nich was vailable

	bioRxiv preprint doi: https://doi.org/10.1101/398487; this version posted September 3, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC 4.0 International license.
1	Endopeptidase regulation as a novel function of the Zur-dependent zinc starvation response
2	
3	Shannon G. Murphy ^{1, 2} , Laura Alvarez ³ , Myfanwy C. Adams ⁴ , Shuning Liu ^{1, 2} , Joshua S. Chappie ⁴ ,
4	Felipe Cava ³ , Tobias Dörr ^{1, 2} #
5	
6	¹ Department of Microbiology, Cornell University, Ithaca, New York, USA.
7	² Weill Institute for Cell and Molecular Biology, Cornell University, Ithaca, New York, USA.
8	³ Laboratory for Molecular Infection Medicine, Department of Molecular Biology, Umeå University,
9	Umeå, Sweden.
10	⁴ Department of Molecular Medicine, College of Veterinary Medicine, Cornell University, Ithaca, NY,
11	USA.
12	
13	Running Head: Endopeptidase regulation via zinc starvation response
14	
15	#Address correspondence to Tobias Dörr, tdoerr@cornell.edu.
16	
17	Abstract
18	The cell wall is a strong, yet flexible, meshwork of peptidoglycan (PG) that gives a bacterium structural
19	integrity. To accommodate a growing cell, the wall is remodeled by both PG synthesis and degradation.
20	Vibrio cholerae encodes a group of three nearly identical zinc-dependent endopeptidases (EPs) that
21	hydrolyze PG to facilitate cell growth. Two of these (<i>shyA</i> and <i>shyC</i>) are housekeeping genes and form a
22	synthetic lethal pair, while the third (<i>shyB</i>) is not expressed under standard laboratory conditions. To

investigate the role of ShyB, we conducted a transposon screen to identify mutations that activate shyB 23

- transcription. We found that *shyB* is induced as part of the Zur-mediated zinc starvation response, a 24
- 25 mode of regulation not previously reported for cell wall lytic enzymes. In vivo, ShyB alone was

2

sufficient to sustain cell growth in low-zinc environments. *In vitro*, ShyB retained its D,D-endopeptidase
activity against purified sacculi in the presence of the metal chelator EDTA at a concentration that
inhibits ShyA and ShyC. This suggests that ShyB can substitute for the other EPs during zinc starvation,
a condition that pathogens encounter while infecting a human host. Our survey of transcriptomic data
from diverse bacteria identified other candidate Zur-regulated endopeptidases, suggesting that this
adaptation to zinc starvation is conserved in other Gram-negative bacteria.

32

33 Importance

34 The human host sequesters zinc and other essential metals in order to restrict growth of potentially harmful bacteria. In response, invading bacteria express a set of genes enabling them to cope with zinc 35 36 starvation. In Vibrio cholerae, the causative agent of the diarrheal disease cholera, we have identified a 37 novel member of this zinc starvation response: a cell wall hydrolase that retains function in low-zinc environments and is conditionally essential for cell growth. Other human pathogens contain homologs 38 39 that appear to be under similar regulatory control. These findings are significant because they represent. to our knowledge, the first evidence that zinc homeostasis influences cell wall turnover. Anti-infective 40 41 therapies commonly target the bacterial cell wall and, therefore, an improved understanding of how the 42 cell wall adapts to host-induced zinc starvation could lead to new antibiotic development. Such therapeutic interventions are required to combat the rising threat of drug resistant infections. 43

44

45 Introduction

The cell wall provides a bacterium with structural integrity and serves as a protective layer
guarding against a wide range of environmental insults. Due to its importance for bacterial survival, the
cell wall is a powerful and long-standing target for antibiotics (1). The wall is composed primarily of
peptidoglycan (PG), a polymer of β-(1,4) linked N-acetylglucosamine (NAG) and N-acetylmuramic acid
(NAM) sugar strands (2) (Fig. 1A). NAM peptide side chains are cross-linked to peptides on adjacent

3

51 strands, enabling the PG to assemble into a meshlike structure called the sacculus (3). In Gram-negative 52 bacteria, the sacculus is a single PG layer that is sandwiched between an inner and an outer membrane 53 (4). This thin wall must be rigid enough to maintain cell shape and to contain high intracellular pressures (3, 5). However, the wall must also be flexible enough to accommodate cell elongation, cell division, 54 and the insertion of *trans*-envelope protein complexes (6). This requirement for both rigidity and 55 56 flexibility necessitates continuous remodeling of the cell wall, which is accomplished by a delicate 57 interplay between PG synthesis and degradation. Inhibition or dysregulation of process can cause growth 58 cessation or cell lysis, rendering the mechanisms of cell wall turnover an attractive target for new 59 antibiotic development (7, 8).

PG synthesis is mediated by Penicillin Binding Proteins (PBPs, the targets for beta-lactam 60 61 antibiotics) and SEDS proteins (9). These proteins collectively catalyze cell wall synthesis through two 62 main reactions: transplycosylation (TG) to elongate the sugar backbone and transpeptidation (TP) to 63 crosslink the peptide stems of adjacent strands (2). Cell wall turnover is mediated by "autolysins", a 64 collective term for diverse and often redundant enzymes (amidases, carboxypeptidases, lytic transglycosylases and endopeptidases) that are able to cleave PG at almost any chemical bond (6). 65 Endopeptidases (EPs), for example, hydrolyze the peptide crosslinks that covalently link adjacent PG 66 67 strands, effectively reversing the TP reaction. EPs are crucial for cell elongation in both Gram-positive and Gram-negative rod-shaped bacteria (10-12), presumably because they create gaps in the PG 68 69 meshwork to allow for the insertion of new cell wall material. Consistent with this proposed role, EP 70 overexpression promotes aPBP activity in *Escherichia coli*, likely through the generation of initiation 71 sites for PG synthesis (13).

While EPs are essential for growth, they are also main drivers of PG degradation after inhibition of PBPs (14, 15). Thus, EP activity must be tightly controlled under normal growth conditions. EPs in two divergent bacterial species (*E. coli* and *Pseudomonas aeruginosa*) are proteolytically degraded to adapt to conditions that require changes in PG cleavage activity (16, 17), such as the transition into

4

stationary phase. In *Bacillus subtilis*, EP expression is regulated by growth-phase dependent sigma
factors (18-21). However, it is not known how EP expression is modified in response to specific
environmental stresses.

79 In this study, we investigate the role of specialized EPs in V. cholerae, the causative agent of the diarrheal disease cholera. V. cholera encodes three nearly identical EPs that are homologous to the well-80 81 characterized D.D-endopeptidase MepM in E. coli (10). Each EP contains a LysM domain that likely binds PG (22) and a Zn^{2+} -dependent M23 catalytic domain that hydrolyzes peptide cross links (23) (Fig. 82 **1B**). We previously showed that two of these (ShyA and ShyC) are housekeeping EPs that are 83 84 collectively essential for growth (12). The gene encoding the third EP, *shyB*, is not transcribed under standard laboratory conditions (LB medium) and thus little is known about its biological function. To 85 86 elucidate the role of ShyB, we conducted a transposon screen to identify mutations that promote *shyB* 87 expression in LB. We found that *shvB* is induced by zinc starvation and, unlike the other two M23 EPs, ShyB enzymatic activity is resistant to treatment with the metal chelator EDTA. These data suggest that 88 ShyB acts as an alternative EP to ensure proper PG maintenance under zinc limiting conditions. 89 Importantly, this represents the first characterization of an autolysin that is controlled by Zur-mediated 90 91 zinc homeostasis and provides insight into how other Gram-negative pathogens might adapt to zinc-92 starvation when colonizing a human host.

93

94 **Results**

95 *shyB* is repressed in LB, but transcribed in minimal medium.

96 The hydrolytic activity of autolysins needs to be carefully controlled to maintain cell wall 97 integrity. We therefore considered it likely that specialized autolysins are transcriptionally regulated and 98 only induced when required. To test this, we examined expression patterns of the LysM/M23 99 endopeptidases using *lacZ* transcriptional fusions. We first compared promoter activity on LB and M9 100 agar, as our previous work showed that a $\Delta shyB$ mutation exacerbates a $\Delta shyA$ growth defect in M9

5

101 minimal medium (12). The $P_{shyA}:lacZ$ and $P_{shyC}:lacZ$ reporters generated a blue colony color on both LB 102 and M9 minimal agar (**Fig. 1C**), meaning that these promoters are actively transcribed on both media. 103 This is consistent with ShyA and ShyC's role as housekeeping EPs (12). In contrast, $P_{shyB}:lacZ$ yielded 104 blue colonies on M9 minimal medium only, indicating that the *shyB* promoter is induced in M9 but 105 repressed in LB.

106

107 *shyB* is induced by zinc starvation.

108 To elucidate the specific growth conditions that favor *shyB* expression, we sought to identify the 109 genetic factors controlling shyB transcription. To this end, we subjected the transcriptional reporter 110 strain to Himar1 mariner transposon mutagenesis and screened for P_{shvB}:lacZ induction (blue colonies) 111 on LB agar. After two independent rounds of mutagenesis (50,000 total colonies), the screen yielded 26 112 blue colored insertion mutants. These were divided into two distinct classes according to colory color: 113 12 dark blue and 14 light blue colonies. Strikingly, arbitrary PCR (24) mapped all 26 transposon 114 insertions to two chromosomal loci involved in zinc homeostasis: vc0378/zur (dark blue colonies) and 115 vc2081-2083/znuABC (light blue colonies) (Fig. 2A). Zur is a fur-family transcriptional regulator and the central repressor in the zinc starvation response (25). In zinc-rich conditions, Zur and its Zn^{2+} 116 117 corepressor bind to promoters containing a "Zur box" and block transcription (26). In low-zinc 118 conditions, Zur dissociates from promoters to induce the zinc starvation response (27). This regulon 119 includes genes encoding zinc uptake systems (i.e. *znuABC*, *zrgABCDE*) (28) and zinc-independent 120 paralogs that replace proteins that ordinarily require zinc for function (i.e. ribosomal proteins) (29). The 121 Zur-controlled znuABC locus encodes V. cholerae's high affinity zinc uptake system (28). To validate the transposon hits, we constructed clean deletions of *zur* and *znuA* in the P_{shvB}:lacZ reporter strain. 122 123 Deletion of either gene resulted in activation of the *shyB* promoter on LB agar, and *shyB* repression was 124 restored by expressing the respective genes in trans (Fig. S1). Thus, shyB is induced under conditions 125 that are expected to either mimic (zur inactivation) or impose (znuA(BC) inactivation) zinc starvation.

