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Influence of core divisome proteins on cell division in Streptomyces venezuelae ATCC 10712

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24 ABSTRACT

The sporulating, filamentous, soil bacterium S. venezuelae ATCC 10712 differentiates under 25 submerged and surface growth conditions. In order to lay a solid foundation for the study of 26 development-associated division for this organism, a congenic set of mutants was isolated, 27individually deleted for a gene encoding either a cytoplasmic (i.e., *ftsZ*) or core inner membrane 28 (i.e., divIC, ftsL, ftsI, ftsO, ftsW) component of the divisome. While ftsZ mutants are completely 29 blocked for division, single mutants in the other core divisome genes resulted in partial, yet similar, 30 31 blocks in sporulation septum formation. Double and triple mutants for core divisome membrane components displayed phenotypes that were similar to the single mutants, demonstrating that the 32 33 phenotypes were not synergistic. Division in this organism is still partially functional without multiple core divisome proteins, suggesting that perhaps other unknown lineage-specific proteins 34 perform redundant functions. In addition, by isolating an *ftsZ2p* mutant with an altered -10 region, 35 the conserved developmentally controlled promoter was also shown to be required for sporulation-36 associated division. Finally, microscopic observation of FtsZ-YFP dynamics in the different 37 mutant backgrounds led to the conclusions that the initial assembly of regular Z rings does not per 38 se require the tested divisome membrane proteins, but that stability of Z rings is dependent on the 39 divisome membrane components tested. The observation is consistent with the interpretation that 40 Z ring instability likely results from and further contributes to the observed defects in sporulation 41 septation in mutants lacking core divisome proteins. 42

43 **INTRODUCTION**

Since the 1960s, *Streptomyces coelicolor* A3(2) has been used as the model system to study the 44 filamentous sporulating soil bacteria in the genus Streptomyces [1]. The streptomycetes are true 45 mycelial organisms of ecological importance, and are responsible for the synthesis of many 46 important biologically active compounds, including a wide range of antibiotics [2]. To facilitate 47 dispersal in the environment, and to enable long-term survival, streptomycetes produce semi-48 dormant spores. Spores are formed as part of a complex developmental life cycle, in which the 49 50 streptomycete initially grows vegetatively as branching hyphae while nutrients are available, forming a vegetative mycelium, also referred to as substrate mycelium. Eventually, specialized 51 52 spore-forming aerial hyphae emerge on the colony surface. The apical parts of such aerial hyphae are partitioned into chains of prespores via synchronous formation of several tens of cell division 53 54 septa in each hypha, and thereafter prespores mature to form pigmented and thick-walled spores with condensed nucleoids and a hydrophobic outer surface layer, before being released into the 55 56 environment (reviewed in e.g. [3-5]). When conditions are favorable, spores germinate and grow to form a new mycelium. The formation of aerial mycelium and spores is governed by a regulatory 57 58 cascade of transcription factors that has been studied mainly in S. coelicolor, but more recently much progress has been made in mapping the developmental regulatory networks in the new model 59 60 organism Streptomyces venezuelae [3-5].

Cell division and its regulation are of central importance in the Streptomyces life cycle, which 61 62 involves two distinct types of division. Both types are formed by a typical bacterial cell division 63 machinery, organized by the tubulin homolog FtsZ [6, 7]. In vegetatively growing hyphae, cell division is infrequent and gives rise to widely-spaced hyphal cross-walls. Intriguingly, vegetative 64 65 hyphae can grow even in the absence of such cross-walls and cell division is not essential for 66 proliferation of these organisms; the key cell division gene ftsZ is dispensable for growth and viability in S. coelicolor and S. venezuelae [8, 9]. The ftsZ-null mutants do not make cross-walls 67 at all, but can still grow as branching hyphae. On the other hand, cell division is absolutely essential 68 69 for spore formation. The septa that divide aerial hyphae into prespore compartments are formed by the same *ftsZ*-dependent core cell division machinery as the vegetative cross-walls. However, 70 71 sporulation septa differ from hyphal cross-walls. Structurally, sporulation septa are thicker,

eventually leading to full constriction and separation of daughter cells, and sporulation septatationis subject to spatial and temporal regulation [10].

Cell division genes are developmentally regulated and directly controlled by several of the key 74 transcriptional regulators that govern morphological differentiation and sporulation. For example, 75 76 ftsZ has a developmentally-regulated promoter, ftsZ2p, that is critical for spore formation [11]. 77 This promoter is repressed by the master regulator BldD in complex with cyclic di-GMP; 78 c-di-GMP-bound BldD negatively controls many important genes related to aerial mycelium 79 formation and sporulation [12, 13]. In S. venezuelae, both the ftsZ2p promoter and promoters for 80 cell division genes ftsW, ftsK, and sepF2 are activated by the proteins WhiA and WhiB, which 81 presumably act as a complex that controls a large regulon of genes involved in spore formation 82 [14, 15]. In addition, ssgB and, indirectly, ssgA are controlled by the developmental regulator 83 BldM [16]. SsgA and SsgB belong to an actinobacteria-specific protein family and affect selection of septation sites in S. coelicolor [17]. Finally, two dynamin-like proteins are involved in septation 84 85 specifically in sporulating aerial hyphae, and developmental upregulation of the corresponding 86 genes, dynA and dynB, depends on the sporulation-specific GntR-family regulator WhiH in S. 87 venezuelae [18].

88 The bacterial cell division process is primarily controlled at the level of assembly of FtsZ into cytokinetic polymers and their formation of a ring-shaped pattern, the Z ring, which serves to 89 recruit and organize most other proteins involved in cell constriction and septum formation [19-90 21]. Interestingly, streptomycetes, as well as Actinobacteria in general, lack obvious homologs of 91 92 most of the proteins that are known to regulate Z-ring formation, including MinC, MinD, SulA, Noc, or to stabilize FtsZ polymers or tether them to the cytoplasmic membrane, like FtsA and 93 ZipA [6, 7]. The exception is SepF, which links FtsZ to membranes and facilitates Z-ring formation 94 in Gram-positive bacteria like Bacillus subtilis [22], and which is present as three homologues in 95 96 S. venezuelae [18]. The regulation of Z-ring assembly in Streptomyces remains poorly understood. During sporulation, SsgA and SsgB proteins have been reported to mark the sites of Z-ring 97 98 formation in S. coelicolor [17], and DynA and DynB are recruited to septation sites and help stabilize Z rings in S. venezuelae [18]. DynA and DynB interact with each other, and DynB also 99 100 interacts with both SsgB and one of the three SepF proteins in S. venezuelae [18]. Analyses with two-hybrid systems have further suggested that the three SepF proteins interact with each other, 101

SepF interacts with FtsZ, and SepF2 interacts with both DynB and SsgB [18], indicating a
sophisticated protein interaction network affecting Z-ring assembly. Additional proteins have been
suggested to affect this critical step in cell division, including SepG and CrgA, but details still
remain unclear [23, 24].

106 Once formed, the Z ring recruits further division proteins that collectively are referred to as the 107 divisome. Present in streptomycetes are orthologues of conserved divisome proteins FtsQ, FtsL, 108 and DivIC(FtsB), which are known to form a complex with a structural and/or regulatory role in Escherichia coli and B. subtilis [19, 25, 26], and FtsW and FtsI, which encode a peptidoglycan 109 transglycosylase of the shape, elongation, division, and sporulation (SEDS) family, and a cognate 110 111 penicillin-binding protein with transpeptidase activity, respectively [27]. In addition, the DNAtranslocase FtsK and ABC-transporter proteins FtsEX are encoded by streptomycete genomes [7]. 112 113 Genetic studies in S. coelicolor show that none of these proteins are essential for growth or viability, which is consistent with the finding that cell division is dispensable in streptomycetes. 114 115 However, although mutants for ftsQ, ftsL, divIC, ftsW, and ftsI are largely defective in spore formation, none of these genes are absolutely needed for cell division, with all mutants being able 116 to form hyphal cross-walls and some sporulation septa [28-31]. For the last four genes, the mutant 117 118 phenotypes were found to be conditional, leading to suppression of the septation defect on minimal 119 medium or low osmolarity medium. Mutants lacking *ftsK* or *ftsEX* show apparently regular 120 sporulation septation, but *ftsK* mutants have a defect in chromosome stability, presumably related to the role of FtsK in clearing trapped chromosomes from the closing septa [7, 32-34]. 121

122 The fact that *Streptomyces* cell division is non-essential, developmentally regulated, disconnected from vegetative growth, and involves previously unknown mechanisms for control of septum 123 formation, make streptomycetes attractive model systems to study the division process and its 124 125 regulation [6, 7]. This distinction is further accentuated by the recent development of live cell 126 imaging systems in the new model organism S. venezuelae that allow time-lapse visualization of the cell division in great detail through the entire life cycle [3, 35]. In order to further establish S. 127 128 venezuelae as a cell division model system, we report the isolation and characterization of null mutants for key cell division genes ftsZ, ftsQ, ftsL, divIC, ftsW, and ftsI for this organism, and we 129 130 clarify the effect of late divisome components on assembly of Z rings and cell division in S. venezuelae. 131

132 METHODS

133 Bacterial strains and growth conditions

The S. venezuelae strains used in this study were derived from S. venezuelae strain ATCC 10712, 134 acquired from Dr. Colin Stuttard (Dalhousie University, Halifax, Canada) (Table S1). S. 135 136 venezuelae strains were cultivated at 30°C on maltose yeast extract medium (MYM) agar plates 137 or in MYM liquid medium [36], as described by Bush et al. [14]. S. venezuelae transconjugants were selected on either MYM or R2S agar after interspecies conjugation, as described previously 138 [37]. Culture conditions and antibiotics followed previously described procedures for 139 streptomycetes [38]. E. coli strain TG1 was used for cloning, construction, and propagation of 140 141 vectors [39]. E. coli strain BW25113/pIJ790 [40-43] was used to create cosmid derivatives containing insertion-deletion mutations. E. coli strain ET12567/pUZ8002 was used for 142 mobilization of oriT-containing cosmids and plasmids into S. venezuelae [37, 38]. E. coli strain 143 BT340 was used to express yeast Flp recombinase in E. coli to excise antibiotic resistance markers 144 flanked by FRT sites [44]. Culture conditions, antibiotic concentrations and genetic manipulations 145 146 generally followed those previously described for *E. coli* [39].

147

148 Plasmids and general DNA techniques

Plasmids and cosmids used in this study are listed in Table S2. DNA restriction and modifying 149 150 enzymes were used according to the manufacturer's recommendation (New England BioLabs). Phusion DNA polymerase (Thermo Fisher Scientific) was used according to the manufacturer's 151 instructions. S. venezuelae total DNA preparations were obtained using the Wizard genomic DNA 152 purification kit (Promega). λRED-mediated recombineering, modified for *Streptomyces*, was used 153 in E. coli to replace S. venezuelae genes on cosmids with mutagenic linear DNA cassettes [43]. 154 Apramycin-resistance gene cassette [oriT acc(3)IV] was amplified by PCR from plasmid pIJ773. 155 The oligonucleotide primers used in this study are listed in Table S3. When necessary, the *bla* gene 156 of the cosmid backbone was replaced by recombineering with a *bla* homology-flanked *oriT* 157 acc(3)IV-cassette from pIJ799. Plasmid pMS82 was used to create genetic complementation 158 plasmids for site-specific integration in the chromosome at the Φ BT1 attachment site [45]. DNA 159

sequences of unmarked in-frame deletions were verified using BigDye cycle sequencing analyzedon an ABI 3130 Prism Genetic Analyzer (Applied Biosystems).

