB-minCD::spec	This work
bMD599	ДропА::kan	This work
bMK005	AponA::cat	17
bMD554	yhdG::Phyperspank-ponA::cat	This work
bMD586	yhdG::Phyperspank-ponA::cat ΔponA::kan	This work
bMD594	yhdG::Pspank-ponA::cat	This work
bMD598	yhdG::Pspank-ponA::cat ΔponA::kan	This work
bMD619	amyE::Pxyl-mreBCD::erm ΔmreBCD, PmreB-minCD::spec yhdG::Pspank-ponA::cat	This work
bMD620	amyE::Pxyl-mreBCD::erm ΔmreBCD, PmreB-minCD::spec yhdG::Pspank-ponA::cat ΔponA::kan	This work
bMD622	amvE::Pxyl-mreBCD::erm yhdG::Pspank-ponA::cat ΔponA::kan	This work
bMD556	yhdG::Phyperspank-rodA::cat	This work
bMD580	yhdG::Phyperspank-rodA::cat ΔrodA::kan	This work
bMD592	Pxyl-rodA::erm	This work
bDR2487	AppH::spec	18
bMD627	Рхуl-rodA::erm ΔpbpH::spec	This work
bMD563	yhdG::Phyperspank-pbpA::cat	This work
bRB776	yhdG::Pspank-pbpH::phleo ДрbpH::spec ДрbpA::erm	18
bMD573	yhdG::Phyperspank-pbpA::cat ΔpbpH::spec	This work
bMD574	yhdG::Phyperspank-pbpA::cat ΔpbpA::erm ΔpbpH::spec	This work
bMD590	Pxyl-pbpA::erm	This work
bMD597	PxyI-pbpA::erm ΔpbpH::spec	This work
bMD557	amyE::Pxyl-mreBCD::erm yhdG::Phyperspank-rodA::cat	This work
bMD571	amyE::Pxyl-mreBCD::erm yhdG::Phyperspank-pbpA::cat	This work
bRB773	yhdG::Pspank-pbpA::phleo ΔpbpH::spec ΔpbpA::cat	18
bMD631	yhdG::Pspank-pbpA::phleo ΔpbpH::spec ΔpbpA::cat Pxyl-rodA::erm	This work
bYS201	HaloTag-11aa-pbpA::cat	This work
bMK385	amyE::Pxyl-mreBCD::erm ΔmreBCD, PmreB-minCD::spec HaloTag-11aa-pbpA::cat	This work
bAM268	ДрbpF ДрbpG ДрbpD ДрonA::kan	19
bAM288	ДрbpF ДрbpG ДрbpD ДрonA::kan amyE::Phyperspank-rodA-His10::spec	19
bMD88	Pxyl-mreC::erm	12
bYS09	mreB-mNeonGreen	20
bYS19	mreB-SW-msfGFP	This work
bYS977	amyE::PxyI-mreB-SW-msfGFP,mreC,mreD::erm	This work
bYS979	amyE::Pxyl-mreB-SW-msfGFP,mreC,mreD::erm, ΔmreBCD, ΔminCD::cat	This work
bYS981	amyE::Pxyl-mreB-SW-msfGFP,mreC,mreD::erm, ΔmreBCD, PmreB-minC,D::spec	This work

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# Table S3 – E. coli strains used in this study

Strain	Relevant genotype	Source
MG1655	Wild-type E. coli	CGSC #6300
NO50	mreB-msfGFP sw	13
NO50 - A20S	NO50 with mreB(A20S)	13
NO50 - L322Q	NO50 with mreB(L322Q)	13
NO50 - ΔA53	NO50 with mreB( $\Delta$ A53)	13
NO50 - S185F	NO50 with mreB(S185F)	13
NO50 - F84V	NO50 with mreB(F84V)	13
NO50 - R74C	NO50 with mreB(R74C)	13
NO50 - S14A	NO50 with mreB(S14A)	14
RM478	ΔrodZ, mreB(S14A)-msfGFP	14
KC407	ΔmrdA csrA::frt, mreB'::msfGFP-mreB" + pww308 Ec mrdA	16
KC408	ΔmrdA csrA::frt, mreB'::msfGFP-mreB" + pww308 St mrdA	16
KC409	ΔmrdA csrA::frt, mreB'::msfGFP-mreB" + pww308 Vc mrdA	16
KC410	ΔmrdA csrA::frt, mreB'::msfGFP-mreB" + pww308 Yp mrdA	16
NO34	csrD::kan, mreB::msfGFP-mreB	13
KC507	csrD::kan, mreB'::msfGFP-mreB'-E276D	15
KC717	csrD::kan, mreB::msfGFP-mreB,ProdZ<>(frt araC PBAD)	15