6

126 If zinc starvation is the factor inducing *shyB* expression in M9, we would expect the P_{shyB} :*lacZ* 127 reporter to be repressed by external zinc addition. Indeed, supplementing M9 with 10 μ M of ZnSO₄ was 128 sufficient to turn off the *shvB* promoter in a wild-type (WT) background (Fig. 2B), whereas repression 129 could not be achieved by adding in other transition metals (iron and manganese) (Fig. S2). In a Δzur 130 background, the *shvB* promoter remained active even when M9 was supplemented with exogenous zinc 131 (Fig. 2B), indicating that Zur is required for P_{shvB} repression. We also found that zinc supplementation 132 somewhat repressed the *shvB* promoter in $\Delta znuA$, suggesting that V. *cholerae* can uptake zinc even in 133 the absence of its primary transporter. Indeed, V. cholerae encodes a second, lower affinity zinc 134 acquisition system (*zrgABCDE*) to maintain zinc homeostasis (28).

135

136 Zur directly binds the *shyB* promoter.

Given Zur's well-defined role as a transcriptional regulator (26) and its requirement for P_{shyB} repression in zinc-rich media, we hypothesized that Zur directly binds the *shyB* promoter. To test this, we retrieved a Zur box sequence logo built from 62 known regulatory targets in Vibrionaceae (30, 31) and aligned it with the *shyB* promoter region. This alignment identified a highly conserved Zur box characterized by an inverted, AT-rich repeat (**Fig. 3A**). We used 5'-RACE to locate the *shyB* transcriptional start site (tss) and found that the putative Zur box overlaps both the -10 region and tss. A bound Zur/Zn²⁺ complex at this position likely prevents RNA polymerase binding and thereby prevents transcription (32).

To determine if Zur binds the *shyB* promoter *in vitro*, we incubated purified Zur with a labeled DNA probe encoding the *P_{shyB}* Zur box. Binding was assessed in the presence of ZnCl₂ using an electrophoretic mobility shift assay (EMSA). As evident by a band shift, Zur formed a complex with the *P_{shyB}* DNA *in vitro* (**Fig. 3B, Lanes 1-2**). To examine DNA binding specificity, a 100-fold molar excess of unlabeled specific (S) or non-specific (NS) competitor DNA was included in the binding reaction. The S competitor, which carries an identical sequence as the labeled probe, effectively sequestered Zur and increased the amount of unbound, labeled probe (**Lane 3**). Meanwhile, the NS competitor was

7

ineffective at binding Zur (Lane 4). These data indicate that the *shyB* promoter contains an authentic
Zur box and we conclude that *shyB* is a novel member of the Zur regulon.

153

154 ShyB supports growth in chelated medium.

As *shyB* is part of the Zur-mediated zinc starvation response, we hypothesized that ShyB endopeptidase activity supports cell growth when zinc availability is low. To induce zinc starvation and robustly derepress the Zur regulon, *V. cholerae* strains were grown in M9 minimal medium supplemented with TPEN (N,N,N',N'-Tetrakis(2-pyridylmethyl)ethylenediamine), an intracellular metal chelator with high affinity for zinc (33). As expected from our genetic analysis, TPEN addition resulted in the production of ShyB protein, which could be reversed by adding zinc (**Fig. S3**).

161 We first tested whether native *shyB* could restore $\Delta shyAC$ growth under zinc-starvation

162 conditions. *shyA* and *shyC* deletions were generated in a parent strain expressing an IPTG-inducible

163 copy of *shyA* (*lacZ*:: P_{tac} :*shyA*), as these genes are conditionally essential in rich media (12). In the

absence of IPTG, we found that chelation with either TPEN or EDTA (a more general divalent metal ion

165 chelator), induced growth of $\Delta shyAC$, but not in the mutant that additionally lacked *shyB* (Fig. 4A; Fig.

166 S4). As expected, chelation-dependent growth of $\Delta shyAC$ could be suppressed by adding zinc (Fig. 4B;

167 Fig. S4). These data suggest that induction of *shyB* alone is sufficient to sustain *V. cholerae* growth, and

168 synthetic lethality of *shyA* and *shyC* is due to the lack of *shyB* expression under laboratory growth

169 conditions. Consistent with this interpretation, we were able to generate a $\Delta shyAC$ knockout in a Δzur

170 background (Fig. S5) or in a strain exogenously overexpressing *shyB* (Fig. S6).

171 A $\Delta shyB$ mutant alone did not exhibit a significant growth defect in M9 TPEN (**Fig. 4C**); 172 however, autolysins often need to be deleted in combination to elicit a substantial phenotype (11). We 173 therefore generated all possible combinations of LysM/M23 endopeptidase deletions to broadly dissect 174 the relevance of zinc concentrations for EP activity. Of these, the $\Delta shyAB$ double mutant failed to grow 175 in the presence of TPEN (**Fig. 4C**). This indicates that ShyC, the only essential LysM/M23 EP in the

8

176 $\Delta shyAB$ mutant, cannot support growth in zinc-starved media. In contrast, only the $\Delta shyAC$ mutant 177 failed to grow in zinc-replete medium and this can be explained by Zur-mediated *shyB* repression (Fig. 178 **4D**). This tradeoff in synthetic lethality partners tentatively suggests that ShyB may function as a 179 replacement for ShyC during zinc starvation. ShyC protein levels, as measured by Western Blot, were 180 not reduced in the presence of TPEN, ruling out the possibility that $\Delta shvAB$ lethality reflects 181 transcriptional downregulation or degradation of ShyC (Fig S3). Rather, these observations suggest that 182 ShyC activity is more sensitive to zinc-chelation than the other EPs. Alternatively, TPEN might induce 183 changes in PG architecture that make it resistant to cleavage by ShvC.

184

185 ShyB is an EDTA-resistant D,D-endopeptidase *in vitro*.

186 ShyB is predicted to be a D,D-endopeptidase but biochemical evidence is lacking. Thus, we measured 187 the *in vitro* hydrolytic activity of each EP against V. cholerae sacculi. Each protein was recombinantly purified without the hydrophobic signal sequence or transmembrane domain (ShyA $_{\Lambda 1-35}$, ShyB $_{\Lambda 1-34}$, and 188 189 $ShyC_{\Lambda_1-33}$) to increase stability *in vitro*. As a negative control, we purified ShyB with a mutation 190 (H370A) in the active site that is expected to abolish activity. EPs were incubated with purified sacculi 191 (see Methods for details) and the soluble PG fragments released by digestion were separated using high 192 pressure liquid chromatography (HPLC) and quantified by spectrophotometry. As predicted, all three 193 enzymes, but not the H370A mutant, hydrolyzed V. cholerae sacculi and generated soluble PG 194 fragments (Fig. 5A; Fig. S7). Sacculi digestion with ShyA and ShyC resulted in a similar profile of PG 195 fragments, indicating similar hydrolytic activity in vitro. In contrast, the ShyB chromatogram showed 196 more peaks with shorter retention times. These observations suggest that ShyB further processes the 197 sacculi into smaller fragments. Consistent with this, ShyB was able to further process PG pre-digested 198 with ShyA or ShyC, while these EPs only slightly modified ShyB-digested PG (Fig. S8). To determine 199 which muropeptides remain in the insoluble pellet after EP digestion, muramidase was used to digest the 200 PG sugar backbone (β 1 \rightarrow 4 linkages). The resulting soluble products produced a single peak,

9

201 corresponding to a M4 monomer. This indicates that all three LysM/M23 EPs exhibit D,D-

202 endopeptidase activity *in vitro* (Fig. 5B).

M23 domains require a coordinated zinc ion to carry out PG hydrolysis (23). However, based on its regulation by Zur, we hypothesized that ShyB evolved to function in zinc-limited environments. To test this, we repeated the *in vitro* PG hydrolysis assays under metal-limited conditions by using the divalent cation chelator EDTA. Strikingly, ShyB retained its activity in the presence of EDTA (1 mM), while ShyA and ShyC activity was completely abolished (**Fig. 5C-D**). This is consistent with results previously obtained for ShyA (12). These data suggest that ShyB has a high affinity for, or can function without, divalent cations like zinc.

210

211 ShyB localizes to the division septum.

212 Endopeptidases often differ in their cellular localization and this is an important determinant of EP function *in vivo* (34). We previously reported that ShyA, a sidewall hydrolase, remains diffuse 213 214 throughout the periplasm, while ShvC localizes to the septum (midcell) during division (12). To 215 investigate the relative role of ShyB, we constructed a functional ShyBmsfGFP translational fusion (Fig. 216 **S9**) and visualized its localization using epifluorescence microscopy. ShyBmsfGFP strongly localized to 217 the septum as cells prepared to divide. (Fig. 6). Septal localization suggests that ShyB and ShyC are 218 involved in cell division. However, neither the $\Delta shvB$ nor $\Delta shvC$ mutant, either alone or in combination, 219 have a division defect and we thus do not know the significance of endopeptidase activity at the septum 220 in V. cholerae.

