Chromosome Dynamics in Bacteria: Triggering Replication at the Opposite Location and Segregation in the Opposite Direction

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ABSTRACT  Maintaining the integrity of the genome is essential to cell survival. In the bacterium Caulobacter crescentus, the single circular chromosome exhibits a specific orientation in the cell, with the replication origin (ori) residing at the pole of the cell bearing a stalk. Upon initiation of replication, the duplicated centromere-like region parS and ori move rapidly to the opposite pole where parS is captured by a microdomain hosting a unique set of proteins that contribute to the identity of progeny cells. Many questions remain as to how this organization is maintained. In this study, we constructed strains of Caulobacter in which ori and the parS centromere can be induced to move to the opposite cell pole in the absence of chromosome replication, allowing us to ask whether once these chromosomal foci were positioned at the wrong pole, replication initiation and chromosome segregation can proceed in the opposite orientation. Our data reveal that DnaA can initiate replication and ParA can orchestrate segregation from either cell pole. The cell reconstructs the organization of its ParA gradient in the opposite orientation to segregate one replicated centromere from the new pole toward the stalked pole (i.e., opposite direction), while displaying no detectable viability defects. Thus, the unique polar microdomains exhibit remarkable flexibility in serving as a platform for directional chromosome segregation along the long axis of the cell.

IMPORTANCE  Bacteria can accomplish surprising levels of organization in the absence of membrane organelles by constructing subcellular asymmetric protein gradients. These gradients are composed of regulators that can either trigger or inhibit cell cycle events from distinct cell poles. In Caulobacter crescentus, the onset of chromosome replication and segregation from the stalked pole are regulated by asymmetric protein gradients. We show that the activators of chromosome replication and segregation are not restricted to the stalked pole and that their organization and directionality can be flipped in orientation. Our results also indicate that the subcellular location of key chromosomal loci play important roles in the establishment of the asymmetric organization of cell cycle regulators.

KEYWORDS  Caulobacter crescentus, DnaA, ParA, centromere, chromosome replication, chromosome segregation

Triggering the onset of chromosome replication under the right conditions is central to cell survival and proliferation. In bacteria, DnaA is the highly conserved initiator of chromosome replication that opens the origin of replication (ori) by forming a helical right-handed polymeric structure (1–5). Once DnaA opens the double-stranded chromosome, the replication machinery assembles at ori and initiates chromosome replication bidirectionally. The subcellular location where DnaA initiates chromosome replication is established by the position of ori inside the cell. Depending on the bacterial species, the subcellular location of ori varies significantly. For instance, ori in Caulobac-
*Caulobacter crescentus* (referred to hereafter as *Caulobacter*) is found at one pole (the stalked pole), whereas in *Escherichia coli* and *Bacillus subtilis*, ori is found near mid-cell. Within a single species, the subcellular location of ori is strictly retained at the same position in nondividing cells and reestablished soon after chromosome replication and segregation initiate in actively dividing cells.

In nondividing *Caulobacter*, ori is retained near the stalked pole by the interaction between the anchoring protein PopZ and the ParB partition protein bound to the centromere-like chromosomal parS (referred to hereafter also as the centromere) (6–8). DnaA initiates replication at ori near the stalked pole (Fig. 1). Once chromosome replication initiates at ori, the replication fork must pass through the centromere chromosomal region (parS, 8 kb away from ori) for chromosome segregation to initiate (9). One of the two newly replicated ParB-coated centromeres is segregated to the opposite pole (also referred to as the new pole) by direct interactions between ParB and the ATPase ParA (9–14). It has been proposed that ParA forms a stable gradient with concentrations gradually decreasing from the new pole to the stalked pole, which are critical for establishing segregation directionality (10–12, 15).

In this report, we asked whether the onset of replication and segregation are restricted to the intrinsic localization of ori and the centromere, which in *Caulobacter* are near the stalked pole. We genetically engineered a *Caulobacter* strain where movement of ori and the centromere can be triggered in the absence of replication initiation (16). Subphysiological levels of DnaA result in translocation of ori away from the stalked pole. In *Caulobacter*, the centromere is the first chromosomal locus to segregate away from the stalked pole (9). Previous analyses of cells expressing subphysiological levels of DnaA (insufficient to initiate replication) revealed a DnaA-dependent and replication-independent segregation of the centromere (16). Subphysiological levels of DnaA cause the unreplicated centromere to move in a ParA-dependent manner from the stalked pole to the new pole (16). We asked whether subphysiological levels of DnaA could also trigger the movement of ori independently of replication. We tracked the localization of ori by constructing a strain with a fluorescent tag to be inserted near ori using the *Yersinia pestis* parS(pMT1) chromosomal sequence and its