162

163 Isolation of strains mutant for cell division genes

164 Using PCR product-directed recombineering of cosmid inserts [43], insertion-deletion mutations 165 were created in which all or crucial portions of the coding regions of S. venezuelae division genes 166 were replaced with an apramycin-resistance cassette (oriT acc(3)IV) from pIJ773 (Table S2). Care 167 was taken when designing the 3' endpoints of in-frame deletions to minimize the potential polar 168 effects on expression of the downstream co-transcribed gene(s). Mutagenized cosmids were 169 confirmed by restriction enzyme digestion and PCR amplification with primers flanking the 170 introduced mutations. These mutagenized cosmids were introduced into S. venezuelae by interspecies conjugation and marked null mutants, generated by double homologous 171 172 recombination events, were identified among primary transformants by their apramycin-resistant, kanamycin-sensitive phenotypes. Genomic DNA from mutant candidates was analyzed by PCR 173 174 amplification using primers flanking the mutations.

Most mutations were designed to introduce unique XbaI and SpeI sites flanking the oriT acc(3)IV175 176 cassette that was inserted into cosmids to generate the marked insertion-deletions. These restriction 177 sites facilitated the isolation of unmarked deletion mutant strains. The mutagenized cosmids were 178 digested with XbaI and SpeI and re-ligated, removing the oriT acc(3)IV cassette, leaving a 6 bp in-179 frame scar with the sequence of ACTAGA. Alternatively, the antibiotic-resistance cassette was 180 removed by site-specific recombination resulting in an 81-bp frt scar for unmarked ftsZ and ftsI 181 mutations. Subsequently, a linear oriT acc(3)IV cassette was used to replace the bla gene on the 182 cosmid backbone allowing conjugation into S. venezuelae and selection of exconjugants by 183 apramycin resistance marker in the vector backbone. Primary exconjugants generated by a single 184 homologous recombination incident were screened by PCR for gene conversion events in which 185 resident wild type alleles were replaced with the introduced unmarked mutagenized ones. 186 Generally, 5-10% of exconjugants had undergone gene conversion events. Exconjugants that were 187 homozygous for the mutant allele were re-streaked without selection to allow loss of integrated 188 cosmids and progeny colonies were screened for apramycin sensitivity, indicating intramolecular 189 homologous recombination events and the loss of the cosmid. All of the mutants chosen for further

characterization were checked by PCR amplification from genomic DNA with primers flanking 190 the introduced mutation to confirm the presence of only the unmarked deletion allele. Double 191 192 mutants were constructed by introducing marked insertion-deletion mutations into unmarked single mutants, as described above for isolating single mutants. A triple mutant for *ftsL*, *ftsQ* and 193 divIC was isolated in a similar fashion from an unmarked double mutant strain. A double mutant 194 strain for the adjacent *ftsL* and *ftsI* genes was obtained by combining recombineering primers used 195 for single mutation isolation (i.e., using the 5' ftsL primer and the 3' ftsI primer). The resulting 196 197 $\Delta ftsIL::apra$ mutation was introduced into the chromosome in the same way as for single mutant isolation. 198

For generation of a non-sporulating strain by manipulating the developmentally controlled *ftsZ2p*, the TAGTGT residues of the -10 motif on cosmid Sv-4-G01 were replaced with an *oriT acc(3)IV* cassette flanked by introduced *Spe*I and *Xba*I sites. Restriction digestion of the mutagenized cosmid with *Spe*I and *Xba*I and re-ligation left ACTAGA in place of the native -10 sequence. Exconjugants were selected as described above for unmarked mutations and identified by a PCR analysis using oligonucleotides specific for each of the two promoter sequences.

205

206 Construction of genetic complementation plasmids

For genetic complementation, a series of DNA fragment inserts were generated from cosmids by restriction digestion or amplification by PCR (Fig. 1a and Fig. S8) and cloned into site-specific integration vectors pMS82. The resulting plasmids were introduced into *S. venezuelae* mutant strains by conjugation and integrated *in trans* into the chromosome at the Φ BT1 attachment site.

211

212 Construction of strains expressing fluorescent FtsZ-YPet fusion proteins

Plasmid pKF351, carrying and *ftsZ-ypet* fusion in a vector that integrates at the Φ C31 attachment site [46], was introduced into relevant mutants by interspecies conjugation, as described above.

215

216 Microscopy

For phase-contrast microscopy, bacteria were grown as confluent patches on MYM agar. Cover
slips were touched to the surface of sporulated patches and material lifted was mounted on pads of
1% agarose in PBS. Samples were visualized using a Nikon Eclipse E400 with a Nikon 100x 1.25
NA oil immersion objective and a MicroPublisher 5.0 RTV high resolution CCD camera
(Qimaging).

For staining of cell wall and nucleoids, cultures were grown on MYM agar or in MYM liquid medium, and samples were fixed in ice-cold methanol for 5 minutes, washed twice in PBS and mounted in 100 µg ml⁻¹ propidium iodide (Molecular Probes) and 10 µg ml⁻¹ WGA-FITC (Molecular Probes) in 50% glycerol. Fixed and stained samples were then spotted onto pads of 2% agarose in PBS and sealed with petroleum jelly. Fluorescence imaging was done with a Leica SP2 TCS confocal microscope using a Leica 63x 1.4 NA glycerin immersion objective.

228 In order to visualize fluorescent FtsZ-YPet fusion protein, cells were grown in liquid MYM, 229 harvested and fixed with 2.28% formaldehyde and 0.018% glutaraldehyde, washed in PBS, and 230 mounted on 1% agarose in PBS. To follow FtsZ dynamics, microfluidics-based time-lapse microscopy was performed using the CellASIC ONIX system and B04A-03 microfluidic plates 231 232 (Merck Millipore), as described previously [35, 47]. The live-cell time lapse experiments were repeated twice for each strain. Imaging was performed on a Zeiss AxioObserver.Z1 microscope 233 with Zeiss Plan-Apochromat 100×/1.4 Oil Ph3 objective, ZEN software (Zeiss) and an ORCA 234 Flash 4.0 LT camera (Hamamatsu). Images and movies were processed using ImageJ/Fiji [48]. 235

For Transmission Electron Microscopy (TEM), cells were grown as lawns on MYM agar and fixed in 2.5% glutaraldehyde in 0.05 M cacodylate buffer (pH 7.2) and incubated for 1 h in 2% osmium tetroxide. Cells were dehydrated by successive transfer in 5 steps from 50% to 100% ethanol. Cells were washed in propylene oxide and then in a 1:1 solution of propylene oxide and Spurr's before being incubated in Spurr's resin overnight at 60°C. Thin sections were stained with 2% uranyl acetate and 1% lead citrate. Samples were visualized using a JEOL JEM-1210 equipped with a Hamamatsu Orca-HR CCD camera.

243 **RESULTS AND DISCUSSION**

244 Generation of *ftsZ*-null mutants

The earliest acting divisome protein FtsZ is required for cell division and viability in most bacteria. 245 246 Previously, it had been demonstrated that *ftsZ*-null strains could be isolated for *S. coelicolor* [8]. However, it was not clear whether this would be a general property of streptomycetes. Therefore, 247 248 the procedure for isolating a deletion-insertion mutation for S. coelicolor ftsZ was replicated in S. venezuelae by replacing 844 bp, starting 16 bp upstream of the ftsZ start codon, with an 249 250 apramycin-resistance cassette (Fig. 1a). The mutants described in this paper were isolated in S. venezuelae ATCC 10712 acquired from Dr. Colin Stuttard (Dalhousie University, Halifax, 251 252 Canada). In three independent experiments, we were able to isolate S. venezuelae null mutants for ftsZ. However, in contrast to the majority of the other cell division mutants isolated in 253 S. venezuelae (described below), obtaining the *ftsZ*-null mutants was not as straightforward. The 254 initial *ftsZ*-null mutant colonies grew very slowly after selection on conjugation plates, therefore, 255 256 waiting up to 8 days before picking colonies was necessary to identify these mutants as "specks" or "flecks" on the agar surface. These primary mutant colonies on conjugation plates did not 257 258 increase in size upon prolonged incubation, nor did they sector. Subsequently, mutants were re-259 streaked several times on selective media. *ftsZ*-null mutants were unhealthy and grew slowly on 260 the nalidixic acid-containing medium used for counter-selection of the conjugation donor E. coli, 261 perhaps contributing to the difficulty when isolating them initially and, similarly, when introducing 262 an empty vector as control for genetic complementation studies (see below). Once nalidixic acid 263 counter-selection was no longer needed, the single colonies were uniform in appearance and the 264 phenotype was stable. Colonies of the purified *ftsZ*-null strains appeared on plates at similar 265 incubation times relative to the wild type, however, plating efficiency was much lower (Figs. S1 266 and S2); mature colonies were smaller and took much longer to develop an aerial mycelium, as 267 judged by the surface of the colonies becoming white. Isolated independent strains had similar 268 microscopic phenotypes and the insertion-deletion mutation of *ftsZ* was confirmed by PCR from 269 genomic DNA. One representative strain was picked for further analysis (DU500). As expected, western blot analysis verified that FtsZ was not detected in a whole cell extract from DU500 (Fig. 270 271S3). As anticipated, phase-contrast and TEM microscopic analyses of the *ftsZ*-null mutant grown on agar showed that aerial hyphae did not differentiate to produce spores and the vegetative hyphae 272

were devoid of the normal cross-walls (Figs. 1b and c). The *ftsZ*-null mutant was also completely 273274 blocked in both cell division and spore production when grown in liquid cultures and little cell material accumulated for the mutant under these growth conditions (data not shown). To confirm 275276 that the observed division phenotype was the result of the introduced mutation, genetic complementation studies were carried out. A restriction fragment containing ftsQ, ftsZ and the 277 278 native ftsZ promoters in the intergenic region between the genes was integrated at the chromosomal att PBT1 site (pJS8; Fig. 1a). The complementation vector rescued the division phenotype of the 279 280 ftsZ-null mutants, as judged by phase-contrast microscopy (Fig. 1b), as well as restored growth 281 and colony size on agar medium (Figs. S2 and S3). We conclude that the deletion-insertion 282 mutation is not polar on downstream gene expression, and FtsZ-dependent cell division is 283 dispensable for growth and viability of S. venezuelae. Nonetheless, an unmarked ftsZ-null strain 284 was also generated and it had an identical microscopic phenotype as the insertion-deletion mutant, 285 but was not used further in this study (DU665, data not shown). The fact that an unmarked null 286 mutant can be isolated by gene conversion (see Methods) argues that the *ftsZ* mutants are difficult 287 to distinguish from background on primary conjugation plates, not that they can only be isolated 288 by very strong selection for marker replacement by double homologous recombination.