KC967	csrD::kan, mreB'::msfGFP-mreB'-D83E	15
KC968	csrD::kan, mreB'::msfGFP-mreB'-R124C	15
KC969	csrD::kan, mreB'::msfGFP-mreB'-A174T	15
AV88	186:: P <sub>Tet</sub> -dCas9, mreB::msfGFP-mreB	This work
AV88 pAV20 G5	AV88 with pAV20 G5	This work
AV88 pAV20 G10	AV88 with pAV20 G10	This work
AV88 pAV20 G14	AV88 with pAV20 G14	This work
AV88 pAV20 G20	AV88 with pAV20 G20	This work
AV88 pcrRNA G10	AV88 with pcrRNA G10	This work
AV88 pcrRNA G11	AV88 with pcrRNA G11	This work
AV88 pcrRNA G14	AV88 with pcrRNA G14	This work
AV88 pcrRNA G20	AV88 with pcrRNA G20	This work

# 1219

# 1220 Table S4 – Plasmids used in this study

pAV20 G5	guide RNA is sgRNA pAV20 G5, 5 matching base pairs against GFP	This work
pAV20 G10	guide RNA is sgRNA pAV20 G10, 10 matching base pairs against GFP	This work
pAV20 G14	guide RNA is sgRNA pAV20 G14, 14 matching base pairs against GFP	This work
pAV20 G20	guide RNA is sgRNA pAV20 G20, 20 matching base pairs against GFP	This work
pcrRNA G10	guide RNA is native crRNA, 10 matching base pairs against GFP	This work
pcrRNA G11	guide RNA is native crRNA, 11 matching base pairs against GFP	This work
pcrRNA G14	guide RNA is native crRNA, 14 matching base pairs against GFP	This work
pcrRNA G20	guide RNA is native crRNA, 20 matching base pairs against GFP	This work

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# 1223 Table S5 – Oligonucleotides used in this study

Oligo	Sequence
oJM028	TTCTGCTCCCTCGCTCAG
oJM029	CAGGGAGCACTGGTCAAC
oMD68	TTTTACAGCTGATCCCATAGCGTAG
oMD69	CTGGACAGGGCTCAGGTCCGGGATCTGGCATGAGGAGAAATAAACCAAAAAGCAAAATC
oMD72	AACATCACCTTTTCTATTTAATCGATCCATTCAAATACAGATGCATTTTATTTC
oMD73	ATACGAACGGTACTGAGCGAGGGGGGGGAGCAGAAGAATTCGAGCTTGCATGCCTG
oMD75	CAGCCGATAACCTTTTATTTGGCTG
oMD96	TTTCACCAGAAGAAAACGTCCAATG
oMD108	ACGAACGGTAGTTGACCAGTGCTCCCTGTCTTGACACTCCTTATTTGATTTTTGAAGAC
oMD116	CAGCCAGAACTGCTATCAATATCACTAC
oMD125	TACGAACGGTAGTTGACCAGTGCTCCCTGTTTACAGAAAGAA
oMD134	CCGAGCAGGGAAGATATTGAAG
oMD191	TTTGGATGGATTCAGCCCGATTG
oMD196	GGGCAAGGCTAGACGGG
oMD197	TCACATACTCGTTTCCAAACGGATC
oMD226	GGTAGTTCCTCCTTAATCGATCCATTCAAATACAGATGCATTTTATTTC
oMD227	TTTGAATGGATCGATTAAGGAGGAACTACCATGTTTGGAATCGGTACTAGAGACC
oMD228	TTTCGGTAAGTCCCGTCTAGCCTTGCCCTTAAGAACGAAC
oMD232	GGTAGTTCCTCCTTAAAGCTTAATTGTTATCCGCTCACAAT
oMD234	ATACGAACGGTACTGAGCGAGGGGGGGGAGCAGAATAATGGATTTCCTTACGCGAAATACG
oMD298	ATACGAACGGTAGTTGACCAGTGCTCCCTGATGTATGTAT
oMD299	ATACGAACGGTACTGAGCGAGGGAGCAGAATGTGATAGAATCAAAGAGAAGAATCTGAC
oMD300	CTCTTCCTTTAACGGCCGATTC
oMD308	ATGTATGTATCTTCCTTTCTTAAAGCAAAAATACC
oMD313	ACAGCATATGCTTTGTCAGATTCTTCTCTTTGATTCTATCACATTAAGAACGAAC
oMD314	AAGAGAAGAATCTGACAAAGCATATGCTGTGTCAGGTTTTTTTGTTTTTCAGGGAGCACTGGTCAAC
oMD315	ATACGAACGGTACTGAGCGAGGGGGGGGGAGCAGAAGCCTGCTTTGTTCTTGACTAAACC
oMD316	TAACAATTAAGCTTTAAGGAGGAACTACCATGAGGAGAAATAAACCAAAAAGCAAAATC
oMD328	CCTGTTTCAGGTTCGGTGTATAC
oMD329	ATACGAACGGTAGTTGACCAGTGCTCCCTGTAAACGTCACTTCCTTTCCAGC
oMD334	TTTGAATGGATCGATTAAGGAGGAACTACCATGTTTGGAATTGGTGCTAGAGAC
oMD364	ATAACAATTAAGCTTTAAGGAGGAACTACCATGAGTCGATATAAGAAACAGCAAAGC
oMD365	ATAACAATTAAGCTTTAAGGAGGAACTACCATGTCAGATCAATTTAACAGCCGTG
oMD367	ACGAACGGTAGTTGACCAGTGCTCCCTGGCCAAGAACAGAAAAGAGGCGGCCTGTATGGC