**RESULTS**

**Subphysiological levels of DnaA result in translocation of ori away from the stalked pole.** In *Caulobacter*, the centromere is the first chromosomal locus to segregate away from the stalked pole (9). Previous analyses of cells expressing subphysiological levels of DnaA (insufficient to initiate replication) revealed a DnaA-dependent and replication-independent segregation of the centromere (16). Subphysiological levels of DnaA cause the unreplicated centromere to move in a ParA-dependent manner from the stalked pole to the new pole (16). We asked whether subphysiological levels of DnaA could also trigger the movement of ori independently of replication. We tracked the localization of ori by constructing a strain with a fluorescent tag to be inserted near ori using the *Yersinia pestis* parS(pMT1) chromosomal sequence and its
corresponding gene encoding ParB(pMT1) (17). This strain is called PM500 [parS(pMT1) vanA::dnaA ΔdnaA xylX::cfp-parB(pMT1)] with fluorescent tag near ori (~1 kb) with dnaA expression regulated by the vanillate promoter. Cells grown in M2G with vanillate were supplemented with xylose (0.3%) for 1 h and synchronized. Swarmer cells were spotted on 1% agarose M2G pads in the presence of vanillate (250 μM) (top row) or absence of vanillate (bottom row). Cells were imaged with phase-contrast and CFP-mediated fluorescence microscopy every 30 min. The time in minutes is shown above the images. The white arrow indicates the location of the stalk. Bars = 1 μm. (B) Percentage of cells with translocated ori to the middle or new pole over a 3-h span of DnaA depletion. Values are means plus standard deviation (SD) (error bars) percentages from three independent experiments. The average number of cells per replicate was 250.

**FIG 2** Translocation of ori in the absence of chromosome replication. (A) Time lapse of indicator strain [PM500; parS(pMT1) vanA::dnaA ΔdnaA xylX::cfp-parB(pMT1)] with fluorescent tag near ori (~1 kb) with dnaA expression regulated by the vanillate promoter. Cells grown in M2G with vanillate were supplemented with xylose (0.3%) for 1 h and synchronized. Swarmer cells were spotted on 1% agarose M2G pads in the presence of vanillate (250 μM) (top row) or absence of vanillate (bottom row). Cells were imaged with phase-contrast and CFP-mediated fluorescence microscopy every 30 min. The time in minutes is shown above the images. The white arrow indicates the location of the stalk. Bars = 1 μm. (B) Percentage of cells with translocated ori to the middle or new pole over a 3-h span of DnaA depletion. Values are means plus standard deviation (SD) (error bars) percentages from three independent experiments. The average number of cells per replicate was 250.

DnaA’s ability to initiate replication is not restricted to the stalked pole. In Caulobacter, DnaA initiates replication only once per cell cycle and only in stalked cells, which have their ori localized at the stalked pole (Fig. 1). To determine whether DnaA can initiate replication at the opposite cell pole, we analyzed cells that had undergone replication-independent translocation of ori. To track replication initiation, we induced dnaA from a vanillate promoter in cells with ori localized to the opposite cell pole and followed the appearance of two newly replicated ori foci (Fig. 3A; Movie S2). We quantified the frequencies of replication initiation based on the initial location of ori prior to induction of dnaA expression (Fig. 3B). Within 30 min of vanillate supplementation, ~63% of cells with ori localized at the new pole had initiated chromosome replication.

**FIG 3** Replication initiation at the new pole. (A) Two newly replicated ori foci are indicated by red and blue stars. (B) Percentage of cells with replicated ori at different locations relative to the stalk. Values are means plus standard deviation (SD) (error bars) from three independent experiments. The average number of cells per replicate was 250.
replication as evidenced by two clearly separated ori foci. Notably, chromosome replication initiated slower in the subpopulation of cells that retained ori at the stalked pole: ~19% of cells displayed two ori foci after 30 min of vanillate supplementation. After the depletion period, DnaA expression was induced by supplementation of vanillate (250 μM). (Bottom) Within 30 min of DnaA repletion, cells were able to initiate chromosome replication, as evidenced by two ori (cyan) foci. Bars = 1 μm. (B) Onset of chromosome replication starting from the stalked pole, mid-cell, or new pole of PM500. The plot represents the mean ± SD percentage of cells with two ori foci from three independent fluorescence microscopy time-lapses. The average number of cells per replicate was 140. Analyses of two-way analysis of variance (ANOVA) between the frequencies of replication at the stalk pole versus new pole are statistically different at the 30-min and 45-min time points (P < 0.001; P < 0.05).