221

222 Zur-controlled endopeptidases are widespread in divergent bacteria.

223 Zur-controlled EPs appear to be widespread in *Vibrionaceae*. Using BLAST homology searches, we

have identified isolates from 30 different non-cholera Vibrio species that contain a ShyB homolog with a

225 Zur box upstream of the gene encoding it (Table S1) (35). To assess the significance of zinc

10

226 homeostasis for EP regulation more broadly, we surveyed published microarray and RNAseq datasets 227 from diverse bacteria for differential EP expression (36-45). Yersinia pestis CO92, the causative agent of 228 plague, encodes a ShyB/MepM homolog (YPO2062) that is significantly up-regulated in a Δzur mutant 229 (36). YPO2062 does not contain its own Zur box, but is adjacent to znuA and may thus be co-transcribed as part of the same operon (Fig. 7). Similarly, *mepM* (b1856) is located adjacent to the *znu* operon in 230 231 laboratory (K12 MG1655) and pathogenic E. coli (Enterohemorrhagic O157:H7 and Enteropathogenic 232 O127:H6). Two independent microarray studies in E. coli, one of which was validated by qPCR, showed that this EP was transcriptionally upregulated in response to zinc starvation (44, 45). These data suggest 233 234 that MepM and its homologs are Zur-regulated EPs. Notably, this *znu*/EP arrangement is conserved in many other Gram-negative pathogens, including Salmonella typhimurium (STM1890), Enterobacter 235 236 cloacae (ECL 0-1442), and Klebsiella pneumoniae (KPK 1913). Lastly, A. baumannii, an important 237 nosocomial pathogen, possesses two M23 endopeptidases differentially transcribed in a Δzur mutant. 238 A1S 3329 is up-regulated and A1S 0820 is down-regulated compared to a wild-type strain (37). 239 suggesting that these EPs are also under zinc starvation control. Collectively, these data suggest that zinc 240 homeostasis and cell wall turnover are linked in a wide array of Gram-negative bacteria.

241

242 Discussion

243 Highly redundant endopeptidases support cell growth.

Endopeptidase activity is essential for cell growth in both Gram-negative and Gram-positive bacteria,
supporting the long-standing hypothesis that autolysins create space in the PG meshwork for the
insertion of new cell wall material (8). As with other autolysins, EPs are highly redundant but exhibit
slight differences in cellular localization (i.e. septal or sidewall) (12, 20, 46), substrate specificity (10,
47) and relative abundance during each growth phase (11, 46). Our previous work in *V. cholerae*identified three LysM/M23 zinc metallo-endopeptidases: two (ShyA and ShyC) are housekeeping
enzymes that are conditionally essential for growth, while the third (ShyB) is not expressed under

11

standard laboratory conditions (12). In this study, we define *shyB* as a new member of the Zur regulon
and demonstrate that ShyB can replace the other EPs *in vivo* when zinc concentrations are limiting. This
is a novel mechanism for regulating autolysins and establishes a link between two essential processes:
cell wall turnover and metal ion homeostasis.

255

256 Zinc availability affects the expression and activity of cell wall hydrolases.

Zur represses *shyB* transcription in zinc-rich growth conditions. This is consistent with our initial observation that the *shyB* promoter is active on M9 and repressed on LB agar. These respective media differ markedly in terms of zinc content; M9 contains no added zinc while LB naturally contains high levels of zinc ions (~12.2 μ M) (48). We found that adding zinc (10 μ M) to M9 represses the *shyB* promoter and hence the zinc starvation response. As a cautionary note, this suggests that *V. cholerae* is starved for zinc in M9, a complication not usually considered when interpreting results obtained in this medium.

Based on its membership in the Zur regulon, it is likely that ShyB evolved to function in low-264 265 zinc environments. Indeed, ShyB endopeptidase activity is resistant to high EDTA concentrations in 266 *vitro*. In an apparent contradiction, the ShyB crystal structure models a zinc ion in the active site (49). It 267 is possible that ShyB has a higher affinity for zinc than the other EPs, but we cannot yet exclude the 268 possibility that ShyB utilizes other metal cofactors. ShyA appears to have an intermediate ability to 269 function in low zinc environments; we found that ShyA can support cell growth in the absence of the 270 other two EPs in TPEN-treated medium (like ShyB), yet EDTA inhibited its activity in vitro. This 271 observation is likely a consequence of the high EDTA concentrations (1 mM) used in the biochemical 272 assays, which do not permit wild type V. cholerae growth. The ability to sustain growth in the presence 273 of TPEN, however, indicates that ShvA function is less affected by metal starvation than ShvC. 274 Since ShyA functions in chelated medium, we tentatively hypothesize that *shyB* is derepressed to 275 compensate for a loss of ShyC activity. This model is supported by localization data and ShyC's

1	2
I	2

276 sensitivity to chelating conditions, both *in vivo* and *in vitro*, that induce *shyB* expression. We did not 277 observe any defects in $\Delta shyB$, $\Delta shyC$, or $\Delta shyBC$ mutants; however, septal EP deletion causes division 278 defects (i.e. filamentation) in other bacteria (20). It is thus possible that the role of septal EPs in *V*. 279 *cholerae* is more prominent under conditions not yet assayed. In our experiments, diffuse ShyA might 280 be present at sufficient concentrations at the septum to alleviate any obvious division defects.

281

282 Bacteria encounter zinc starvation while infecting a host.

283 Proteins that retain function in low-zinc conditions likely play important roles in pathogenesis as 284 bacteria encounter zinc-starvation inside the human host (50). Vertebrates and other organisms sequester metals to restrict the growth of potentially harmful bacterial, a defense strategy referred to as "nutritional 285 286 immunity" (50). In response, bacteria employ zinc-starvation responses to maintain essential cellular 287 processes (51). Zinc importers (*znuABC* and *zrgABCDE*), for example, are critical for host colonization 288 and infection in V. cholerae (28), A. baumannii (52), pathogenic E. coli (53, 54), Salmonella enterica 289 (55), and others (56). It is tempting to speculate that ShyB, a rather unusual addition to the Zur regulon, 290 supports PG remodeling in a zinc-depleted host environment. Consistent with this idea, *shvB* is located 291 on a mobile genomic island (VSP-II) that is strongly associated with the current (seventh) Cholera 292 pandemic (57). The current pandemic strain emerged in the 1960's and, owing to its higher spread 293 capability, replaced its pandemic predecessors (58). This suggests that VSP-II (and thus possibly ShvB) 294 conferred a fitness advantage to pathogenic V. cholerae.

Importantly, Zur-controlled M23 endopeptidases do not appear to be confined to *V. cholerae*. Diverse bacteria, including notable human pathogens, possess a conserved *shyB/mepM/yebA* homolog adjacent to the Zur-controlled *znu* operon. Transcriptomic data from both *Y. pestis* and *E. coli* support the prediction that this EP is upregulated along with the zinc importer. The conservation of zincregulated EPs in divergent Gram-negative pathogens suggests that there may be a widely conserved mechanism for maintaining cell wall homeostasis in low zinc environments. Importantly, this may

13

301 confer an important adaptation to host-induced zinc starvation. These findings in *V. cholerae* will inform
302 future investigations examining the interplay between cell wall turnover and zinc homeostasis.

303

304 **Experimental Procedures**

305 Bacterial growth conditions.

306 Cells were grown by shaking (200 rpm) at 37°C in 5 mL of LB medium unless otherwise indicated. M9 307 minimal medium with glucose (0.4%) was prepared with ultrapure Mili-Q water to minimize zinc 308 contamination. When appropriate, antibiotics were used at the following concentrations: streptomycin 309 (200 μ g mL⁻¹), ampicillin (100 μ g mL⁻¹), and kanamycin (50 μ g mL⁻¹). IPTG (200 μ M) was added to all 310 liquid and solid media if required to sustain *V. cholerae* growth. X-gal (40 μ g mL⁻¹) was added to plates 311 for blue-white screening.

312

313 Plasmid and strain construction.

All genes were PCR amplified from V. cholerae El Tor N16961 genomic DNA. Plasmids were built 314 315 using isothermal assembly (59) with the oligonucleotides summarized in Table S2. The suicide vector 316 pCVD442 was used to make gene deletions via homologous recombination (60); 700 bp regions 317 flanking the gene of interest were amplified for Δzur (SM89/90 + SM91/92), $\Delta znuA$ (SM107/108 + 318 SM109/110), and *\DeltaznuABC* (SM93/94, SM95/96) and assembled into XbaI digested pCVD442. 319 Endopeptidase deletion constructs were built as described previously (12). Chromosomal delivery 320 vectors (pJL-1 and pTD101) were used to insert genes via double cross-over into native lacZ. To 321 construct the shyB transcriptional reporter, 500 bp upstream of shyB were amplified (SM1/2) and assembled into NheI-digested pAM325 to yield a P_{shyB}:lacZ fusion. This fusion was amplified (SM3/4) 322 323 and cloned into StuI-digested pJL-1 (61). To complement gene deletions, zur (SM99/100) and znuA 324 (SM113/114) were cloned into SmaI-digested pBAD: a chloramphenicol resistant, arabinose-inducible plasmid. To construct the ShyBmsfGFP C-terminal translational fusion, shyB (SM181/63) and msfGFP 325

14

326 (SM65/66) were amplified with an overhang encoding a 10 amino acid flexible linker

327 (gctggctccgctgctggttctggcgaattc). These fragments were assembled into SmaI-digested pTD101, which
328 positions the fusion under an IPTG-inducible promoter. In a similar manner, pTD101(*shyB*) was
329 constructed with SM181/182 and pTD100(*shyA*) was built as previously described (12). An additional
330 chromosomal delivery vector (pSGM100) was built for crossover into VC0817. *shyB* (SM141/SM55)
331 was placed under arabinose-inducible control by cloning into SmaI-digested pSGM100. All assemblies
332 were initially transformed into *E. coli* DH5a λpir and then into SM10 λpir for conjugation into *V*.
333 *cholerae.*

334 All strains are derivatives of V. cholerae El Tor N16961 (WT). To conjugate plasmids into V. cholerae, SM10 \pir donor strains carrying pCVD442, pTD101, PJL-1, or pSGM100 plasmids were 335 336 grown in LB/ampicillin. Recipient V. cholerae strains were grown overnight in LB/streptomycin. 337 Stationary phase cells were pelleted by centrifugation (6,500 rpm for 3 min) and washed with fresh LB 338 to remove antibiotics. Equal ratios of donor and recipient (100μ L: 100μ L) were mixed and spotted onto LB agar plates. After a 4-hour incubation at 37°C, cells were streaked onto LB containing streptomycin 339 340 and ampicillin to select for cross-over recipients. Colonies were purified and cured through two rounds 341 of purification on salt free sucrose (10%) agar with streptomycin. Insertions into native *lacZ* (via pJL-1, 342 pTD101) were identified by blue-white colony screening on X-gal plates. Gene deletions (via 343 pCVD442) were checked via PCR screening with the following primers: $\Delta shvA$ (TD503/504), $\Delta shvB$ 344 (SM30/31), ΔshyC (TD701/702), Δzur (SM122/123), ΔznuA (SM119/120), and ΔznuABC (SM119/121). 345