	GAAAAGAGGCGGCCTGTATGGCCGTCAAGAATTAAAAAGATAACTTCTGTATTTCGTCAG
oMD369	
oMD370	GAAAAGAGGCGGCCTGTATGGCCGTTAATTTGTTTTTCAATGGATGAGTTGATTGTTTG
oMD371	CGGCCATACAGGCCG
oMD372	GCTGACGGTAAAATCGGCAATATC
oMD379	TGTTACATATTGCTGCTTTTTGGTCTTCACATGTATGTAT
oMD380	GTGAAGACCAAAAAGCAGCAATATG
oMD381	ATACGAACGGTACTGAGCGAGGGAGCAGAAAAAAAAAAA
oMD382	ATACGAACGGTAGTTGACCAGTGCTCCCTGTGTGATAGAATCAAAGAGAAGAATCTGAC
oMD386	GTTTTCGCTGCTATATGACTGATTAGC
oMD387	ATACGAACGGTACTGAGCGAGGGAGCAGAATATGCAGACAGCCTTTACAGAGG
oMD388	ATACGAACGGTAGTTGACCAGTGCTCCCTGTCTATCCCGCCTTACATTTTCATCG
oMD389	TGTTGTATGGATTTTCATCTTTACAGGTG
oMD393	ATGGATCGATTAAATAGAAAAGGTGATGTTATGAGGAGAAATAAAC
oMD394	TTTGAATGGATCGATTAAGGAGGAACTACCATGAGTCGATATAAGAAACAGCAAAGC
oMD395	TGCACAAAAAAAGACAGCCTCTG
oMK001	GCCTTATCCTTTCCTCCGCC
oMK002	CTGAGCGAGGGAGCAGAACATCTCAACCTTTCGTTAATCAACC
oMK005	TACGAACGGTAGTTGACCAGTGCTCCCTGGTACACAATAAAAAACCTCCCGTTTTAACAG
oMK006	CGTGTACAAGCAAAGCAGAATGAAC
oMK221	TTCTTTCGGTAAGTCCCGTCTAGCCTTGCCCTTACTCATCTCTCAATTCTTTCT
oYS34	TTCTTCCCCTTTTCGCATACTTGAGCCGCTGATTTCCATTTTAATCGCTTCAGCCG
oYS35	GCGATTAAAATGGAAATCAGCGGCTCAAGTATGCGAAAAAGGGGAAGAATTGTTTAC
oYS36	AGCTTCTGCAGATCCGCCAGGTGCTCCAGATTTGTAAAGTTCATCCATTCCATGC
oYS37	GATGAACTTTACAAATCTGGAGCACCTGGCGGATCTGCAGAAGCTCCTGAA
oYS598	GCCAGATCCCGGACCTGAGCCCTGTCCAGAGCCGCCGCTGATTTCTAAGGTAGAAAG
oYS599	GAGTGTCTTTTCTTTCTGTAAATAGAAAAGGTGATGTTATGGCAGAAATCGGTACTGGC
V162	GGTGCTTTTTTTGAACGAAAACTCACGTTAAGGGATTTTG
V163	TAGCTGTCAACACTGAGCGTCAGACCCC
V164	ACGCTCAGTGTTGACAGCTAGCTCAGTCCTAGGTATAATACTAG
V165	CGAGACCTAGACTGGTCTCACTAGTATTATACCTAGGACTGAGCTAGCT
V166	ACTAGTGAGACCAGTCTAGGTCTCGTTTAGAGCTAGAAATAGCAAGTTAAAATAAGG
V167	GAGTTTTCGTTCAAAAAAAGCACCGACTCGGT
V272	ctagtCACCACGAACAGAGAATTTGgt
V273	ctagtGTGGTGGAACAGAGAATTTGgt
V274	ctagtGTGGTGCTTGAGAGAATTTGgt
V275	ctagtGTGGTGCTTGTCTCTATTTGgt
V279	tagtAGACCCACGGAAGTAACGGA
V280	taaaacCAAATTCTCTGTTCGTGGTGa
V281	taaaacCAAATTCTCTGTTCCACCACa
V282	taaaacCAAATTCTCTCAAGCACCACa
V283	taaaacCAAATAGAGACAAGCACCACa
V287	aaacTCCGTTACTTCCGTGGGTCT

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