289 Using the same procedure that is described above, we have also isolated an *ftsZ*-null mutant 290 (DU669) in the S. venezuelae strain NRRL B-65442 obtained from Dr. Mark Buttner (John Innes 291 Centre, Norwich, UK) [49]. The mode of growth of this *ftsZ*-null mutant DU699 (NRRL B-65442 292 background) in the absence of cell division and hyphal cross-walls has been described elsewhere 293 [9]. As observed for other cell division mutants (data not shown), the macroscopic and microscopic phenotypes of *ftsZ*-null mutants in the two independent wild type backgrounds were essentially 294 indistinguishable and could be genetically complemented, revealing no overt differences at the 295 296 phenotypic level between the S. venezuelae parent strains obtained from different sources. All 297 experiments described in the rest of this paper were carried out in the ATCC 10712 strain background. (Nonetheless, about a dozen core divisome mutant strains were also isolated in the 298 299 NRRL B-65442 background. Their strain designations and genotypes are listed in Table S1.)

300

The developmentally regulated promoter of *ftsZ* is required for sporulation-associated cell division in *S. venezuelae*

For S. coelicolor, three promoters for ftsZ have been mapped to the 288 bp intergenic region 303 304 between ftsZ and ftsQ, and one of them, ftsZ2p, is developmentally regulated [11]. It has been shown that BldD, a transcriptional regulator that plays a key role in Streptomyces development, 305 306 binds to the developmentally-regulated ftsZ2p promoter and acts to repress expression of ftsZduring vegetative growth [12]. In S. venezuelae it has been shown that expression of ftsZ is 307 308 dependent on WhiA and WhiB, which are transcriptional regulators required for the transition from growth of aerial hyphae to sporulation [14, 15]. ChIP experiments indicated that a WhiA binding 309 310 target lies around 158 bp from the predicted start codon of ftsZ [14], and in this region is a sequence that is identical to the -10 region TAGTGT of the S. coelicolor ftsZ2p [11] and S. griseus P_{spo} [50]. 311 The intergenic region upstream of *ftsZ* is highly conserved between *S. coelicolor* and *S. venezuelae* 312 with sequence conservation at the three mapped promoter regions, including the ftsZ2p promoter 313 314 (Fig. S4a). To test whether the importance of this developmental promoter for sporulation was also retained in S. venezuelae, a strain was generated that was mutant for this presumed ftsZ2p promoter 315 316 region. In the unmarked mutant, the TAGTGT residues at the -10 region of this presumed promoter were changed to ACTAGA (Fig. 4b). The resulting strain (DU523) had reduced plating efficiency 317 318 compared to the wild type, but grew robustly and formed an abundant aerial mycelium (Fig. S1). 319 However, the *ftsZ2p* mutant was unable to efficiently convert the aerial hyphae to spore chains 320 during growth on solid medium, and mature spores were observed much less frequently compared 321 to the wild type. Instead, longer spore-like compartments of irregular length were produced from 322 aerial hyphae (Fig. 1b). Thus, this promoter mutant can form functional division septa that result in complete division events with cell separation that lead to formation of spore-like aerial hyphal 323 fragments. However as shown by the absence of regularly septated spore chains, the mutant has a 324 greatly reduced frequency of cell division compared to wild type. The phenotype of the 325 326 S. venezuelae ftsZ2p mutant is consistent with a failure to up-regulate the expression of ftsZ, and 327 this up-regulation is required for developmentally-associated cell division, similarly to what has been observed in S. coelicolor and S. griseus [11, 50]. 328

329

330 Null mutants for core divisome genes

In order to clarify the roles of some of the conserved core divisome proteins for *S. venezuelae*,
unmarked in-frame null mutants for *ftsQ* (strain DU629), *divIC* (strain DU613), *ftsL* (strain

DU520), ftsW (strain DU521) and ftsI (strain DU679) were isolated (Fig. 1a). These genes are 333 broadly conserved among bacteria and their products are membrane proteins required for 334 coordinating the cytoplasmic Z ring with the peptidoglycan synthesis machinery [19, 25]. In other 335 336 bacteria, FtsQ, FtsL and DivIC form a subcomplex that is recruited to the divisome. A 1 MDa 337 complex containing those proteins, along with FtsZ, has been identified for E. coli [51]. Likewise, 338 FtsI and FtsW form a subcomplex involved in septal peptidoglycan synthesis as a transpeptidase and transglycosylase, respectively [27]. While it has not been demonstrated directly for 339 340 *Streptomyces*, it is reasonable to expect that the protein subcomplexes are conserved.

Since most of the core divisome genes are part of the complex *dcw* gene cluster (Fig. 1a) and are 341 342 likely co-transcribed with other cell wall biosynthetic genes, unmarked in-frame deletions were generated to avoid polar effects on downstream genes. The mutants were readily isolated and 343 344 showed consistent macroscopic and microscopic phenotypes that were strikingly similar to one another on MYM agar, but were distinct from the *ftsZ*-null and *ftsZ2p* mutants. In contrast to the 345 346 ftsZ-null mutant, plating efficiency was unaffected in these mutants (Fig. S1), suggesting that vegetative cross-wall formation was not severely impaired. Likewise, aerial mycelium 347 348 development was essentially unaffected, however, aerial hyphae were not efficiently converted into chains of spores. By 48 hrs of incubation on solid medium, the majority of aerial mycelium 349 350 had been converted into spores for the wild type. For the ftsQ, ftsL, divIC, ftsW and ftsI null mutants, a mixture of spores, hyphal fragments and frequent lysed compartments were observed 351 (Figs. 1b and S5). Aerial hyphae often contained frequent and regularly-spaced constrictions, 352 reminiscent of sporulation septa (Fig. 1b). Some separated spores were produced and fragments of 353 varying lengths were also observed for each of these mutants, showing that the products of these 354 genes are not absolutely required for cell division for S. venezuelae (Figs. 1b and S5). Each 355 356 mutation was genetically complemented using site-specific integration plasmids with inserts shown in Figure 1a. The complemented strains sporulate similar to the wild type parent indicating 357 that the phenotypes were associated with the introduced null mutations (Figs. 1b and S1). 358 359 Combining *ftsQ*, *divIC* and *ftsL* mutations as either double mutants or as a triple mutant did not 360 have a synthetic effect on the observed phenotypes, as judged by phase-contrast microscopy (Fig. 361 2), suggesting that the loss of all the parts of the putative subcomplex formed by their gene products 362 is no more deleterious than the loss of any one component. This result is consistent with the 363 interpretation that missing any one component must inactivate the remainder of the tripartite

364 complex. In addition, a double mutant lacking both ftsW and ftsI was also constructed and the 365 phenotype was indistinguishable from the individual *ftsW* and *ftsI* mutants (Fig. 2), suggesting that 366 the loss of both parts of the putative subcomplex formed by their gene products is no more 367 deleterious than the loss of either one component. Finally, deleting both adjacent *ftsL* and *ftsI* genes 368 together resulted in a mutant with a similar phenotype to an *ftsI* single mutant (Fig. 2), indicating 369 that removal of parts of both putative divisome subcomplexes is no more deleterious than the loss of one part. The similarity of the core mutant phenotypes and lack of synergism when combining 370 371 divisome mutations seems to support a model where there is no apparent hierarchy of assembly of the core divisome components in S. venezuelae. Further experimentation will be needed to clarify 372 373 the situation and define each contribution.

Spores could be isolated from the aerial mycelium of surface-grown cultures despite the fact that 374 375 development-associated division was impaired for core divisome single mutants with a reduction 376 in the number of spores produced relative to the wild type. In order to quantify the severity of 377 reduction in sporulation-associated division in these core divisome mutants, measurements were 378 made of mature spores and hyphal fragments harvested in a typical fashion from agar plates after 379 4 days incubation. For the wild type, the material harvested consisted almost entirely of spores 380 with an average length of 1.00 (± 0.23) μ m (Fig. S6). In contrast, the average lengths of the spores 381 and hyphal fragments for the ftsQ, divIC and ftsL mutants were similar at 2.13 (±1.76) µm, 2.75 382 (± 2.67) µm and 2.49 (± 3.92) µm, respectively, suggesting that in aerial hyphae when development-383 associated division resulting in cell separation occurred, every other to every third septum was 384 formed with cell separation in some aerial hyphae. In contrast, the average spore-type compartment 385 lengths were greater for the ftsW and ftsI mutants, 7.18 (±8.32) µm and 5.73 (±6.00) µm, 386 respectively (Fig. S6), suggesting that in some hyphae with development-associated division 387 leading to cell separation, every sixth to seventh septum may have been completed all the way to 388 detachment of cells.

Phase-contrast microscopy showed that the divisome mutants were capable of division leading to cell separation, but did not provide detail on the septum morphology when division failed. Of the five isolated single mutants, *ftsL* and *ftsI* single mutants were selected for observation by electron microscopy as representative examples of mutants affecting the putative FtsQ-DivIC-FtsL complex and the FtsW-FtsI complex. TEM analysis for an *ftsL*-null mutant indicated that the evenly-spaced constrictions in aerial hyphae observed by light microscopy represented complete
invaginations with very thick peptidoglycan (Fig. 1c), while *ftsI*-null mutants produced more
normal looking septa (Fig. 1c), but at a lower frequency.

While extremely rare in the wild type strain, branching within nascent spore chains was commonly observed for the *ftsQ*, *divIC*, *ftsL*, *ftsW* and *ftsI* mutants. Both the cell division defect and the observed branching phenotype in aerial hyphae were rescued in genetic complementation studies, confirming that the phenotypes are associated with the deletion of these genes and not the result of unlinked mutations (Fig. 1b).

402

403 Core divisome proteins are not absolutely required for genome segregation in *S. venezuelae*

To further characterize the cell division and sporulation defects in *ftsZ*, *ftsZ2p*, *ftsQ*, *ftsL*, *divIC*, *ftsW* and *ftsI* mutants of *S. venezuelae*, the cell wall was stained using WGA-FITC and nucleoids were stained by propidium iodide (Figs. 3 and S7). Cell wall staining confirmed that hyphae of the *ftsZ*-null mutant had no signs of invagination, vegetative cross walls or sporulation septa. In addition, there was no evidence of DNA condensation either (Fig. 3). In contrast, partial nucleoid condensation and segregation was observed for the *ftsZ2p* developmental promoter mutant (Fig. 3).

Consistent with observations from phase-contrast and TEM microscopy, ftsQ, ftsL, divIC, ftsW 411 and *ftsI* mutants showed very similar patterns of cell wall and DNA staining (Figs. 3 and S7). In 412 413 aerial hyphae with constrictions visible by light microscopy, ladders of nascent septal wall material could be seen. However, these ladders were often not as regular as the evenly-spaced ones seen 414 415 for the wild type. DNA segregation was not grossly affected by the loss of any of these division 416 genes, but often was less uniform than for the wild type. Overall, under the laboratory conditions that we tested, we conclude that S. venezuelae is able to lay down cell division septa and segregate 417 418 their genomes even in the absence of the core divisome genes ftsQ, ftsL, divIC, ftsW or ftsI. Future 419 avenues of research will be necessary to understand why these genes are conserved, yet their products are not essential for septum formation for this filamentous bacterium. 420

422 Assembly of FtsZ into ladder-like arrays of Z rings in sporogenic hyphae does not require 423 the core divisome genes *ftsQ*, *ftsL*, *divIC*, and *ftsW*

Next, using a subset of the mutants, we investigated to what extent the core divisome mutations 424 affected the localization and dynamics of FtsZ rings in S. venezuelae. In order to do this, ftsZ-ypet 425 426 (pKF351) was introduced into the Φ BT1 *att* site of *ftsQ*, *ftsL*, *divIC*, and *ftsW* mutants, as well as 427 the wild type strain, leading to production of FtsZ fused to the yellow fluorescent protein YPet in 428 addition to the native FtsZ. In vegetative hyphae sampled from standard liquid medium cultures at different stages along the growth curve, we observed an apparently normal distribution of Z rings 429 in vegetative hyphae (not shown, but see also microfluidics data below). Further, sporulating 430 431 hyphae with multiple, closely and regularly-spaced Z rings were observed in both the wild type strain and the *ftsQ*, *ftsL*, *divIC*, and *ftsW* mutants (Fig. 4), albeit examples of sporulating hyphae 432 433 were observed at a lower frequency in the mutants than in the wild type.