Centromeres are effectively segregated in the opposite direction. On the basis of our results that DnaA is able to initiate replication from the new pole (Fig. 3), we then asked about chromosome segregation. Can the partitioning system ParABS initiate segregation of the centromere from the new pole toward the stalked pole, which in this case would be in the opposite direction? To test this, we used a Caulobacter strain in which the native parB gene was replaced with the fusion gene encoding cyan fluorescent protein (CFP)-ParB and in which the only copy of dnaA was regulated under the vanillate promoter (PM109) (16). In Caulobacter, the partitioning protein ParB binds directly to the centromere (9, 18). Thus, we can track centromere movement by using cells expressing a functional fusion protein CFP-ParB. When the vanillate inducer is removed from the growth medium of strain PM109, cells are exposed to subphysiological levels of DnaA insufficient to initiate chromosome replication but sufficient to trigger movement of the unreplicated centromere (16). Fluorescent imaging of PM109...
cells depleted of DnaA (incubated without vanillate for 3 h) revealed the localization of a single CFP-ParB focus with the following distribution: ~37% at or near the new pole, ~54% at or near the stalked pole, and ~10% at around mid-cell (Fig. S2), consistent with previous analyses (16). One potential explanation for the difference in frequencies of replication-independent parS-ori translocation between strains PM109 and PM500 (~37% versus ~75%) may have to do with expression of the Y. pestis parB(pMT1). Although we did not observe defects on doubling times and/or viability from expression of parB(pMT1) in Caulobacter, our translocation frequency data suggest that Y. pestis ParB may influence the activity of Caulobacter’s ParA in chromosome segregation. From here on, we use only Caulobacter strains with its native parB fluorescently tagged.

To determine the ability of ParABS to trigger segregation of the centromere in the opposite direction (from the new pole toward the stalked pole), we tracked the movement of CFP-ParB in PM109 cells with translocated unreplicated centromeres subsequent to the addition of vanillate (Fig. 4). Two clearly separated centromeres were observed soon after dnaA expression was reestablished, irrespective of the initial localization of centromere (Fig. 4A; Movie S3). Upon induction of dnaA expression, we observed a similar pattern in the rates of replication initiation when tracking the number of parS centromeres. The appearance of two CFP-ParB foci occurred slightly sooner in cells with a centromere at the new pole compared to cells with a centromere at the stalked pole (Fig. S3).
Notably, cells with two CFP-ParB foci were able to segregate their centromeres to the cell poles, irrespective of the initial localization of centromere prior to dnaA induction (Fig. 4A). Quantification of these data revealed that the rate at which centromeres were segregated to the cell poles were significantly faster in cells with centromeres that departed from the mid-cell (Fig. 4B). These results can potentially be explained based on the shorter distance that the replicated centromeres had to travel from mid-cell to reach the cell poles. The rates of centromere segregation that initiated from either the stalked pole or the new pole were not statistically different. These data suggest that the partitioning protein ParA was able to quickly rearrange its gradient in order to segregate centromeres from mid-cell or from the new pole with no significant delays.

Active ParA is required for centromere segregation in the opposite direction. Caulobacter cells expressing ParA variants unable to hydrolyze ATP cannot segregate their centromere to the cell poles (9, 10, 14). To determine whether centromere segregation observed from the new pole or mid-cell is ParA dependent, we tracked CFP-ParB localization in a merodiploid strain that carries the wild-type allele of parA at the native locus and a dominant-negative mutant parA allele unable to hydrolyze ATP expressed from a xylose-inducible promoter. This dominant-negative allele contains a missense mutation in the ATPase domain (ParA^{D44A}) of ParA that inhibits chromosome segregation (19). To test segregation in the opposite direction, we first allowed cells to translocate their unreplicated centromeres by growing them in growth media devoid of vanillate. The growth media were then supplemented with xylose and vanillate so that replication initiation was induced in the presence of the dominant-negative ParA^{D44A}. Our data revealed that >80% of cells expressing wild-type ParA were able to segregate the centromeres to the poles, as evidenced by one CFP-ParB focus at each pole (Fig. 5A). However, cells expressing the dominant-negative ParA^{D44A} after replication initiated at the opposite cell pole failed to segregate their replicated centromeres as evidenced by two CFP-ParB near each other (Fig. 5A and B). These data strongly suggest that the segregation of the centromere in the opposite direction, from the new pole to the stalked pole, requires an active chromosome segregation machinery.