346 Transposon mutagenesis and arbitrary PCR.

The *shyB* transcriptional reporter was mutagenized with Himar1 mariner transposons, which were
delivered via conjugation by an SM10 λpir donor strain carrying pSC189 (62). The recipient and donor
were grown overnight in LB/streptomycin and LB/ampicillin, respectively. Stationary phase cells were
pelleted by centrifugation (6,500 rpm for 3 min) and washed with fresh LB to remove antibiotics. Equal

15

351 ratios of donor and recipient (500 µL:500 µL) were mixed and spotted onto 0.45 µm filter disks adhered 352 to pre-warmed LB plates. After a 4-hour incubation at 37°C, cells were harvested by aseptically 353 transferring the filter disks into conical tubes and vortexing in fresh LB. The cells were spread onto LB 354 agar containing streptomycin to kill the donor strain, kanamycin to select for transposon mutants, and X-355 gal to allow for blue-white colony screening. Plates were incubated at 30°C overnight followed by two 356 days at room temperature. To identify the transposon insertion site, purified colonies were lysed via 357 boiling and used directly as a DNA template for arbitrary PCR. As described elsewhere, this technique amplifies the DNA sequence adjacent to the transposon insertion site through successive rounds of PCR 358 359 (24). Amplicons were Sanger sequenced and high quality sequencing regions were aligned to the N16961 genome using BLAST (35). 360 361 362 5' rapid amplification of cDNA ends. The *shvB* transcription start site was identified with 5' rapid amplification of cDNA ends (5' RACE). To 363 obtain a *shyB* transcript, *Azur* was grown in LB at 37°C until cells reached mid-exponential phase 364 365 $(OD_{600} = 0.5)$ and RNA was extracted using Trizol and acid:phenol chloroform (Ambion). DNA 366 contamination was removed through two RO1 DNase (Promega) treatments and additional acid:phenol chloroform extractions. cDNA synthesis was performed with MultiScribe reverse transcriptase 367 (ThermoFisher) and a *shvB* specific primer (SM270). cDNA was column purified and treated with 368 369 terminal transferase (New England BioLabs) to add a homopolymeric cytosine tail to the 3' end. The

370 cDNA was amplified through two rounds of touchdown PCR with a second gene-specific primer

371 (SM271) and the Anchored Abridged Primer (ThermoFisher). The PCR product was Sanger sequenced

373

372

374 Electrophoretic mobility shift assay.

using primer SM271.

16

375 The LightShift Chemiluminescent EMSA kit (ThermoFisher) was used to detect Zur-promoter binding. 376 41 bp complimentary oligos (SM264/265) containing the putative *shvB* Zur box, with and without a 5' biotin label, were annealed according to commercial instructions (Integrated DNA Technologies). 20 uL 377 378 binding reactions contained buffer, Poly dI-dC (50 ng μ L⁻¹), ZnCl₂ (5 μ M), labeled probe (1 pmol), and 379 purified Zur (600 nM). Unlabeled specific or non-specific competitor oligos were added in 100-fold 380 molar excess. Reactions were incubated on ice for 1 hour, electrophoresed on a 6% DNA retardation gel 381 (100 V, 40 min), and wet transferred to a Biodyne B membrane (100 V, 30 min) (ThermoFisher) in a cold room. The membrane was developed using chemiluminescence according to the manufacturer's 382 383 instructions and imaged using a Bio-Rad ChemiDoc MP imaging system.

384

385 **Protein expression and purification.**

386 DNA encoding N-terminally truncated LysM/M23 endopeptidases (ShyA $_{\Delta 1-35}$, ShyB $_{\Delta 1-34}$, and ShyC $_{\Delta 1-33}$)

and full length Zur was PCR amplified from genomic DNA, while template for the ShyB H370A

388 mutation was commercially synthesized (Integrated DNA Technologies). Shy constructs were cloned

into pCAV4, and Zur into pCAV6, both modified T7 expression vectors that introduce an N-terminal

390 6xHis-NusA tag (pCAV4) or 6xHis-MBP tag (pCAV6) followed by a Hrv3C protease site upstream of

the inserted sequence. Constructs were transformed into BL21(DE3) cells, grown at 37°C in Terrific

392 Broth supplemented with carbenicillin (100 mg mL^{-1}) to an OD₆₀₀ of 0.8-1.0, and then induced with

393 IPTG (0.3 mM) overnight at 19°C. ZnCl₂ (50 μM) was added during Zur induction. Cells were

harvested via centrifugation, washed with nickel loading buffer (NLB) (20 mM HEPES pH 7.5, 500 mM

395 NaCl, 30 mM imidazole, 5% glycerol (v:v), 5 mM β –Mercaptoethanol), pelleted in 500mL aliquots, and 396 stored at -80°C.

397 Pellets were thawed at 37°C and resuspended in NLB supplemented with PMSF (10 mM),

398 DNAse (5 mg), MgCl₂ (5 mM), lysozyme (10 mg mL⁻¹), and one tenth of a complete protease inhibitor

399 cocktail tablet (Roche). All buffers used in Zur purification were supplemented with ZnCl₂ (1 µM). Cell

1	7

400	suspensions were rotated at 4°C, lysed via sonication, centrifuged, and the supernatant was syringe
401	filtered using a 0.45 μ M filter. Clarified samples were loaded onto a NiSO ₄ charged 5 mL HiTrap
402	chelating column (GE Life Sciences), and eluted using an imidazole gradient from 30 mM to 1M.
403	Hrv3C protease was added to the pooled fractions and dialyzed overnight into cation exchange loading
404	buffer (20 mM HEPES pH 7.5, 50 mM NaCl, 1 mM EDTA, 5% glycerol (v:v), 1 mM DTT). Cleaved
405	Shy proteins were loaded onto a 5 mL HiTrap SP HP column and cleaved Zur was loaded onto a 5mL
406	HiTrap Heparin HP column (GE Life Sciences). All constructs were eluted along a NaCl gradient from
407	50mM to 1M. Fractions were concentrated and injected onto a Superdex 75 16/600 equilibrated in Size
408	Exclusion Chromotography buffer (20 mM HEPES pH7.5, 150 mM KCl, 1 mM DTT). Zur dimers
409	coeluted with MBP on the sizing column and were subsequently incubated with amylose resin (New
410	England BioLabs) at 4°C and Zur was collected from a gravity column. Final purified protein
411	concentrations were determined by SDS-PAGE and densitometry compared against BSA standards:
412	ShyA, 5.72 mg mL ⁻¹ ; ShyB, 5.72 mg mL ⁻¹ ; ShyB H320A, 2.35 mg mL ⁻¹ ; ShyC, 17.93 mg mL ⁻¹ ; Zur,
413	0.31 mg mL^{-1} .

414

415 Sacculi digestion assay.

Peptidoglycan from stationary phase V. cholerae cells was extracted and purified via SDS boiling and 416 417 muramidase digestion (63). 10 µL of sacculi and 10 µg of enzyme were mixed in 50 µL buffered solution (50 mM Tris-HCl pH 7.5, 100 mM NaCl) in the absence or presence of 1 mM EDTA. 418 419 Digestions were incubated for 16 h at 37°C. Soluble products were harvested and the remaining pellet 420 was further digested with muramidase. All soluble products were reduced with sodium borohydride, 421 their pH adjusted, and injected into a Waters UPLC system (Waters, Massachusetts, USA) equipped 422 with an ACQUITY UPLC BEH C18 Column, 130 Å, 1.7 µm, 2.1 mm × 150 mm (Waters) and a dual wavelength absorbance detector. Eluted fragments were separated at 45°C using a linear gradient from 423

18

- 424 buffer A [formic acid 0.1% (v/v)] to buffer B [formic acid 0.1% (v/v), acetonitrile 40% (v/v)] in a 12 425 min run with a 0.175 ml min⁻¹ flow, and detected at 204 nm.
- 426

427 Growth curve analysis.

- 428 Strains were grown overnight in LB/streptomycin with IPTG. Cells were washed in 1X phosphate
- 429 buffered solution (PBS) and subcultured 1:10 into M9 glucose plus IPTG. After 2 hours shaking at 37°C,
- 430 cells were washed and subcultured 1:100 into M9 glucose containing combinations of TPEN (250 nM),
- 431 ZNSO4 (1 μM), and IPTG (200 μM). The growth of each 200 μL culture in a 100-well plate was
- 432 monitored by optical density (OD₆₀₀) on a Bioscreen C plate reader (Growth Curves America).
- 433

434 Microscopy and image analysis.

- 435 Cells were imaged on an agarose patch (0.8% agarose in M9 minimal medium) using a Leica DMi8
 436 inverted microscope. To image the ShyBmsfGFP fusion, cells were exposed to 490 nm for 300 ms.
- 437 Image analysis, including cell selection and subpixel quantification of fluorescent signal as a function
- 438 distance from the midcell, was performed in Oufti (64).
- 439

440 Acknowledgements

We thank all the members of the Doerr and Chappie Labs for helpful discussions and assistance with
this work. We thank members of the John Helmann Lab for lending their expertise and reagents. We
thank the faculty, staff, and students at the Weill Institute for Cell and Molecular Biology (WICMB) for
their support. Research in the Doerr Lab is supported by start-up funds from Cornell University.
Research in the Chappie Lab is supported by the National Institute of Health (NIH). Research in the
Cava lab is supported by MIMS, the Knut and Alice Wallenberg Foundation (KAW), the Swedish
Research Council and the Kempe Foundation.