In order to observe more clearly how mutations in core divisome genes affect FtsZ dynamics and 434 Z-ring formation, we used microfluidics-based fluorescence live cell imaging, as described 435 previously [35, 47]. Representative micrographs of FtsZ ladders formed in the wild type and *ftsQ*. 436 437 and *ftsW* mutants under these conditions are shown in Fig. 5. Time lapse images were also acquired 438 to visualize the FtsZ dynamics. In the wild type, during vegetative stage, typical Z rings are observed that are highly dynamic as shown by their movement along the hyphae before they 439 stabilize at fixed positions and then increase in fluorescence intensity (Movie S1). Presumably, 440 441 these observed intense Z rings mark sites of vegetative cross-wall formation. A very similar pattern 442 was seen for formation of Z rings in early growth timepoints for vegetative hyphae of the *ftsQ* and ftsW mutants (Movies S2 and S3, respectively). FtsZ dynamics were also visualized during the 443 sporulation stage for the wild type parent, where examples of the assembly of evenly-spaced Z 444 rings in ladder-like patterns could be seen in sporogenic hyphae (Movie S1, >10 hours of growth 445 446 in this sample). Intriguingly, similar development-associated FtsZ dynamics were observed in the mutants for ftsQ (Movie S2) and ftsW (Movie S3) and the assembly of evenly-spaced FtsZ ladders 447 448 occurred. Closer inspection of the stability FtsZ ladder persistence was accomplished by constructing a montage from timepoint images for the wild type parent and the ftsQ and ftsW449 450 mutants (Fig. 6). The FtsZ ladders persist for approximately two hours for the wild type, but the ladders do not show the same dynamics for the mutants. Certain FtsZ rings are lost over time in 451

the mutants, with the rungs of FtsZ ladders in the ftsW mutant being the least stable. These relative FtsZ ladder stabilities correlate with the average lengths of mature spores that can be harvested from surface grown cultures (Fig. S6), with the spores for ftsQ and ftsW mutants being approximately 2X and 7X the length of those produced by the wild type.

456

457 Physiological relevance of cumulative results

458 Overall, the results clearly show that the products of the divisome genes *ftsQ*, *ftsL*, *divIC*, and *ftsW* are not required for Z-ring assembly, for the single Z rings that are formed in vegetative hyphae 459 (normally leading to hyphal cross-walls in the wild type), and in sporogenic hyphae, where ladders 460 461 of regularly spaced Z rings are typically formed as part of sporulation septation. In some 462 instances, Z-ring formation appeared essentially normal in the *ftsQ*, *ftsL*, *divIC*, and *ftsW* mutants. 463 The results are consistent with the previously observed ability of corresponding mutants in S. 464 coelicolor to form cross-walls and septa, at least under certain conditions [28-31]. It has been 465 speculated previously that *ftsW* may be required for Z-ring assembly, and may provide a membrane 466 attachment for FtsZ in both S. coelicolor and Mycobacterium smegmatis [31, 52]. Our results 467 presented here for S. venezuelae show that ftsW is not required for Z-ring formation.

468 The fact that core cell division proteins FtsQ, FtsL, DivIC, FtsW and FtsI are not strictly required 469 for cell division in *Streptomyces* spp. gives rise to interesting questions to be investigated in future 470 studies. For example, how is it possible to carry out cell division in the absence of the FtsQ-FtsL-471 DivIC complex? Either the divisome in *Streptomyces* spp. can be stable and functional without these proteins, or there are other proteins that can replace or reinforce these core divisome proteins. 472 In the latter case, such proteins would be pertinent to identify. Interestingly, co-473 474 immunoprecipitation experiments identified 63 FtsQ-interacting proteins for Mycobacterium tuberculosis and may point to homologs for further investigation [53]. 475

Similarly, how can cell division occur in the absence of FtsW and its cognate transpeptidase FtsI?
The transpeptidase of FtsI has been shown to be nonessential for some other gram-positive
bacteria, although the protein is still physically required [54-56]. In the absence of FtsI, perhaps
FtsW functions with one or more of the many PBPs encoded for *S. venezuelae*. FtsI transpeptidase
activity can be supplied by other PBPs in *B. subtilis* [56]. FtsW co-purifies with two different PBPs

481 in a potential trimeric complex in E. coli [57]. FtsW has recently been identified as a peptidoglycan 482 transglycosylase (essentially a peptidoglycan polymerase) [27], in similarity to related RodA 483 SEDS proteins [58-60]. These are essential activities for formation of a cell division septum, and 484 the results presented here suggests that another peptidoglycan polymerase likely is recruited to Z rings at division sites in order for the S. venezuelae ftsW mutant to form septa. As one possibility, 485 486 perhaps transglycosylase activity can be provided by an autonomous bifunctional class A PBP and 487 not by a SEDS protein. In support of that notion, evidence for intimate participation of bifunctional 488 PBPs in septum peptidoglycan synthesis has been accumulating [61]. Recent evidence suggests 489 that pneumococcal peptidoglycan is synthesized, in part, by bifunctional PBPs [62]. As another possibility, one of the other three SEDS proteins encoded by streptomycete genomes [29, 31] may 490 491 be active either at the same time as FtsW and/or induced in the absence. It will be interesting to 492 see which protein(s) functions in a *ftsW* mutant and how it would be recruited to the divisome. Of final note, we also have constructed strains individually expressing EFGP fusions to each protein 493 494 of the FtsQ-FtsL-DivIC or FtsI-FtsW complexes and the fluorescent localization signals are not strong enough to publish (data not shown). Potentially, the weak fluorescence signal is indicative 495 496 of a low intracellular concentration. Again, future work will have to be done to learn if small 497 amounts of the proteins are needed for normal function, if another protein can substitute, or if 498 multiple SEDS-PBP pairs work simultaneously during sporulation septum formation.

499 SUMMARY AND CONCLUSIONS

500 In this study, we have established the contributions of known central cell division proteins in the 501 coordinated process of sporulation septation in *S. venezuelae*. Knowing the null phenotypes for 502 mutants lacking known players in cell division will be essential for future studies as we continue 503 to peel back the novel lineage-specific layers of controls evolved to govern the concerted 504 development-associated control of essential cell biological processes in streptomycetes.

505 In this study, we have taken advantage of the benefits of *S. venezuelae* to visualize the synchronous 506 events being orchestrated within sporogenic hyphae by live-cell time-lapse microscopy because 507 this species undergoes differentiation under submerged growth conditions. The data show that 508 ladder-like assemblages of evenly-spaced FtsZ rings typically form in all of the characterized core 509 divisome mutants. Thus, the tested divisome components are not required for that early 510 coordinated event. However, once formed the Z-rings appear to be unstable and a number of rings prematurely disband. The loss of coordination results in irregular spacing between completed septaand irregular spore size, as seen in the divisome mutants.

Evidence has accumulated for subcomplex formation of FtsQ-FtsL-DivIC and FtsW-FtsI before 513 participation in the divisome. For S. venezuelae, combining mutations of genes encoding these 514 components do not result in synthetic phenotypes. The result is consistent with the interpretation 515 516 that the loss of any one component disrupts the function of the subcomplex. While the subcomplexes are not absolutely required, they do contribute to the stability of the synchronous 517 518 tandem arrays of divisome complexes as visualized by FtsZ-YPet. Recently, analysis of bacterial dynamins DynA and DynB for S. venezuelae showed that they interact with the division machinery 519 520 [18], contribute to Z-ring stability and mutants encoding those proteins have somewhat similar phenotypes as the divisome mutants reported here. Future work will be needed to understand how 521 522 these components interact and are regulated to synchronously coordinate sporulation septum formation. 523

524

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536

537 **Conflicts of interest**

538 The authors declare no conflict of interests.

539 **REFERENCES**

Hopwood DA. Forty years of genetics with *Streptomyces*: from *in vivo* through 540 1. in vitro to in silico. Microbiology. 1999;145:2183-202. Doi: 10.1099/00221287-145-9-541 2183 542 Barka EA, Vatsa P, Sanchez L, Gaveau-Vaillant N, Jacquard C, Meier-2. 543 Kolthoff JP, et al. Taxonomy, physiology, and natural products of Actinobacteria. 544 Microbiol Mol Biol Rev. 2016;80:1-43. Doi: 10.1128/MMBR.00019-15 545 Bush MJ, Tschowri N, Schlimpert S, Flärdh K, Buttner MJ. c-di-GMP 546 3. signalling and the regulation of developmental transitions in streptomycetes. *Nat Rev* 547 Microbiol. 2015;13:749-60. Doi: 10.1038/nrmicro3546 548 McCormick JR, Flärdh K. Signals and regulators that govern Streptomyces 549 4. development. FEMS Microbiol Rev. 2012;36:206-31. Doi: 550 Elliot MA, Flärdh K. Streptomycete spores. eLS. Chichester: John Wiley & 551 5. Sons Ltd; 2020. Doi: 10.1002/9780470015902.a0000308.pub3 552 6. **Jakimowicz D, van Wezel GP.** Cell division and DNA segregation in 553 Streptomyces: how to build a septum in the middle of nowhere? Mol Microbiol. 554 2012;85:393-404. Doi: 10.1111/j.1365-2958.2012.08107.x 555 **McCormick JR.** Cell division is dispensable but not irrelevant in *Streptomyces*. 556 7. *Curr Opin Microbiol.* 2009;12:689-98. Doi: 557 558 8. McCormick JR, Su EP, Driks A, Losick R. Growth and viability of Streptomyces coelicolor mutant for the cell division gene ftsZ. Mol Microbiol. 559 560 1994;14:243-54. Doi: 561 Santos-Beneit F, Roberts DM, Cantlay S, McCormick JR, Errington J. 9. A mechanism for FtsZ-independent proliferation in Streptomyces. Nat Commun. 562 2017;8:1378. Doi: 10.1038/s41467-017-01596-z 563 Wildermuth H, Hopwood DA. Septation during sporulation in *Streptomyces* 564 10. 565 coelicolor. J Gen Microbiol. 1970;60:57-9. Doi: Flärdh K, Leibovitz E, Buttner MJ, Chater KF. Generation of a non-566 11. sporulating strain of *Streptomuces coelicolor* A₃(2) by the manipulation of a 567 developmentally controlled *ftsZ* promoter. *Mol Microbiol*. 2000;38:737-49. Doi: 568 569 den Hengst CD, Tran NT, Bibb MJ, Chandra G, Leskiw BK, Buttner 12. MJ. Genes essential for morphological development and antibiotic production in 570 Streptomyces coelicolor are targets of BldD during vegetative growth. Mol Microbiol. 571 572 2010;78:361-79. Doi: 10.1111/j.1365-2958.2010.07338.x Tschowri N, Schumacher MA, Schlimpert S, Chinnam NB, Findlay KC, 573 13. Brennan RG, et al. Tetrameric c-di-GMP mediates effective transcription factor 574 dimerization to control Streptomyces development. Cell. 2014;158:1136-47. Doi: 575 10.1016/j.cell.2014.07.022 576 Bush MJ, Bibb MJ, Chandra G, Findlay KC, Buttner MJ. Genes required 14. 577 for aerial growth, cell division, and chromosome segregation are targets of WhiA before 578 sporulation in *Streptomyces venezuelae*. *MBio*. 2013;4:e00684-13. Doi: 579 580 10.1128/mBio.00684-13 Bush MJ, Chandra G, Bibb MJ, Findlay KC, Buttner MJ. Genome-wide 581 15. chromatin immunoprecipitation sequencing analysis shows that WhiB is a transcription 582 583 factor that cocontrols its regulon with WhiA to initiate developmental cell division in Streptomyces. MBio. 2016;7:e00523-16. Doi: 10.1128/mBio.00523-16 584