Relocalization of the centromere locus triggers rearrangement of the ParA gradient. In Caulobacter, ParA forms a visible gradient with concentrations gradually decreasing from the new pole to the stalked pole (10–12, 20) (Fig. 1). Our observation that the centromere could be segregated in the opposite direction suggested that cells with centromeres at the new pole rearrange the gradient of ParA. To determine whether ParA could change the orientation of its gradient, we assessed the localization patterns of ParA using the background of a parA merodiploid strain that carries the wild-type allele of parA at the native locus and a fluorescently tagged parA (ParA-mCherry) under the inducible promoter for xylose (19). We found that the simultaneous overexpression of ParA (native ParA plus ParA-mCherry) and DnaA depletion resulted in chromosome replication initiation in approximately 45% of cells, suggestive of the coregulation of DnaA and ParA observed in B. subtilis (21). However, all cells that retained their unreplicated centromere at the stalked pole displayed the ParA-mCherry gradient that resembles the gradient of wild-type cells (Fig. 6). Notably, cells with unreplicated centromeres that had translocated to the new pole displayed a flipped pattern of ParA-mCherry (high levels at the stalked pole and low levels at the new pole). When the centromere was localized at mid-cell, ParA-mCherry displayed what appears to be two separate gradients with high levels starting from both cell poles (Fig. 6). Thus, our data suggest that the subcellular localization of parS-ori plays a role in dictating the organization of ParA.

PopZ subcellular localization patterns based on parS localization. The swarmer exhibits a single PopZ focus at the pole bearing the flagellum. As the swarmer cell differentiates into a stalked cell and initiates DNA replication, a second PopZ focus is established at the opposite pole (new pole) (6–8). The directionality of ParA’s function in centromere segregation from the stalked pole to the new pole has been proposed to be influenced by the localization of the anchoring protein PopZ (19, 22).
To determine whether the onset of chromosome replication/segregation from the new cell pole altered PopZ localization dynamics, we tracked the localization of PopZ by using cells expressing a functional fusion protein mCherry-PopZ. In our control experiment with cells grown in the presence of vanillate (the dnaA inducer), mCherry-PopZ exhibited foci at each pole upon the onset of chromosome replication and segregation (6–8, 16). In cells with translocated unreplicated centromeres, mCherry-PopZ localization was dependent on the localization of CFP-ParB bound to the parS centromere (Fig. 7A). About 90% of cells with CFP-ParB at the stalked pole displayed a single mCherry-PopZ focus also localized at the stalked pole. In cells with CFP-ParB localized at the new pole, ~90% displayed PopZ-CFP foci located at each pole. Notably, cells with CFP-ParB localized at mid-cell displayed an equal combination of cells with either one mCherry-PopZ focus localized at the stalked pole or mCherry-PopZ foci localized at each pole. Upon the induction of chromosome replication by the addition of the inducer vanillate, cells with bipolar localization of PopZ remained bipolar (Fig. 7B). Regardless of where the centromere was localized when replication initiation was induced by vanillate supplementation, ~100% of cells displayed bipolar localization of mCherry-PopZ subsequent to the onset of chromosome replication. Our data suggest that the initiation of chromosome replication and segregation influence the bipolar localization of PopZ.

**Effects in viability from initiating chromosome replication/segregation from outside the stalked pole.** To determine whether initiating chromosome replication and/or segregation from outside the stalked pole altered the viability of *Caulobacter*, we analyzed CFU of cells that had undergone ori or centromere translocation away from the stalked pole in the absence of replication (Fig. 8). Cells of strains PMS00
fluorescent label near ori) and PM109 (fluorescently labeled ParB-centromere) were spotted immediately after DnaA depletion (1h to 3h) on plates containing the inducer for dnaA expression (supplemented with vanillate). Our data revealed no significant differences between cells that initiated chromosome replication and segregation from the new pole (or mid-cell) compared to wild-type conditions. Our data suggest that cells can recover relatively quickly after initiating chromosome replication/segregation from outside the stalked pole.

**FIG 6** Localization of ParA depends on the subcellular localization of parS-ParB. Translocation of centromere to the new pole in DnaA-depleted cells results in flipped ParA-mCherry gradient (red). Green foci represent parS-CFP-ParB (centromeres) localized at the stalked pole, mid-cell, or at the new pole. White arrows indicate locations of the stalks. PM503 cells (parB::cfp-parB, dnaA::T1, vanA::dnaA xylX::parA-mCherry) were synchronized and depleted of DnaA for 3 h. The culture was also supplemented with xylose (0.1%) during the time of DnaA depletion. After DnaA depletion, 2 μl of cells was mounted on 1% agarose pad and imaged using phase-contrast fluorescence microscopy. Micrographs were spliced to show cells with a single parS-CFP-ParB focus grouped based on the subcellular location of that focus (stalked pole, mid-cell, new pole). Bars = 1 μm.