448

449

450 <u>References</u>

451	1.	Schneider T, Sahl H-G. 2010. An oldie but a goodie – cell wall biosynthesis as antibiotic targe
452		pathway. Int J Med Microbiol 300 :161–169.

453	2.	Typas A, Banzhaf M, Gross CA, Vollmer W. 2012. From the regulation of peptidoglycan

- 454 synthesis to bacterial growth and morphology. Nat Rev Microbiol **10**:123–136.
- 455 3. Höltje JV. 1998. Growth of the stress-bearing and shape-maintaining murein sacculus of
- 456 *Escherichia coli*. Microbiol Mol Biol Rev **62**:181–203.
- 457 4. Silhavy TJ, Kahne D, Walker S. 2010. The bacterial cell envelope. Cold Spring Harb Perspect
 458 Biol 2:a000414.
- 459 5. Cabeen MT, Jacobs-Wagner C. 2005. Bacterial cell shape. Nat Rev Microbiol 3:601–610.
- 460 6. Vollmer W, Joris B, Charlier P, Foster S. 2008. Bacterial peptidoglycan (murein) hydrolases.
 461 FEMS Microbiol Rev 32:259–286.
- 462 7. Tomasz A. 1979. The mechanism of the irreversible antimicrobial effects of penicillins: how the
 463 beta-lactam antibiotics kill and lyse bacteria. Annu Rev Microbiol 33:113–137.
- 464 8. Vollmer W. 2012. Bacterial growth does require peptidoglycan hydrolases. Mol Microbiol
 465 86:1031–1035.
- 466 9. Cho H, Wivagg CN, Kapoor M, Barry Z, Rohs PDA, Suh H, Marto JA, Garner EC,

467 Bernhardt TG. 2016. Bacterial cell wall biogenesis is mediated by SEDS and PBP polymerase
468 families functioning semi-autonomously. Nature Microbiology 2016 1:10 1:16172.

10.

469

Singh SK, SaiSree L, Amrutha RN, Reddy M. 2012. Three redundant murein endopeptidases

n	n
2	υ
	_

470 471		catalyse an essential cleavage step in peptidoglycan synthesis of <i>Escherichia coli</i> K12. Mol Microbiol 86 :1036–1051.
, ,,,		WICTOOLOG 60 .1050–1051.
472	11.	Hashimoto M, Ooiwa S, Sekiguchi J. 2011. The synthetic lethality of lytE cwlO in Bacillus
473		subtilis is caused by lack of d,l-endopeptidase activity at the lateral cell wall. J Bacteriol 194:796-
474		803.
475	12.	Dörr T, Cava F, Lam H, Davis BM, Waldor MK. 2013. Substrate specificity of an elongation-
476		specific peptidoglycan endopeptidase and its implications for cell wall architecture and growth of
477		Vibrio cholerae. Mol Microbiol 89:949–962.
478	13.	Lai GC, Cho H, Bernhardt TG. 2017. The mecillinam resistome reveals a role for
479		peptidoglycan endopeptidases in stimulating cell wall synthesis in Escherichia coli. PLoS Genet
480		13 :e1006934.
481	14.	Dörr T, Davis BM, Waldor MK. 2015. Endopeptidase-Mediated Beta Lactam Tolerance. PLoS
482		Pathog 11:e1004850.
483	15.	Kitano K, Tuomanen E, Tomasz A. 1986. Transglycosylase and endopeptidase participate in
484		the degradation of murein during autolysis of <i>Escherichia coli</i> . J Bacteriol 167 :759–765.
485	16.	Singh SK, Parveen S, SaiSree L, Reddy M. 2015. Regulated proteolysis of a cross-link-specific
486		peptidoglycan hydrolase contributes to bacterial morphogenesis. Proc Natl Acad Sci USA
487		112 :10956–10961.
488	17.	Srivastava D, Seo J, Rimal B, Kim SJ, Zhen S, Darwin AJ. 2018. A Proteolytic Complex
489		Targets Multiple Cell Wall Hydrolases in Pseudomonas aeruginosa. mBio 9:e00972–18.

\mathbf{r}	1
2	

490 18. Britton RA, Eichenberger P, Gonzalez-Pastor JE, Fawcett P, Monson R, Losick R,

- 491 **Grossman AD**. 2002. Genome-wide analysis of the stationary-phase Sigma factor (Sigma-H)
- 492 Regulon of *Bacillus subtilis*. J Bacteriol **184**:4881–4890.
- 493 19. Ishikawa S, Hara Y, Ohnishi R, Sekiguchi J. 1998. Regulation of a new cell wall hydrolase
- 494 gene, cwlF, which affects cell separation in *Bacillus subtilis*. J Bacteriol **180**:2549–2555.
- 495 20. Ohnishi R, Ishikawa S, Sekiguchi J. 1999. Peptidoglycan hydrolase LytF plays a role in cell
 496 separation with CwlF during vegetative growth of *Bacillus subtilis*. J Bacteriol 181:3178–3184.
- 497 21. Yamaguchi H, Furuhata K, Fukushima T, Yamamoto H, Sekiguchi J. 2004. Characterization
- 498 of a new *Bacillus subtilis* peptidoglycan hydrolase gene, yvcE (named cwlO), and the enzymatic
 499 properties of its encoded protein. J Biosci Bioeng **98**:174–181.
- 500 22. Buist G, Steen A, Kok J, Kuipers OP. 2008. LysM, a widely distributed protein motif for
 501 binding to (peptido)glycans. Mol Microbiol 68:838–847.
- Rawlings ND, Barrett AJ, Thomas PD, Huang X, Bateman A, Finn RD. 2018. The MEROPS
 database of proteolytic enzymes, their substrates and inhibitors in 2017 and a comparison with
 peptidases in the PANTHER database. Nucleic Acids Res 46:D624–D632.
- 505 24. O'Toole GA, Pratt LA, Watnick PI, Newman DK, Weaver VB, Kolter R. 1999. Genetic
 506 approaches to study of biofilms. Methods Enzymol 310:91–109.
- 507 25. Patzer SI, Hantke K. 1998. The ZnuABC high-affinity zinc uptake system and its regulator Zur
 508 in *Escherichia coli*. Mol Microbiol 28:1199–1210.
- 509 26. Gilston BA, Wang S, Marcus MD, Canalizo-Hernández MA, Swindell EP, Xue Y,
- 510 Mondragón A, O'Halloran TV. 2014. Structural and Mechanistic Basis of Zinc Regulation

511 Across the *E. coli* Zur Regulon. PLoS Biol **12**:e1001987.

512 27. Shin J-H, Helmann JD. 2016. Molecular logic of the Zur-regulated zinc deprivation response in
513 *Bacillus subtilis*. Nat Commun 7:1–9.

514 28. Sheng Y, Fan F, Jensen O, Zhong Z, Kan B, Wang H, Zhu J. 2015. Dual zinc transporter

- 515 systems in *Vibrio cholerae* promote competitive advantages over gut microbiome. Infect Immun
 516 83:3902–3908.
- 517 29. Panina EM, Mironov AA, Gelfand MS. 2011. Comparative genomics of bacterial zinc

518 regulons: Enhanced ion transport, pathogenesis, and rearrangement of ribosomal proteins. Proc

- 519 Natl Acad Sci USA **100**:9912–9917.
- 30. Novichkov PS, Kazakov AE, Ravcheev DA, Leyn SA, Kovaleva GY, Sutormin RA, Kazanov
 MD, Riehl W, Arkin AP, Dubchak I, Rodionov DA. 2013. RegPrecise 3.0 A resource for
 genome-scale exploration of transcriptional regulation in bacteria. BMC Genomics 14:745.
- 523 31. Crooks GE, Hon G, Chandonia J-M, Brenner SE. 2004. WebLogo: a sequence logo generator.
 524 Genome Res 14:1188–1190.
- 32. Napolitano M, Rubio MÁ, Camargo S, Luque I. 2013. Regulation of internal promoters in a
 zinc-responsive operon is influenced by transcription from upstream promoters. J Bacteriol
 195:JB.01488–12–1293.
- Arslan P, Di Virgilio F, Beltrame M, Tsien RY, Pozzan T. 1985. Cytosolic Ca2+ homeostasis
 in Ehrlich and Yoshida carcinomas. A new, membrane-permeant chelator of heavy metals reveals
 that these ascites tumor cell lines have normal cytosolic free Ca²⁺. J Biol Chem 260:2719–2727.
- 531 34. Yamamoto H, Kurosawa S-I, Sekiguchi J. 2003. Localization of the vegetative cell wall

532		hydrolases LytC, LytE, and LytF on the Bacillus subtilis cell surface and stability of these
533		enzymes to cell wall-bound or extracellular proteases. J Bacteriol 185:6666–6677.
534	35.	Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. 1990. Basic local alignment search
535		tool. J Mol Biol 215 :403–410.
536	36.	Li Y, Qiu Y, Gao H, Guo Z, Han Y, Song Y, Du Z, Wang X, Zhou D, Yang R. 2009.
537		Characterization of Zur-dependent genes and direct Zur targets in Yersinia pestis. BMC Microbiol
538		9 :128.
539	37.	Mortensen BL, Rathi S, Chazin WJ, Skaar EP. 2014. Acinetobacter baumannii response to
540		host-mediated zinc limitation requires the transcriptional regulator Zur. J Bacteriol
541		196 :JB.01650–14–2626.
542	38.	Maciąg A, Dainese E, Rodriguez GM, Milano A, Provvedi R, Pasca MR, Smith I, Palù G,
543		Riccardi G, Manganelli R. 2007. Global analysis of the Mycobacterium tuberculosis Zur (FurB)
544		regulon. J Bacteriol 189:730–740.
545	39.	Pawlik M-C, Hubert K, Joseph B, Claus H, Schoen C, Vogel U. 2012. The zinc-responsive
546		regulon of Neisseria meningitidis comprises seventeen genes under control of a Zur element. J
547		Bacteriol 194 :JB.01091–12–6603.
548	40.	Mazzon RR, Braz VS, da Silva Neto JF, do Valle Marques M. 2014. Analysis of the
549		Caulobacter crescentus Zur regulon reveals novel insights in zinc acquisition by TonB-dependent
550		outer membrane proteins. BMC Genomics 15:734.
551	41.	Gaballa A, Wang T, Ye RW, Helmann JD. 2002. Functional analysis of the Bacillus subtilis
552		Zur regulon. J Bacteriol 184:6508-6514.