585 16. Al-Bassam MM, Bibb MJ, Bush MJ, Chandra G, Buttner MJ. Response
586 regulator heterodimer formation controls a key stage in *Streptomyces* development.
587 *PLoS Genet.* 2014;10:e1004554. Doi: 10.1371/journal.pgen.1004554

588 17. Willemse J, Borst JW, de Waal E, Bisseling T, van Wezel GP. Positive 589 control of cell division: FtsZ is recruited by SsgB during sporulation of *Streptomyces*. 590 *Genes Dev*. 2011;25:89-99. Doi:

591 18. Schlimpert S, Wasserstrom S, Chandra G, Bibb MJ, Findlay KC,

Flärdh K, et al. Two dynamin-like proteins stabilize FtsZ rings during *Streptomyces* sporulation. *Proc Natl Acad Sci USA*. 2017;114:E6176-E83. Doi:

594 10.1073/pnas.1704612114

595 19. **Du S, Lutkenhaus J.** Assembly and activation of the *Escherichia coli* divisome. 596 *Mol Microbiol.* 2017;105:177-87. Doi: 10.1111/mmi.13696

597 20. **Haeusser DP**, **Margolin W**. Splitsville: structural and functional insights into 598 the dynamic bacterial Z ring. *Nat Rev Microbiol*. 2016;14:305-19. Doi:

599 10.1038/nrmicro.2016.26

600 21. **Ortiz C, Natale P, Cueto L, Vicente M.** The keepers of the ring: regulators of 601 FtsZ assembly. *FEMS Microbiol Rev.* 2016;40:57-67. Doi: 10.1093/femsre/fuv040

602 22. Duman R, Ishikawa S, Celik I, Strahl H, Ogasawara N, Troc P, et al.

603 Structural and genetic analyses reveal the protein SepF as a new membrane anchor for 604 the Z ring. *Proc Natl Acad Sci USA*. 2013;110:E4601-E10. Doi:

605 10.1073/pnas.1313978110

23. Zhang L, Willemse J, Claessen D, van Wezel GP. SepG coordinates
sporulation-specific cell division and nucleoid organization in *Streptomyces coelicolor*. *Open Biol.* 2016;6:150164. Doi: 10.1098/rsob.150164

Del Sol R, Mullins JG, Grantcharova N, Flärdh K, Dyson P. Influence of
CrgA on assembly of the cell division protein FtsZ during development of *Streptomyces coelicolor. J Bacteriol.* 2006;188:1540-50. Doi:

612 25. Buddelmeijer N, Beckwith J. A complex of the *Escherichia coli* cell division
613 proteins FtsL, FtsB and FtsQ forms independently of its localization to the septal region.
614 *Mol Microbiol.* 2004;52:1315-27. Doi: 10.1111/j.1365-2958.2004.04044.x

615 26. Boes A, Olatunji S, Breukink E, Terrak M. Regulation of the peptidoglycan
616 polymerase activity of PBP1b by antagonist actions of the core divisome proteins FtsBLQ
617 and FtsN. *mBio*. 2019;10:e01912-18. Doi: 10.1128/mBio.01912-18

618 27. Taguchi A, Welsh MA, Marmont LS, Lee W, Sjodt M, Kruse AC, et al.

619 FtsW is a peptidoglycan polymerase that is functional only in complex with its cognate

penicillin-binding protein. *Nat Microbiol.* 2019;4:587-94. Doi: 10.1038/s41564-0180345-x

Bennett JA, Aimino RM, McCormick JR. Streptomyces coelicolor genes *ftsL* and *divIC* play a role in cell division but are dispensable for colony formation. J
Bacteriol. 2007;189:8982-92. Doi:

625 29. Bennett JÁ, Yarnall J, Cadwallader AB, Kuennen R, Bidey P,

626 **Stadelmaier B, et al.** Medium-dependent phenotypes of *Streptomyces coelicolor* with 627 mutations in *ftsI* or *ftsW. J Bacteriol*. 2009;191:661-4. Doi:

628 30. McCormick JR, Losick R. Cell division gene *ftsQ* is required for efficient

- 629 sporulation but not growth and viability in *Streptomyces coelicolor* A3(2). *J Bacteriol*.
- 630 1996;178:5295-301. Doi:

631 31. Mistry BV, Del Sol R, Wright C, Findlay K, Dyson P. FtsW is a

dispensable cell division protein required for Z-ring stabilization during sporulation
septation in *Streptomyces coelicolor*. *J Bacteriol*. 2008;190:5555-66. Doi:

634 10.1128/JB.00398-08

635 32. Ausmees N, Wahlstedt H, Bagchi S, Elliot MA, Buttner MJ, Flardh K.

SmeA, a small membrane protein with multiple functions in *Streptomyces* sporulation
including targeting of a SpoIIIE/FtsK-like protein to cell division septa. *Mol Microbiol.*2007;65:1458-73. Doi: 10.1111/j.1365-2958.2007.05877.x

- 639 33. Dedrick RM, Wildschutte H, McCormick JR. Genetic interactions of *smc*,
 640 *ftsK*, and *parB* genes in *Streptomyces coelicolor* and their developmental genome
 641 segregation phenotypes. *J Bacteriol*. 2009;191:320-32. Doi:
- 642 34. Wang L, Yu Y, He X, Zhou X, Deng Z, Chater KF, et al. Role of an FtsK643 like protein in genetic stability in *Streptomyces coelicolor* A3(2). *J Bacteriol.*644 2007;189:2310-8. Doi:
- 645 35. Schlimpert S, Flärdh K, Buttner M. Fluorescence time-lapse imaging of the
 646 complete *S. venezuelae* life cycle using a microfluidic device. *J Vis Exp.* 2016:e53863.
 647 Doi: 10.3791/53863
- 648 36. **Stuttard C.** Temperate phages of *Streptomyces venezuelae*: lysogeny and 649 specificity shown by phages SV1 and SV2. *Microbiology*. 1982;128:115-21. Doi:
- Bibb MJ, Domonkos A, Chandra G, Buttner MJ. Expression of the chaplin
 and rodlin hydrophobic sheath proteins in *Streptomyces venezuelae* is controlled by
 sigma(BldN) and a cognate anti-sigma factor, RsbN. *Mol Microbiol*. 2012;84:1033-49.
 Doi: 10.1111/j.1365-2958.2012.08070.x
- 654 38. Kieser T, Bibb MJ, Buttner MJ, Chater KF, Hopwood DA. Practical
 655 Streptomyces Genetics. Norwich, UK: The John Innes Foundation; 2000.
- 656 39. Sambrook J, Fritsch EF, Maniatis T. Molecular Cloning: A Laboratory
 657 Manual. Second ed. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press;
 658 1989.
- 659 40. Datsenko KA, Wanner BL. One-step inactivation of chromosomal genes in
 660 *Escherichia coli* K-12 using PCR products. *Proc Natl Acad Sci U S A*. 2000;97:6640-5.
 661 Doi: 10.1073/pnas.120163297
- 662 41. Gust B, Challis GL, Fowler K, Kieser T, Chater KF. PCR-targeted
 663 *Streptomyces* gene replacement identifies a protein domain needed for biosynthesis of
 664 the sesquiterpene soil odor geosmin. *Proc Natl Acad Sci USA*. 2003;100:1541-6. Doi:
 665 10.1073/pnas.0337542100
- 666 42. Datsenko KA, Wanner BW. One-step inactivation of chromosomal genes in
 667 *Escherichia coli* K-12 using PCR products. *Proc Natl Acad Sci USA*. 2000;97:6640-5.
 668 Doi: 10.1073/pnas.120163297
- 669 43. Gust B, Challis GL, Fowler K, Kieser T, Chater KF. PCR-targeted
 670 Streptomyces gene replacement identifies a protein domain needed for biosynthesis of
 671 the sesquiterpene soil odor geosmin. Proc Natl Acad Sci USA. 2003;100:1541-6. Doi:
 672 10.1073/pnas.0337542100
- 673 44. Cherepanov PP, Wackernagel W. Gene disruption in *Escherichia coli:* TcR
 674 and KmR cassettes with the option of Flp-catalyzed excision of the antibiotic-resistance
 675 determinant. *Gene.* 1995;158:9-14. Doi: 10.1016/0378-1119(95)00193-a

Gregory MA, Till R, Smith MC. Integration site for Streptomyces phage 676 45. 677 phiBT1 and development of site-specific integrating vectors. J Bacteriol. 2003;185:5320-3. Doi: 10.1128/jb.185.17.5320-5323.2003 678 679 46. Donczew M, Mackiewicz P, Wrobel A, Flärdh K, Zakrzewska-680 Czerwinska J, Jakimowicz D. ParA and ParB coordinate chromosome segregation 681 with cell elongation and division during *Streptomyces* sporulation. Open Biol. 682 2016;6:150263. Doi: 10.1098/rsob.150263 Sen BC, Wasserstrom S, Findlay KC, Söderholm N, Sandblad L, von 683 47. 684 **Wachenfeldt C, et al.** Specific amino acid substitutions in β strand S2 of FtsZ cause spiraling septation and impair assembly cooperativity in Streptomyces. Mol Microbiol. 685 686 2019;112:184-98. Doi: 10.1111/mmi.14262 687 Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, 48. 688 **Pietzsch T, et al.** Fiji: an open-source platform for biological-image analysis. Nat 689 Methods. 2012;9:676-82. Doi: 10.1038/nmeth.2019 Bush MJ, Chandra G, Al-Bassam MM, Findlay KC, Buttner MJ. BldC 690 49. 691 delays entry into development to produce a sustained period of vegetative growth in 692 Streptomyces venezuelae. MBio. 2019;10:e02812-18. Doi: 10.1128/mBio.02812-18 693 Kwak J, Dharmatilake AJ, Jiang H, Kendrick KE. Differential regulation 50. of *ftsZ* transcription during septation of *Streptomyces griseus*. J Bacteriol. 694 2001;183:5092-101. Doi: 695 696 Trip EN, Scheffers DJ. A 1 MDa protein complex containing critical 51. 697 components of the Escherichia coli divisome. Sci Rep. 2015;5:18190. Doi: 698 10.1038/srep18190 Datta P, Dasgupta A, Bhakta S, Basu J. Interaction between FtsZ and FtsW 699 52. 700 of Mycobacterium tuberculosis. J Biol Chem. 2002;277:24983-7. Doi: 701 10.1074/jbc.M203847200 Wu KJ, Zhang J, Baranowski C, Leung V, Rego EH, Morita YS, et al. 702 53. Characterization of conserved and novel septal factors in Mycobacterium smegmatis. J 703 704 Bacteriol. 2018;200:e00649-17. Doi: 10.1128/JB.00649-17 Morales Angeles D, Liu Y, Hartman AM, Borisova M, Borges AD, de 54. 705 Kok N, et al. Pentapeptide-rich peptidoglycan at the *Bacillus subtilis* cell-division site. 706 Mol Microbiol. 2017;104:319-33. Doi: 10.1111/mmi.13629 707 Peters K, Schweizer I, Beilharz K, Stahlmann C, Veening JW, 708 55. 709 Hakenbeck R, et al. Streptococcus pneumoniae PBP2x mid-cell localization requires the C-terminal PASTA domains and is essential for cell shape maintenance. Mol 710 Microbiol. 2014;92:733-55. Doi: 10.1111/mmi.12588 711 Sassine J, Xu MZ, Sidiq KR, Emmins R, Errington J, Daniel RA. 712 56. Functional redundancy of division specific penicillin-binding proteins in *Bacillus* 713 subtilis. Mol Microbiol. 2017;106:304-18. Doi: 10.1111/mmi.13765 714 Leclercq S, Derouaux A, Olatunji S, Fraipont C, Egan AJ, Vollmer W, 715 57. et al. Interplay between Penicillin-binding proteins and SEDS proteins promotes 716 bacterial cell wall synthesis. Sci Rep. 2017;7:43306. Doi: 10.1038/srep43306 717 58. Cho H, Wivagg CN, Kapoor M, Barry Z, Rohs PDA, Suh H, et al. 718 Bacterial cell wall biogenesis is mediated by SEDS and PBP polymerase families 719 720 functioning semi-autonomously. Nat Microbiol. 2016;1:16172. Doi:

721 10.1038/nmicrobiol.2016.172

- 722 59. Emami K, Guyet A, Kawai Y, Devi J, Wu LJ, Allenby N, et al. RodA as
- the missing glycosyltransferase in *Bacillus subtilis* and antibiotic discovery for the
- peptidoglycan polymerase pathway. *Nat Microbiol*. 2017;2:16253. Doi:
- 725 10.1038/nmicrobiol.2016.253
- 726 60. Meeske AJ, Riley EP, Robins WP, Uehara T, Mekalanos JJ, Kahne D,
- et al. SEDS proteins are a widespread family of bacterial cell wall polymerases. *Nature*.
 2016;537:634-8. Doi: 10.1038/nature19331
- 729 61. **Pazos M, Peters K, Casanova M, Palacios P, VanNieuwenhze M**,
- 730 **Breukink E, et al.** Z-ring membrane anchors associate with cell wall synthases to
- 731 initiate bacterial cell division. *Nat Commun.* 2018;9:5090. Doi: 10.1038/s41467-018-
- 732 07559-2
- 733 62. Straume D, Piechowiak KW, Olsen S, Stamsas GA, Berg KH, Kjos M,
- et al. Class A PBPs have a distinct and unique role in the construction of the
- pneumococcal cell wall. *Proc Natl Acad Sci USA*. 2020;117:6129-38. Doi:
- 736 10.1073/pnas.1917820117
- 737

738 FIGURE LEGENDS

Figure 1. Construction and complementation of *S. venezuelae* strains mutant for core division genes.

(a) A physical map of the *dcw* cluster in *S. venezuelae* and the genetic locus of *divIC*. Maps 741 of two regions of the S. venezuelae chromosome are shown that contain genes encoding core 742 proteins of the divisome. In each of the two loci, all genes are in the same orientation as the 743 divisome genes. Regions replaced with an apramycin-resistance cassette or an unmarked in-744 frame deletion mutation are shown above the maps. DNA fragments used for constructing 745 genetic complementation plasmids are shown below the map. (b) Phase-contrast microscopy of 746 wild type and mutant phenotypes and mutant phenotypes following genetic 747 748 complementation. All images are phase-contrast micrographs of cover slip impressions from cultures grown for 2 days at 30°C on MYM agar. The top row contains wild type S. venezuelae 749 strain containing the empty complementation vector on the left (wt). Immediately adjacent are 750 shown seven division mutants containing the empty complementation vector pMS82. In the 751 bottom row are shown the seven division mutants containing a complementing fragment cloned 752 into pMS82 which restores sporulation to wild type levels. Scale bar, 5 μ m. (c) Transmission 753 electron micrographs reveal septation and cell wall defects in the *ftsZ*, *ftsI* and *ftsL* mutants. 754 Cells were grown for 2 days at 30°C on MYM agar and thin sections were viewed by 755 756 transmission electron microscopy. Mainly spores were observed for the wild type strain (wt). No examples of vegetative cross-walls and sporulation septa were observed for the *ftsZ*-null mutant. 757 White arrow heads indicate formed unresolved sporulation septa in aerial hyphae for the *ftsL and* 758 ftsI mutants. Scale bar, 500 nm. 759

Figure 2. Double and triple divisome mutants do not have additive or synergistic division phenotypes.

The strains were grown for two days on MYM agar medium at 30°C. Shown are phase-contrast images from impression coverslips of aerial hyphae for double and triple mutant strains. Aerial hyphae of double and triple mutants frequently contain evenly-spaced constrictions as do the

single mutants. The double and triple mutant phenotypes are strikingly similar to the single

mutants (Fig. 1b) and do not result in synthetic division phenotypes. Scale bar, 5 μ m.

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Figure 3. DNA segregation and cell wall phenotypes of *S. venezuelae* division mutants.

Cells were grown for 2 days at 30°C on MYM agar and cover slips were pressed onto confluent lawns. Samples of aerial hyphae were stained for cell wall (green) and DNA (red) and viewed by epifluorescence microscopy. The top row contains corresponding DIC light images. Wild type samples contained mainly spores and spore chains. Examples of aerial hyphae of mutant strains $\Delta ftsZ$, $ftsZ\Delta 2p$, $\Delta ftsL$ and $\Delta ftsI$ are shown. Scale bar, 5 µm.

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Figure 4. Z-ring assembly in sporogenic hypha of *S. venezuelae* divisome mutants.

Batch cultures were grown in a standard fashion in liquid MYM at 30°C and samples were fixed by formaldehyde treatment before cells were mounted for microscopy. Representative micrographs of sporulating hyphae with FtsZ ladders are shown, visualized using YPet-tagged FtsZ. Shown are the wild type control strain and the indicated divisome mutants into which plasmid pKF351[P_{ftsZ} -ftsZ-ypet] had been introduced. Scale bars, 2 µm.

Figure 5. Live-cell imaging of Z-ring assembly in sporogenic hypha of *S. venezuelae* wild type and $\Delta ftsQ$ and $\Delta ftsW$ mutants.

Cultures were grown in MYM at 30°C using a microfluidic system. Representative micrographs of unfixed sporulating hyphae with FtsZ ladders are shown, visualized using YPet-tagged FtsZ. Shown are the wild type control strain and the indicated divisome mutants into which plasmid $pKF351[P_{ftsZ}-ftsZ-ypet]$ had been introduced. Scale bars, 2 µm.

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790 Figure 6. FtsQ and FtsW stabilize Z-rings during sporulation-specific cell division.

791 Shown is a montage of representative time series documenting FtsZ dynamics during spore

formation. Strains were grown in liquid MYM at 30°C using a microfluidic system. Fluorescence

images of FtsZ-YPet signal were obtained from time-lapse microscopy (top) and the

corresponding phase-contrast images are also shown (bottom). Shown are montages of the wild

type control strain and the indicated divisome mutants into which plasmid pKF351[P_{ftsZ} -ftsZ-

ypet] has been introduced. Time intervals between images were kept at 20 min. In addition, zero

797 min was considered as the time wherein the shown hypha had undergone arrest of tip extension

before sporulation septation began. Scale bars, $2 \mu m$.



(b)



(c)







Phase contrast

YPet















AftsW/ftsZ-ypet







AdivIC/ftsZ-ype1







ftsZ⁺/ftsZ-ypet



∆ftsW/ftsZ-ypet





[min]

ftsZ⁺/ftsZ-ypet

 $\Delta ftsQ/ftsZ-ypet$

∆ftsW/ftsZ-ypet

Phase contrast



[min]



Supplementary material

Influence of core divisome proteins on cell division in

Streptomyces venezuelae ATCC 10712

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Table S1. S. venezuelae Strains

Strain Genotypes [†]	ATCC 10712	NRRL B- 65442	Comments [‡]
attB _{ФВТ1} ::pMS82	DU522	DU732	cloning vector integrated by site specific recombination, Hyg ^R
$attB_{\Phi BT1}$::pJS26 ($divIC^+$)	DU620		<i>divIC</i> ⁺ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi BT1}$::pJS7 (ftsIL ⁺)	DU528		<i>ftsIL</i> ⁺ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi BT1}$::pJS10 (ftsL ⁺)	DU538		$ftsL^+$ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi BT1}$::pJS32 (Pdcw-ftsQ ⁺)	DU626		$ftsQ^+$ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi BT1}$::pJS6 ($ftsW^+$)	DU524		$ftsW^+$ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi BT1}$::pJS8 (ftsZ ⁺)	DU534		$ftsZ^+$ genetic complementation vector integrated <i>in trans</i>
$attB_{\Phi C31}$::pKF280 (ypet)	DU576	DU733	ypet cloning vector integrated by site specific recombination, Apra ^R
$attB_{\Phi C31}$::pKF351 (ftsZ-ypet)	DU578	DU724	ftsZ-ypet fusion vector integrated in trans by site specific recombination
ΔdivIC::apra	DU518	DU519	471 bp deletion of <i>divIC</i> including the start and stop codons
ΔdivIC	DU613		471 bp deletion of <i>divIC</i> including the start and stop codons
$\Delta divIC attB_{\Phi BT1}$::pMS82	DU614		cloning vector integrated by site specific recombination in DU613
$\Delta divIC attB_{\Phi BT1}$::pJS26 ($divIC^+$)	DU615		<i>divIC</i> ⁺ genetic complementation vector integrated <i>in trans</i> in DU613
$\Delta divIC attB_{\Phi C31}$::pKF280	DU634		ypet cloning vector integrated by site specific recombination DU613
$\Delta divIC attB_{\Phi C31}$::pKF351 (ftsZ-ypet)	DU635		ftsZ-ypet fusion vector integrated in trans by site specific recombination in DU613
$\Delta divIC \Delta ftsL$	DU592		Mutation in DU613 introduced into DU520
$\Delta divIC \Delta ftsL \Delta ftsQ$	DU628		Mutation in DU629 introduced into DU592
$\Delta divIC \Delta ftsQ$	DU636		Mutation in DU613 introduced into DU629
ΔftsI::apra	DU508	DU509	1,140 bp deletion beginning 60 bp downstream of the <i>ftsI</i> start codon
$\Delta ftsI::frt$	DU679		1,140 bp deletion beginning 60 bp downstream of the <i>ftsI</i> start codon with insertion of 81 bp <i>frt</i> scar
$\Delta ftsI::frt attB_{\Phi BT1}::pMS82$	DU683		cloning vector integrated by site specific recombination in DU679
$\Delta ftsI::frt attB_{\Phi BT1}::pJS7 (ftsIL^+)$	DU691		<i>ftsIL</i> ⁺ genetic complementation vector integrated <i>in trans</i> in DU679
$\Delta ftsI::frt attB_{\Phi C31}::pKF280$	DU681		ypet cloning vector integrated by site specific recombination DU679
$\Delta ftsI::frt attB_{\Phi C31}::pKF351 (ftsZ-ypet)$	DU680		ftsZ-ypet fusion vector integrated in trans by site specific recombination in DU679
ΔftsIL::apra	DU516	DU517	1,817 bp deletion beginning 5 bp downstream of the <i>ftsL</i> start codon through 984 bp downstream of the <i>ftsI</i> start codon
$\Delta ftsIL::apra \ attB_{\Phi BT1}::pMS82$	DU574		cloning vector integrated by site specific recombination in DU516
$\Delta ftsIL::apra attB_{\Phi BT1}::pJS7 (ftsIL^+)$	DU570		pJS7 integrated at the Φ BT1 attachment site in DU516
ΔftsI::apra ΔftsW	DU594		Mutation in DU521 introduced into DU508
Δ <i>ftsL</i> ::apra	DU510	DU511	352 bp deletion beginning 5 bp downstream of the <i>ftsL</i> start codon
ΔftsL	DU520		352 bp deletion beginning 5 bp downstream of the <i>ftsL</i> start codon
$\Delta ftsL attB_{\Phi BT1}$::pMS82	DU544		cloning vector integrated by site specific recombination in DU520
$\Delta ftsL attB_{\Phi BT1}$::pJS10 (ftsL ⁺)	DU546		$ftsL^+$ genetic complementation vector integrated <i>in trans</i> in DU520
$\Delta ftsL attB_{\Phi C31}$::pKF280	DU588		ypet cloning vector integrated by site specific recombination DU520
$\Delta ftsL attB_{\Phi C31}$::pKF351 (ftsZ-ypet)	DU590		ftsZ-ypet fusion vector integrated in trans by site specific recombination in DU520
$\Delta ftsL \Delta ftsQ::apra$	DU622		Mutation in DU520 introduced into DU502
ΔftsQ::apra	DU502	DU503	810 bp deletion beginning at the second codon of $ftsQ$
$\Delta ftsQ$	DU629		810 bp deletion beginning at the second codon of $ftsQ$
$\Delta ftsQ$ attB _{$\Phi BT1::pMS82$}	DU630		cloning vector integrated by site specific recombination in DU629