553	42.	Schröder J, Jochmann N, Rodionov DA, Tauch A. 2010. The Zur regulon of Corynebacterium
554		glutamicum ATCC 13032. BMC Genomics 11:12.
555	43.	Kallifidas D, Pascoe B, Owen GA, Strain-Damerell CM, Hong H-J, Paget MSB. 2010. The
556		zinc-responsive regulator Zur controls expression of the coelibactin gene cluster in Streptomyces
557		coelicolor. J Bacteriol 192:608–611.
558	44.	Sigdel TK, Easton JA, Crowder MW. 2006. Transcriptional response of Escherichia coli to
559		TPEN. J Bacteriol 188:6709-6713.
560	45.	Hensley MP, Gunasekera TS, Easton JA, Sigdel TK, Sugarbaker SA, Klingbeil L, Breece
561		RM, Tierney DL, Crowder MW. 2012. Characterization of Zn(II)-responsive ribosomal
562		proteins YkgM and L31 in E. coli. J Inorg Biochem 111:164–172.
563	46.	Fukushima T, Afkham A, Kurosawa S-I, Tanabe T, Yamamoto H, Sekiguchi J. 2006. A New
564		d,l-endopeptidase gene product, YojL (renamed CwlS), plays a role in cell separation with LytE
565		and LytF in Bacillus subtilis. J Bacteriol 188:5541-5550.
566	47.	Smith TJ, Blackman SA, Foster SJ. 2000. Autolysins of Bacillus subtilis: multiple enzymes
567		with multiple functions. Microbiology 146 :249–262.
568	48.	Takahashi H, Oshima T, Hobman JL, Doherty N, Clayton SR, Iqbal M, Hill PJ, Tobe T,
569		Ogasawara N, Kanaya S, Stekel DJ. 2015. The dynamic balance of import and export of zinc in
570		Escherichia coli suggests a heterogeneous population response to stress. J R Soc Interface
571		12 :20150069–20150069.
572	49.	Ragumani S, Kumaran D, Burley SK, Swaminathan S. 2008. Crystal structure of a putative
573		lysostaphin peptidase from Vibrio cholerae. Proteins: Struct, Funct, Bioinf 72:1096–1103.

574	50.	Kehl-Fie TE, Skaar EP. 2010. Nutritional immunity beyond iron: a role for manganese and zinc.
575		Current Opinion in Chemical Biology 14:218–224.
576	51.	Ma L, Terwilliger A, Maresso AW. 2015. Iron and zinc exploitation during bacterial
577		pathogenesis. Metallomics 7:1541–1554.
578	52.	Hood MI, Mortensen BL, Moore JL, Zhang Y, Kehl-Fie TE, Sugitani N, Chazin WJ,
579		Caprioli RM, Skaar EP. 2012. Identification of an Acinetobacter baumannii zinc acquisition
580		system that facilitates resistance to calprotectin-mediated zinc sequestration. PLoS Pathog
581		8 :e1003068.
582	53.	Sabri M, Houle S, Dozois CM. 2009. Roles of the extraintestinal pathogenic Escherichia coli
583		ZnuACB and ZupT Zinc transporters during urinary tract infection. Infect Immun 77:1155–1164.
584	54.	Gabbianelli R, Scotti R, Ammendola S, Petrarca P, Nicolini L, Battistoni A. 2011. Role of
585		ZnuABC and ZinT in Escherichia coli O157:H7 zinc acquisition and interaction with epithelial
586		cells. BMC Microbiol 11:36.
587	55.	Ammendola S, Pasquali P, Pistoia C, Petrucci P, Petrarca P, Rotilio G, Battistoni A. 2007.
588		High-affinity Zn ²⁺ uptake system ZnuABC is required for bacterial zinc homeostasis in
589		intracellular environments and contributes to the virulence of Salmonella enterica. Infect Immun
590		75 :5867–5876.
591	56.	Cerasi M, Ammendola S, Battistoni A. 2013. Competition for zinc binding in the host-pathogen
592		interaction. Front Cell Infect Microbiol 3 .
593	57.	O'Shea YA, Reen FJ, Quirke AM, Boyd EF. 2004. Evolutionary genetic analysis of the
594		emergence of epidemic Vibrio cholerae isolates on the basis of comparative nucleotide sequence
595		analysis and multilocus virulence gene profiles. J Clin Microbiol 42:4657–4671.

596	58.	Hu D, Liu B, Feng L, Ding P, Guo X, Wang M, Cao B, Reeves PR, Wang L. 2016. Origins of		
597		the current seventh cholera pandemic. Proc Natl Acad Sci USA 113:E7730-E7739.		
598	59.	Gibson DG, Young L, Chuang R-Y, Venter JC, Hutchison CA III, Smith HO. 2009.		
599		Enzymatic assembly of DNA molecules up to several hundred kilobases. Nat Methods 6:343–		
600		345.		
601	60.	Donnenberg MS, Kaper JB. 1991. Construction of an eae deletion mutant of enteropathogenic		
602		<i>Escherichia coli</i> by using a positive-selection suicide vector. Infect Immun 59 :4310–4317.		
603	61.	Butterton JR, Beattie DT, Gardel CL, Carroll PA, Hyman T, Killeen KP, Mekalanos JJ,		
604		Calderwood SB. 1995. Heterologous antigen expression in Vibrio cholerae vector strains. Infect		
605		Immun 63 :2689–2696.		
606	62.	Chiang SL, Rubin EJ. 2002. Construction of a mariner-based transposon for epitope-tagging and		
607		genomic targeting. Gene 296 :179–185.		
608	63.	Alvarez L, Hernandez SB, de Pedro MA, Cava F. 2016. Ultra-sensitive, high-resolution liquid		
609		chromatography methods for the high-throughput quantitative analysis of bacterial cell wall		
610		chemistry and structure. pp. 11–27. In Bacterial Cell Wall Homeostasis. Humana Press, New		
611		York, NY, New York, NY.		
612	64.	Paintdakhi A, Parry B, Campos M, Irnov I, Elf J, Surovtsev I, Wagner CJ. 2016. Oufti: an		
613		integrated software package for high-accuracy, high-throughput quantitative microscopy analysis.		
614		Mol Microbiol 99 :767–777.		
615	65.	Apweiler R, Bairoch A, Wu CH, Barker WC, Boeckmann B, Ferro S, Gasteiger E, Huang		
616		H, Lopez R, Magrane M, Martin MJ, Natale DA, O'Donovan C, Redaschi N, Yeh LSL.		
617		2004. UniProt: the universal protein knowledgebase. Nucleic Acids Res 32:D115-D119.		

27

66. Fong C, Rohmer L, Radey M, Wasnick M, Brittnacher MJ. 2008. PSAT: A web tool to 619 compare genomic neighborhoods of multiple prokaryotic genomes. BMC Bioinformatics 9:170.

620

621 Figure Legends

622 Fig 1. *shyB* is a LysM/M23 endopeptidase that is transcribed in minimal medium.

- 623 (A) Model of the peptidoglycan sacculus indicating EP cleavage sites (B) The V. cholerae genome
- encodes three endopeptidases (ShyA, ShyB, ShyC) possessing a hydrophobic region (gray), a PG
- binding domain (LysM, pink), and metallo-endopeptidase domain (M23, green). Protein domains were
- 626 annotated using UniProt (65). (C) *lacZ* transcriptional reporters for each endopeptidase spotted onto LB
- 627 (top row) and M9 minimal (bottom row) agar containing X-gal. A blue colony color indicates that the
- 628 promoter is actively transcribed. Wild-type (WT) and $\Delta lacZ$ strain are included as positive and negative
- 629 controls, respectively.
- 630

631 Fig 2. *shyB* transcription is regulated by zinc homeostasis.

632 (A) The *shyB* transcriptional reporter (*lacZ::P_{shyB}:lacZ*) was mutagenized with a Himar1 mariner 633 transposon and screened for *shyB* induction (blue colonies) on LB agar containing X-gal and selective 634 antibiotics (see Methods). Representative dark blue (black arrow) and light blue (white arrows) colonies 635 are shown. Approximate Tn insertion sites identified by arbitrary PCR are shown (triangles). (B) The 636 *shyB* transcriptional reporter in a wild-type, Δzur , or $\Delta znuABC$ background were grown on M9 X-gal 637 agar without (top row) or with (bottom row) 10 μ M ZnSO₄.

638

639 Fig 3. Zur directly binds the *shyB* promoter.

640 (A) The *shyB* promoter, annotated with a 5'-RACE transcription start site (+1) and putative -10 region

641 (box), was aligned with a *Vibrio* Zur sequence logo (30, 31). The inverted AT-rich repeat in the putative

642	Zur-box is underlined with black arrows. (B) A chemiluminescent probe containing the putative <i>shyB</i>
643	Zur box was incubated with purified Zur in the presence of $ZnCl_2$ (5 μ M). Zur binding specificity was
644	tested by adding 100-fold molar excess of unlabeled specific (S, Lane 3) or non-specific (NS, Lane 4)
645	competitor DNA. Samples were electrophoresed on a 6% DNA retardation gel to separate unbound
646	(black arrow) and bound probe (white arrow).
647	
648	Fig 4. <i>shyB</i> supports cell growth in chelated medium and is conditionally essential in a $\Delta shyA$
649	mutant.
650	Mid-exponential cultures of the indicated V. cholerae mutants were washed to remove IPTG before
651	being diluted 1:100 into M9 glucose containing streptomycin plus TPEN (250 nM) in the absence (A,C)
652	or presence (B,D) of ZnSO ₄ (1 µM). Growth of each strain was monitored by optical density (OD600) in

653 a Bioscreen C 100-well plate. Error bars report standard error of the mean (SEM) for three independent

biological replicates. (A-B) Log-transformed growth curves are shown for WT (green circle). $\Delta shvAC$

lacZ:: P_{tac} : *shyA* (blue square), and $\Delta shyABC$ *lacZ*:: P_{tac} : *shyA* (red triangle). (C-D) In a similar growth

experiment, growth rates of WT (solid black), single mutants (solid gray) and double mutants (striped) 656

657 were calculated from exponential phase and normalized to the average WT growth rate (%). Statistical

658 difference relative to WT was assessed using two-way analysis of variance (ANOVA) followed by

Dunnett's multiple comparison test (***, p-value < 0.001). 659

660

654

655

661 Fig 5. ShyB retains endopeptidase activity in EDTA.

V. cholerae sacculi was digested with 10 µg of purified ShyA, ShyB, or ShyC for 16 h at 37°C in the 662 absence (A) or presence (C) of 1mM EDTA. The soluble products released by digested sacculi were 663 separated by size via HPLC and quantified by absorbance (204 nm). (**B**, **D**) The remaining insoluble 664 665 pellet was digested with muramidase and soluble products were separated by HPLC.