$\Delta ftsQ attB_{\Phi BT1}$::pJS32 (Pdcw-ftsQ ⁺)	DU631		$ftsQ^+$ genetic complementation vector integrated <i>in trans</i> in DU629
$\Delta ftsQ$ attB _{ΦC31} ::pKF280	DU632		ypet cloning vector integrated by site specific recombination DU629
$\Delta ftsQ$ attB _{ΦC31} ::pKF351 (ftsZ-ypet)	DU633		ftsZ-ypet fusion vector integrated in trans by site specific recombination in DU629
ΔftsW::apra	DU512	DU513	1,076 bp deletion beginning 140 bp downstream of the <i>ftsW</i> start codon
$\Delta ftsW$	DU521		1,076 bp deletion beginning 140 bp downstream of the <i>ftsW</i> start codon
$\Delta ftsW attB_{\Phi BT1}$::pMS82	DU548		cloning vector integrated by site specific recombination in DU521
$\Delta ftsW attB_{\Phi BT1}$::pJS6 (ftsW ⁺)	DU550		pJS6 integrated at the ΦBT1 attachment site in DU521
$\Delta ftsW attB_{\Phi C31}$::pKF280	DU584		ypet cloning vector integrated by site specific recombination DU521
$\Delta ftsW attB_{\Phi C31}$::pKF351 (ftsZ-ypet)	DU586		<i>ftsZ-ypet</i> fusion vector integrated <i>in trans</i> by site specific recombination in DU521
ΔftsZ::apra	DU500	DU669 [1]	844 bp deletion beginning 16 bp upstream of the <i>ftsZ</i> start codon
$\Delta ftsZ::frt$	DU665		844 bp deletion beginning 16 bp upstream of <i>ftsZ</i> with insertion of 81 bp <i>frt</i> scar
$\Delta ftsZ::apra attB_{\Phi BT1}::pMS82$	DU637	DU671	cloning vector integrated by site specific recombination in DU500/DU699
$\Delta ftsZ::apra attB_{\Phi BT1}::pJS8 (ftsZ^+)$	DU536	DU670	$ftsZ^+$ genetic complementation vector integrated in <i>trans</i> in DU500/DU699
ftsZ2p::apra	DU504	DU505	6 bp deletion of the -10 site (TAGTGT) of a developmentally regulated promoter
$ftsZ\Delta 2p$	DU523		TAGTGT of -10 site of <i>ftsZ2p</i> replaced with ACTAGA
$ftsZ\Delta 2p \ attB_{\Phi BT1}$::pMS82	DU552		cloning vector integrated by site specific recombination in DU523
$ftsZ\Delta 2p \ attB_{\Phi BT1}$::pJS8 ($ftsZ^+$)	DU554		<i>ftsZ</i> ⁺ genetic complementation vector integrated <i>in trans</i> in DU523

[†]Unmarked gene deletions contain an in-frame six base scar (ACTAGA) ‡All strains were made for this study.

Vector/	Description [†]	Reference or source
Sv-4-G01	Source of <i>ftsI</i> , <i>ftsL</i> , <i>ftsW</i> , <i>ftsO</i> , and <i>ftsZ</i> (division and cell wall locus, <i>dcw</i>)	M. Bibb. unpublished
Sv-5-C06	Source of <i>divIC</i>	M. Bibb, unpublished
pIJ773	Source of <i>oriT apra</i> cassette	[2]
pIJ799	Source of cassette with oriT apra flanked with bla homology	[2]
pKF280	Control plasmid with promoter-less <i>ypet</i> , integrates at $attB_{\Phi C_{31}}$ (Apra ^R)	[3]
pKF351	Derivative of pKF280 containing <i>ftsZ-ypet</i>	[3]
pMS82	Cloning vector for genetic complementation, integrates at $attB_{\Phi BT_1}$ (Hyg ^R)	[4]
pJK1	Δ <i>ftsZ::apra-oriT</i> , Sv-4-G01 derivative	This study
pJK3	$ftsZ\Delta 2p::apra-oriT$, Sv-4-G01 derivative	This study
pJK5	Δ <i>ftsI::apra-oriT</i> , Sv-4-Go1 derivative	This study
pJK6	Δ <i>ftsL::apra-oriT</i> , Sv-4-G01 derivative	This study
pJK7	Δ <i>ftsW::apra-oriT</i> , Sv-4-G01 derivative	This study
pJK9	Δ <i>ftsIL::apra-oriT</i> , Sv-4-G01 derivative	This study
pJK23	$ftsZ\Delta 2p \ bla::apra-oriT,$ Sv-4-G01 derivative	This study
pJK26	Δ <i>ftsL bla::apra-oriT</i> , Sv-4-G01 derivative	This study
pJK27	Δ <i>ftsW bla::apra-oriT</i> , Sv-4-G01 derivative	This study
pJS6	pMS82 containing <i>ftsW</i> (5,434 bp <i>Pvu</i> II fragment)	This study
pJS7	pMS82 containing <i>ftsI</i> (5,686 bp <i>Pvu</i> II fragment)	This study
pJS8	pMS82 containing <i>ftsZ</i> (3,900 bp <i>Bam</i> HI- <i>Pst</i> I fragment)	This study
pJS10	pMS82 containing <i>ftsL</i> (2,869 bp <i>Kpn</i> I fragment)	This study
pJS18	$\Delta divIC::apra-oriT$, Sv-4-Co6 derivative	This study
pJS24	$\Delta divIC \ bla::apra-oriT$, Sv-4-Co6 derivative	This study
pJS26	pMS82 containing <i>divIC</i> (679 bp PCR fragment beginning 187 bp upstream of <i>divIC</i>)	This study
pJS30	$\Delta ftsQ::apra-oriT$, Sv-4-G01 derivative	This study
pJS32	pMS82 carrying <i>P</i> _{dcw} - <i>ftsQ</i> (1095 bp PCR fragment containing 257 bp of <i>Pdcw</i> and <i>ftsQ</i>)	This study
pJS39	$\Delta ftsQ \ bla::apra-oriT, Sv-4-Go1 \ derivative$	This study
pJS40	Δ <i>ftsZ::frt bla::apra-oriT</i> , Sv-4-G01 derivative	This study
pJS41	Δ <i>ftsI::frt bla::apra-oriT</i> , Sv-4-Go1 derivative	This study

Table S2.	Cosmids	and	plasmids	used i	in 1	the study	

[†]Apr^R, apramycin resistance; Hyg^R, Hygromycin resistance; Unmarked gene deletions contain an in-frame six base scar (ACTAGA)

Name	Sequence	Application
	GCGGCGCGAACCAACGCGCGGCGACGACACGTA	
SVftsZ60	ACTCGAGATTCCGGGGGATCCGTCGACC	ftsZ deletion in cosmid
	TGATGTTGGCCTCGGGGGGGGGGGGCCTCGCTGAC	Sv-4-G01
SVftsZ59	CAGCTGTGTAGGCTGGAGCTGCTTC	
	AGGTTCGGCGTGTTCGTTGAACGTGCGCCACTTG	
SV2pUP66	TCGACTACTAGT ATTCCGGGGGATCCGTCGACC	<i>ftsZ2p</i> "-10" deletion in cosmid
	GGTTACCAGTGTCTCTGTTCGCTGGACTCTTCCG	Sv-4-G01
SV2pDOWN65	AACAGGTCTAGA TGTAGGCTGGAGCTGCTTC	
	CGGGGCGCCGAGCGGATCCGGGAAGACGTCCAG	
SVftsL64	TGAGCACTAGTATTCCGGGGGATCCGTCGACC	<i>ftsL</i> deletion in cosmid
	CCGGCTTCGAGGGCGTCGGGGACGGCTTCGGGG	Sv-4-G01
SVftsL65	CCTCCGCTCTAGATGTAGGCTGGAGCTGCTTC	
	TCCACGAGCGGGGCGCGGAAGGCCTGGGACCGGC	
SVftsW64	CGCTCACTAGTATTCCGGGGGATCCGTCGACC	ftsW deletion in cosmid
	CGGCGCAGGGCCAGGGCCGCTTTCGCGGCGGGT	Sv-4-G01
SVftsW63	TCCTGTCTAGATGTAGGCTGGAGCTGCTTC	
	GCGCCGCCGCGTTCCCGGACCCGCCCGGCCCGC	
SVftsI60	GCGCCCCATTCCGGGGGATCCGTCGACC	<i>ftsI</i> deletion in cosmid
	GCGTCAGGTACCAGGTCTCGTGGTCGACGTCGTC	Sv-4-G01
SVftsI59	CTTGAATGTAGGCTGGAGCTGCTTC	
	AGGCGTAGCGCGGCGGCTGAGGGCAGGAGGCGC	
ftsQSXFor	CAGGTGACTAGTATTCCGGGGGATCCGTCGACC	<i>ftsQ</i> deletion in cosmid
	CTGGCCAACCAGGGTGCTGGCCAGGGGTGATAC	Sv-4-G01
ftsQSXRev	CCGTCATCTAGATGTAGGCTGGAGCTGCTTC	
	GTGCGGGGACGTCCGCGTGAACAGGGGAGGCGA	
divICSXF	CACGACACTAGTATTCCGGGGGATCCGTCGACC	<i>divIC</i> deletion in cosmid
	TTCTCTCGGTTGCCTTGCTCGTCTGGTGCGGAGG	Sv-5-C06
divICSXR	AGGGGTCTAGATGTAGGCTGGAGCTGCTTC	
	CGCTCGCCCGTACCTCCGCGCCGGATCTGAGAG	Unmarked deletion of <i>dcw</i>
SVdcwPSXF	GGCGCAACTAGTATTCCGGGGGATCCGTCGACC	locus between <i>Pdcw</i> and <i>ftsQ</i> in
	CGCGCCGCTCTTCTCGGCGGTCGTCGGTCCGGCT	cosmid Sv-4-G01
SVdcwPSXQR	GCCACTCTAGATGTAGGCTGGAGCTGCTTC	
22ftsQEcoRV	GGAATTCGATATCGAGGGGACAAAGAACCGCAT	PCR amplify Pdcw-ftsQ for
22ftsQSpeI	GGAATTCACTAGTGCTGGCCAGGGGTGATACCC	complementation
divICcompEcoRV	GGAATTCGATATCCATCGAGGAGATCCTCGAC	PCR amplify <i>divIC</i> with native
divICcompSpeI	GGAATTC ACTAGT GTTTCTCTCGGTTGCCTTGC	promoter for complementation

Table S3. Oligonucleotides used in this study



Figure S1. Macroscopic phenotypes of isolated S. venezuelae cell division mutants.