666

	not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC 4.0 International license.
667	Fig 6. ShyBmsfGFP localizes to the midcell during division.
668	V. cholerae expressing a C-terminal fluorescent fusion ($lacZ::P_{tac}$ -shyBmsfGFP) was grown overnight in
669	M9 + IPTG (200 μ M). (A) The ShyBmsfGFP fusion was imaged on an agarose patch (0.8% agarose in
670	M9 minimal medium) with 300 ms exposure at 490 nm. (B) A heat map showing intensity of fluorescent
671	signal as a function of distance from the midcell ("demograph") was generated from over 1,800 cells in
672	Oufti (64).
673	
674	Fig 7. ShyB/MepM homologs are adjacent to the Zur-controlled <i>znu</i> operon in many Gram-
675	negative pathogens.
676	Gene neighborhood alignments generated by Prokaryotic Sequence Homology Analysis Tool (PSAT)
677	from 7 different Gram-negative bacteria (66). Arrows indicate the approximate location of the
678	bidirectional promoter and site of Zur-binding in the znu operon. Asterisks indicate that co-transcription
679	of znu and the downstream M23 endopeptidase is supported by transcriptomic data.
680	
681	Fig S1. $\triangle zur$ and $\triangle znuABC$ deletions induce the <i>shyB</i> promoter on LB agar.
682	Clean deletions of Δzur and $\Delta znuA$ in the P_{shyB} : lacZ transcriptional reporter were complemented with an
683	arabinose-inducible (pBAD) plasmid carrying the respective gene in trans. Strains were plated onto LB
684	agar containing x-gal (40 μ g mL ⁻¹), chloramphenicol (10 μ g mL ⁻¹), and arabinose (0.2%). Plates were
685	incubated overnight and then at room temperature for 2 days.
686	

Fig S2. *shyB* promoter is repressed by exogenous zinc, but not by other transition metals. 687

The P_{shvB}:lacZ transcriptional reporter was plated on M9 X-gal agar containing 10 µM of ZnSO₄, FeSO₄, 688

or MnCl₂. Plates were incubated overnight and then at room temperature for 2 days. 689

690

Fig S3. Western Blot of ShyB and ShyC protein levels in high and low-zinc media. 691

\mathbf{a}	1	٦	
5	ι	J	
-		-	

692	N16961 strains encoding ShyB
693	(0.4%) with added TPEN (250 nM) or TPEN plus ZnSO4 (1 μ M). Cells were harvested at mid-log
694	(OD600 = 0.4) and lysed via SDS boiling and sonication. Western blot was performed using standard
695	techniques. Blots were developed using a mouse anti-FLAG F1804 primary antibody (Sigma Aldrich)
696	and Goat anti-Mouse IR CW800 secondary antibody (LI-COR Biosciences). Blots were imaged using a
697	Lycor Odyssey CLx imager.
698	
699	Fig S4. EDTA-induced <i>shyB</i> expression restores growth to $\triangle shyAC$.
700	Wt (green), $\Delta shyAC lacZ::P_{tac}$ -shyA (blue), and $\Delta shyABC lacZ::P_{tac}$ -shyA (red) strains were grown in
701	M9 glucose (0.4%) containing (A) EDTA (30 μ M) (solid lines) or (B) EDTA plus ZnSO ₄ (60 μ M)
702	(dashed lines). Growth of each 200 μL culture was measured by optical density (600 nm) in a Bioscreen
703	C 100-well plate. Error bars report standard error of the mean (SEM) for three biologically independent
704	replicates.
705	
706	Fig S5. <i>zur</i> deletion restores growth to $\triangle shyAC$ in LB medium.
707	Overnight cultures (grown in LB/streptomycin at 37°C) were subcultured 1:100 into fresh media and
708	grown at 37°C until mid-log phase. $\Delta zur lacZ::P_{tac}-zur$ (blue) and $\Delta zur \Delta shyAC lacZ::P_{tac}-zur$ (red) were
709	diluted 1:100 into LB (solid lines) or in LB plus IPTG (200 μ M) (dashed lines). Growth of each strain
710	was monitored by optical density (OD600) in a Bioscreen C 100-well plate. Error bars report standard
711	error of the mean (SEM) for three biologically independent replicates.
712	
713	Fig S6. Inducible <i>shyB</i> expression rescues growth of $\triangle shyABC$.
714	Strains were grown overnight in LB/streptomycin plus IPTG (200uM) at 37°C. Cells were washed,
715	subcultured 1:10 into M9 glucose (0.4%), and grown at 37°C for 2 hours. Wt (dotted lines) and
716	$\Delta shyABC \ lacZ::P_{tac}-shyA \ vc1807::P_{ara}-shyB$ (solid lines) strains were diluted 1:100 in M9 glucose

31

at

717	(0.4%) (orange), with 200 μ M IPTG (green) or with 0.2% arabinose (black). Growth of each 200 μ L		
718	culture was measured by optical density (600 nm) in a Bioscreen C 100-well plate. Error bars report		
719	standard error of the mean (SEM) for three biologically independent replicates.		
720			
721	Fig S7. A point mutation in the ShyB active site abolishes endopeptidase activity <i>in vitro</i> .		
722	Purified ShyB, and ShyB H370A were incubated with purified V. cholerae sacculi for 16 h at 37°C. (A)		
723	The soluble products released by digested were separated by HPLC and quantified by absorbance (204		
724	nm). (B) The remaining pellet was digested with muramidase and the soluble products were separated		
725	by HPLC and quantified by absorbance.		
726			
727	Fig S8. Sequential digestion of V. cholerae sacculi by Shy endopeptidases.		
728	10 µg of purified (A) ShyA, (B) ShyB, and (C) ShyC were incubated with V. cholerae sacculi for 16 h a		
729	37°C, followed by secondary digestion a different endopeptidase. The soluble products released by		
730	digested sacculi were separated by size via HPLC and quantified by absorbance (204 nm).		
731			
732	Fig S9. ShyBmsfGFP translational fusion rescues growth of $\Delta shyAB$ in TPEN-chelated medium.		
733	Mid-exponential cultures of WT (blue), $\Delta shyAB$ (red), $\Delta shyAB$ lacZ:: P_{tac} -shyB (green), and $\Delta shyAB$		
734	<i>lacZ::P_{tac}-shyB</i> (orange, purple) were washed and subcultured 1:100 into M9 containing TPEN (250		
735	nM) with and without IPTG (200 μ M). Growth of each culture at 37°C was measured by optical density		
736	(600 nm). Error bars report standard error of the mean (SEM) for three biologically independent		
737	replicates.		
738			
739	Table S1. Summary of ShyB homologs that contain an upstream, canonical Zur box.		
740			

741 Table S2. Summary of oligonucleotides used in this study.

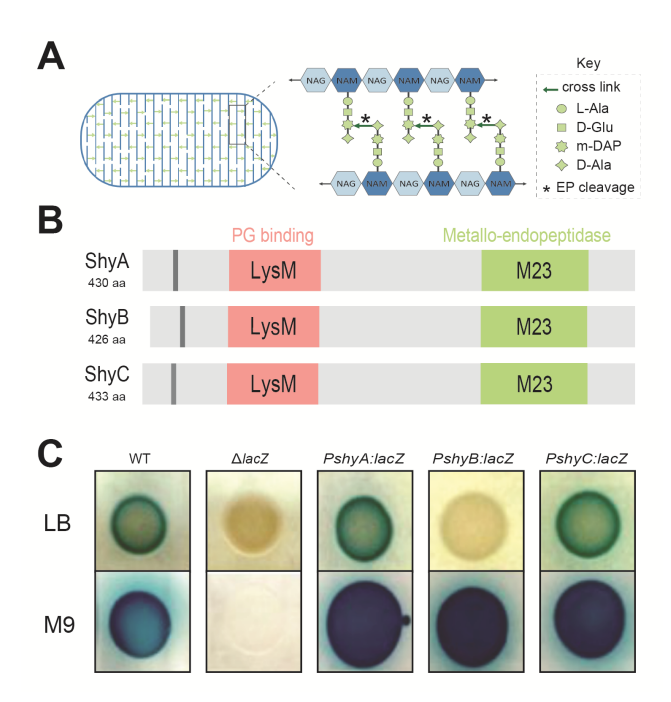


Fig 1. *shyB* is a LysM/M23 endopeptidase that is transcribed in minimal medium.

(A) Model of the peptidoglycan sacculus indicating EP cleavage sites (B) The *V. cholerae* genome encodes three endopeptidases (ShyA, ShyB, ShyC) possessing a hydrophobic region (gray), a PG binding domain (LysM, pink), and metallo-endopeptidase domain (M23, green). Protein domains were annotated using UniProt (1). (C) *lacZ* transcriptional reporters for each endopeptidase spotted onto LB (top row) and M9 minimal (bottom row) agar containing X-gal. A blue colony color indicates that the promoter is actively transcribed. Wild-type (WT) and $\Delta lacZ$ strain are included as positive and negative controls, respectively.