Shown are patches of wild type *S. venezuelae* ATCC 10712 and congenic divisome mutant strains after growth on MYM agar for 4 days at 30°C. Pairs of mutant strains are shown containing either empty integration vector (pMS82) or a cognate genetic complementation vector, respectively. The wild type parent strain shown also contains the empty vector pMS82 (WT).



Figure S2. Colony phenotypes of *ftsZ*-null mutant, complemented null mutant and wild type strains.

Shown are streak plates of *ftsZ*-null mutant (DU500), wild type and *ftsZ*-null mutant containing the genetic complementation vector pJS8 (DU537) grown for 2 days on MYM agar at 30°C.



Figure S3. Western blot analysis of wild type and *ftsZ*-null mutant strains using polyclonal anti-FtsZ antibody.

Cultures of wild type and *ftsZ*-null mutant (DU500) were grown overnight in MYM liquid medium. Cell extracts were obtained by bead mill homogenization and normalized amounts of total protein was fractionated in duplicate on 12% polyacrylamide-SDS gel. Fractionated proteins were either stained with Coomassie brilliant blue (lower panel) or transferred to a solid support for a western blot analysis (upper panel). FtsZ was detected using a polyclonal antibody raised against FtsZ from *S. coelicolor* [5] and a secondary antibody conjugated with alkaline phosphatase.

(a)

S. venezuelae S. coelicolor	CGGGTATcacccCtGGCCAGCACCCTGGTTGGcCAGCGCTACGGgTGATCACATAGGG gaCGGGTATacgtgCaGGCCAGCACCCTGGTTGGgCAGCGCTACGGcTGATCACATAGGG	58 60
	ftsZ3p ##	
S. venezuelae	TGAAAAGAAA AA CGGGAGGTTCGGCGTGTTCGTTGAACGTGCGCCACTTGTCGACTTAGT	118
S. coelicolor	TGAAAAGAAA AA CGGGAGGTTCGGCGTGTTCGTTGAACGTGCGCCACTTGTCGACTTAGT	120
	ftsZ2p ftsZ1p ###	
S. venezuelae	GTCCTGT TCG GAAGAGTCCAgcGAACAGAGACACTGGTAACCCTAAACTTCAaCGT TA GG	178
S. coelicolor	GTCCTGT TCG GAAGAGTCCAagGAACAGACACACTGGTAACCCTAAACTTCAgCGT TA GG	180
S. venezuelae	GTTtGGGTCGGCGtTtCGGACCGTCCCAATCGGCATCcGTCGgaGcGgCGCGa AcC	234
S. coelicolor	GTTcGGGTCGGCGcTaCGGACCGTCCCAATCGGCATCaGTCGtcGgGtCGCGgggggcAtC	240
S. venezuelae	AacGCgcgGcgac GACACGTAACTCGAGGCGAGAGGCCTTCGAC	278
S. coelicolor	AgtGCttcGgcggCCGGGCGACACGTAACTCGAGGCGAGAGGCCTTCGAC	290

(b)



Figure S4. Comparison of the nucleotide sequences for *S. coelicolor* and *S. venezuelae ftsQ-ftsZ* intergenic regions containing the developmental *ftsZ2p* promoter and the sequence of the constructed *S. venezuelae ftsZ2p* promoter mutation.

(a) Nucleotide sequences of the entire intergenic regions upstream of *ftsZ* from *S. venezuelae* (278 bases) and *S. coelicolor* (290 bases) were aligned and the 3 promoters mapped in *S. coelicolor* are indicated (hash marks and bolded transcription start sites). Conserved sequences are in capital letters and divergent sequences are in small letters. (b) Five of the six residues at the -10 sequence of *ftsZ2p* (TAGTGT) for *S. venezuelae* were mutated to ACTAGA in this study (-10 sequences are bolded). The introduction of this *ftsZd2p* mutation into *S. venezuelae* ATCC 10712 created strain DU523.



Figure S5. Spores and aerial hyphae are less robust in mutants for *ftsQ*, *ftsL*, *divIC*, *ftsW* and *ftsI* than in the wild type.

Strains were grown for 4 days on MYM agar at 30°C. Impression coverslip lifts were prepared and representative phase-contrast micrographs are shown for wild type and mutant strains. Arrowheads indicate frequent lysed regions of hyphae and spore compartments for the mutants, which are not typically observed for the wild type. Scale bar, 5 μ m.



Figure S6. The average length of mature spore and spore-like cells resulting from completed development-associated division events from wild type and mutant strains. Strains were grown for 4 days on MYM agar and spores and spore-like cells were harvested in a standard fashion. Samples for the wild type, containing pMS82 (empty vector), and mutant strains, containing either pMS82 or a genetic complementation vector containing the cognate divisome gene, were spotted onto pads of 1% agarose and images were captured by phase-contrast microscopy. The lengths of spores and spore-like cells were measured. The data represent averages from 3 technical replicates. Bars show the standard deviation. For each strain, N = 750. A one-way ordinary ANOVA analysis with Dunnett's multiple comparisons test showed that the mutants all differ from the wild type with p<0.0001. The average spore length of each divisome mutant is significantly different from the wild type while the average spore length of each complemented strain is not.



Figure S7. Developmental genome segregation is not overtly affected in core divisome mutants.

Strains were grown for 2 days on MYM agar at 30°C and impression coverslip lifts were made. The top row contains corresponding DIC light images. Samples of aerial hyphae were stained for cell wall (green) and DNA (red) and viewed by epifluorescence microscopy. Wild type samples contained mainly spores. Examples of aerial hyphae of mutant strains are shown. Aerial hyphae of the mutants typically contain segregated nucleoids with few anucleate compartments. Scale bar represents 5 μ m.



Figure S8. Intermediate plasmid constructions used to make a complementation plasmid for a $\Delta ftsQ$ mutant.

A complementation plasmid was constructed to put expression of the *ftsQ* gene directly under the control of the native *dcw* locus promoter (pJS32). Starting with cosmid Sv-4-G01, recombineering was used to delete 9 *dcw* genes located between *ftsQ* and the *dcw* promoter generating pJS42. The unique *SpeI* and *XbaI* restriction sites, introduced in the process of making pJS42, were digested and religated resulting in pJS57. pJS57 was used as template for PCR to generate complementation plasmid pJS32.

Movie legends

Movie S1: Time-lapse fluorescence microscopy of FtsZ-YPet in sporulating wild-type *S*. *venezuelae*. Shown is a time-lapse microscopy experiment consisting of fluorescence images of FtsZ-YPet (right) and the corresponding phase-contrast images (left) of DU578 ($ftsZ^+$ $attB_{\phi C31}$::pKF351[P_{ftsZ} -ftsZ-ypet]). Cells were grown at 30°C in MYM medium in a microfluidic system and monitored by fluorescence microscopy. After an initial period (4-6 hours) of vegetative growth, spent medium was administered to induce sporulation. The time interval between each frame is 10 min. Experiment was run two times. Scale bar, 2 µm.

Movie S2: Time-lapse fluorescence microscopy of FtsZ-YPet in sporulating *S. venezuelae* $\Delta ftsQ$ mutant. Shown is a time-lapse microscopy experiment consisting of fluorescence images of FtsZ-YPet (left) and the corresponding phase-contrast images (right) of DU633 ($\Delta ftsQ$ $attB_{\phi C3}$::pKF351[P_{ftsZ} -ftsZ-ypet]). Cells were grown at 30°C in MYM medium in a microfluidic system and monitored by fluorescence microscopy. After an initial period (4-6 hours) of vegetative growth, spent medium was administered to induce sporulation. The time interval between each frame is 10 min. Experiment was run two times. Scale bar, 5 µm.

Movie S3: Time-lapse fluorescence microscopy of FtsZ-YPet in sporulating *S. venezuelae* $\Delta ftsW$ mutant. Shown is a time-lapse microscopy experiment consisting of fluorescence images of FtsZ-YPet (right) and the corresponding phase-contrast images (left) of DU586 ($\Delta ftsW$ $attB_{\phi C31}$::pKF351[P_{ftsZ} -ftsZ-ypet]). Cells were grown at 30°C in MYM medium in a microfluidic system and monitored by fluorescence microscopy. After an initial period (4-6 hours) of vegetative growth, spent medium was administered to induce sporulation. The time interval between each frame is 10 min. Experiment was run two times. Scale bar, 5 µm.

Supplementary references

- 1. Santos-Beneit F, Roberts DM, Cantlay S, McCormick JR, Errington J. A mechanism for FtsZ-independent proliferation in *Streptomyces. Nat Commun.* 2017;8:1378. Doi: 10.1038/s41467-017-01596-z
- 2. **Gust B, Challis GL, Fowler K, Kieser T, Chater KF.** PCR-targeted *Streptomyces* gene replacement identifies a protein domain needed for biosynthesis of the sesquiterpene soil odor geosmin. *Proc Natl Acad Sci USA*. 2003;100:1541-6. Doi: 10.1073/pnas.0337542100
- 3. **Donczew M, Mackiewicz P, Wrobel A, Flärdh K, Zakrzewska-Czerwinska J, Jakimowicz D.** ParA and ParB coordinate chromosome segregation with cell elongation and division during *Streptomyces* sporulation. *Open Biol.* 2016;6:150263. Doi: 10.1098/rsob.150263
- 4. **Gregory MA, Till R, Smith MC.** Integration site for *Streptomyces* phage phiBT1 and development of site-specific integrating vectors. *J Bacteriol*. 2003;185:5320-3. Doi: 10.1128/jb.185.17.5320-5323.2003
- 5. Schwedock J, McCormick JR, Angert EA, Nodwell JR, Losick R. Assembly of the cell division protein FtsZ into ladder-like structures in the aerial hyphae of *Streptomyces coelicolor. Mol Microbiol.* 1997;25:847-58. Doi: 10.1111/j.1365-2958.1997.mmi507.x