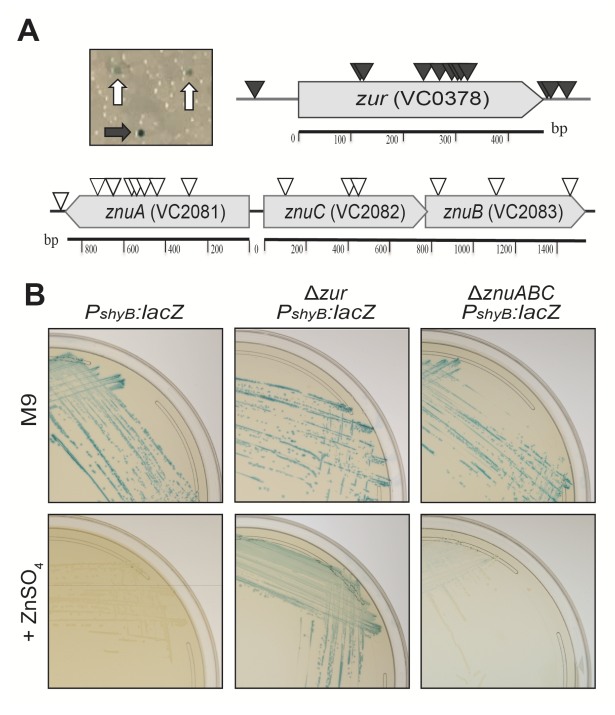


Fig 2. shyB transcription is regulated by zinc homeostasis.

(A) The *shyB* transcriptional reporter ($lacZ::P_{shyB}:lacZ$) was mutagenized with a Himar1 mariner transposon and screened for *shyB* induction (blue colonies) on LB agar containing X-gal and selective antibiotics (see Methods). Representative dark blue (black arrow) and light blue (white arrows) colonies are shown. Approximate Tn insertion sites identified by arbitrary PCR are shown (triangles). (B) The *shyB* transcriptional reporter in a wild-type, Δzur , or $\Delta znuABC$ background were grown on M9 X-gal agar without (top row) or with (bottom row) 10 μ M ZnSO₄.

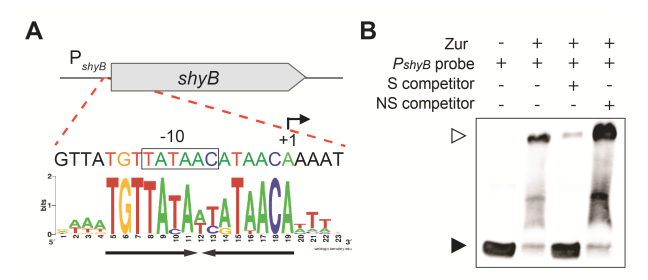


Fig 3. Zur directly binds the *shyB* promoter.

(A) The *shyB* promoter, annotated with a 5'-RACE transcription start site (+1) and putative -10 region (box), was aligned with a *Vibrio* Zur sequence logo (30, 31). The inverted AT-rich repeat in the putative Zur-box is underlined with black arrows. (B) A chemiluminescent probe containing the putative *shyB* Zur box was incubated with purified Zur in the presence of $ZnCl_2$ (5 μ M). Zur binding specificity was tested by adding 100-fold molar excess of unlabeled specific (S, Lane 3) or non-specific (NS, Lane 4) competitor DNA. Samples were electrophoresed on a 6% DNA retardation gel to separate unbound (black arrow) and bound probe (white arrow).

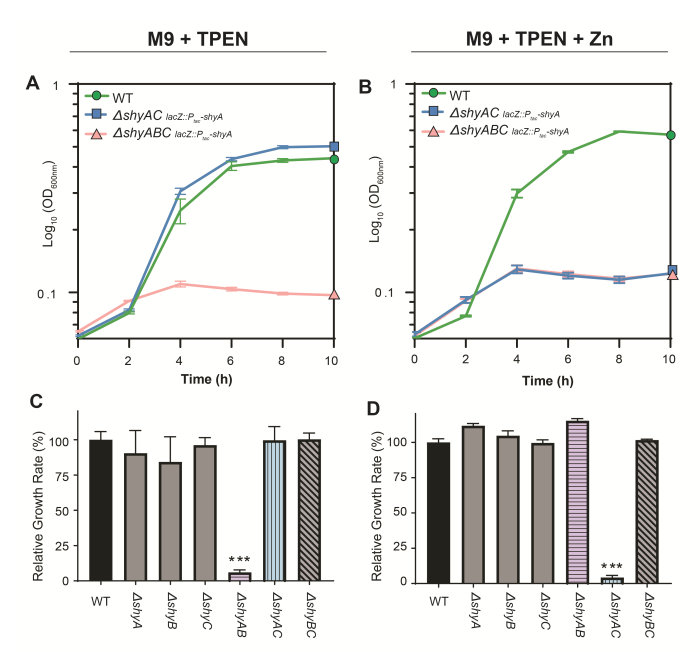


Fig 4. *shyB* supports cell growth in chelated medium and is conditionally essential in a $\triangle shyA$ mutant.

Mid-exponential cultures of the indicated *V. cholerae* mutants were washed to remove IPTG before being diluted 1:100 into M9 glucose containing streptomycin plus TPEN (250 nM) in the absence (A,C) or presence (B,D) of ZnSO₄ (1 μ M). Growth of each strain was monitored by optical density (OD600) in a Bioscreen C 100-well plate. Error bars report standard error of the mean (SEM) for three independent biological replicates. **(A-B)** Log-transformed growth curves are shown for WT (green circle), $\Delta shyAC$ *lacZ::P_{tac}:shyA* (blue square), and $\Delta shyABC$ *lacZ::P_{tac}:shyA* (red triangle). **(C-D)** In a similar growth experiment, growth rates of WT (solid black), single mutants (solid gray) and double mutants (striped) were calculated from exponential phase and normalized to the average WT growth rate (%). Statistical difference relative to WT was assessed using two-way analysis of variance (ANOVA) followed by Dunnett's multiple comparison test (***, p-value < 0.001).

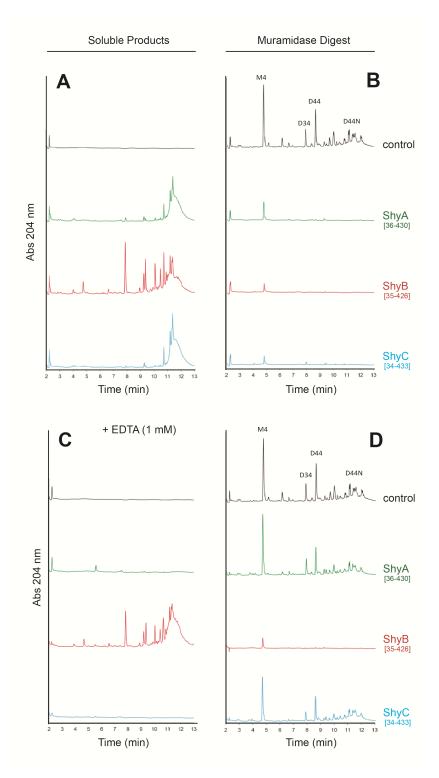


Fig 5. ShyB retains endopeptidase activity in EDTA.

V. cholerae sacculi was digested with 10 μ g of purified ShyA, ShyB, or ShyC for 16 h at 37°C in the absence (**A**) or presence (**C**) of 1mM EDTA. The soluble products released by digested sacculi were separated by size via HPLC and quantified by absorbance (204 nm). (**B**, **D**) The remaining insoluble pellet was digested with muramidase and soluble products were separated by HPLC.

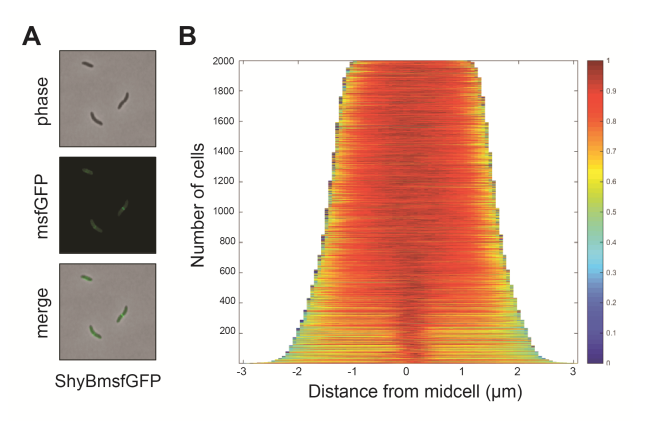


Fig 6. ShyBmsfGFP localizes to the midcell during division.

V. cholerae expressing a C-terminal fluorescent fusion ($lacZ::P_{tac}$ -shyBmsfGFP) was grown overnight in M9 + IPTG (200 μ M). (A) The ShyBmsfGFP fusion was imaged on an agarose patch (0.8% agarose in M9 minimal medium) with 300 ms exposure at 490 nm. (B) A heat map showing intensity of fluorescent signal as a function of distance from the midcell ("demograph") was generated from over 1,800 cells in Oufti (64).

_	терМ	znuA	znuC	znuB
Escherichia coli K12 MG1655 * Escherichia coli 0157:H7 Escherichia coli 0127:H6 Yersinia pestis CO92 * Salmonella typhimurium LT2 Enterobacter cloacae ATCC 13047 Klebsiella pneumoniae 342	Z2908 E2348C_1981	b1857 Z2909 E2348C_1982 YPO2061 STM1891 ECL_01441 KPK_1912	b1858 Z2910 E2348C_1983 YPO2060 STM1892 ECL_01440 KPK_1911	b1859 Z2911 E2348C_1984 YPO2059 STM1893 ECL_01439 KPK_1910

Fig 7. ShyB/MepM homologs are adjacent to the Zur-controlled *znu* operon in many Gramnegative pathogens.

Gene neighborhood alignments generated by Prokaryotic Sequence Homology Analysis Tool (PSAT) from 7 different Gram-negative bacteria (66). Arrows indicate the approximate location of the bidirectional promoter and site of Zur-binding in the *znu* operon. Asterisks indicate that co-transcription of *znu* and the downstream M23 endopeptidase is supported by transcriptomic data.