**FIG 7** Localization of PopZ based on the subcellular organization of parS-ParB. (A and B) Localization of PopZ (red) in DnaA-depleted cells (A) and in 1 h DnaA replete cells (B). Green foci represent CFP-ParB (centromeres). Bars = 1 μm. The graphs display quantification of localization of PopZ in cells depleted of DnaA (A) and DnaA replete cells (B). PM247 (parB::cfp-parB, dnaA::T1, vanA::dnaA xylX::mCherry-popZ) were grown in the absence of vanillate (DnaA depletion) for 3 h and then supplemented with vanillate (DnaA repletion) for 1 h. Cultures were supplemented with xylose (0.1%) to induce the expression of mCherry-PopZ for 1 h prior to isolation of swarmer cells. Phase-contrast fluorescent micrographs were obtained just before and after 1 h of the addition of vanillate. The data represent three independent experiments. The average number of cells per replicate was 200. The bar graphs show the mean plus SD values.
DISCUSSION

By inducing the movement of ori and centromere away from their intrinsic subcellular locations in a replication-independent manner, we have shown that the molecular machinery involved in chromosome replication and segregation are remarkably flexible with respect to their subcellular orientation. We found that the activity of DnaA and ParA are not restricted to a single polar microdomain and can successfully induce chromosome replication and segregation from the opposite polar microdomain composed of distinct client proteins. These results suggest that the chromosomal loci ori and parS play key roles in the localization of cell cycle regulators. Cells that initiate chromosome replication and segregation from outside the stalked pole displayed no detectable viability defects compared to wild-type conditions. Therefore, our data reveal the ability of the cell to rapidly reorganize chromosome orientation along with the set of proteins involved in chromosomal replication and segregation.

DnaA’s activity as replication initiator and cell asymmetry. In the absence of membrane-bound organelles, bacteria rely on proteins organized in gradients to establish cellular polarity and perform asymmetric functions. One example of such organization is the phosphorylated regulator CtrA (CtrA→P), which binds ori and inhibits DnaA from initiating replication at one cell pole (23–25). In predivisional cells, a phospho-signaling relay at the cell poles has been proposed to generate an asymmetric concentration gradient of CtrA→P with the highest levels at the new pole that gradually decrease toward the stalked pole (26). Alterations to this proposed asymmetric concentration gradient of CtrA→P eliminate the asymmetric regulation of DnaA, resulting in cells initiating replication from the new pole (26). Our data revealed that DnaA can also trigger replication initiation from the new pole in cells with altered subcellular location of ori (Fig. 3). Our results can be explained by the undetected levels of CtrA in cells depleted of DnaA (27, 28). This is because DnaA is a transcriptional regulator of gcrA, and GcrA is a transcriptional regulator of ctrA (27–29). Consequently, the expression of ctrA is indirectly dependent on the levels of DnaA. Thus, cells with ori localized at the new pole with depleted levels of DnaA are likely to have no CtrA→P or a minimal CtrA→P gradient that is insufficient to inhibit replication initiation from either pole.

Regulation of periodicity of DnaA activity. Most of the regulators of DnaA activity that have been identified thus far are negative regulators that prevent the overinitiation of chromosome replication. However, positive regulators that trigger DnaA to initiate replication with such efficient periodicity remain limited. This periodicity of DnaA activity is maintained even in E. coli cells that were artificially designed to have two spatially separated ori foci. In those cells, DnaA productively initiated replication synchronously from both ori foci (30). In E. coli and Helicobacter pylori, a recruiter of DnaA has been characterized that promotes the assembly of polymeric DnaA at ori (31–33). Notably, constitutive expression of dnaA in Caulobacter has been shown to

FIG 8 Depletion of DnaA for 3 h does not alter the viability of Caulobacter. (A and B) CFU assays of PM500 cells [parS(PlMT1) vanA dnaAΔ dnaA sytX cfp-parB(pMT1)] (A) and PM109 cells [ΔvanA parB cfp-parB dnaAΔ dnaA (Δ vanA dnaA)] (B). The cultures (3 ml) grown to an OD600 of ~0.3 were washed three times with 1× M2 salts as described in Materials and Methods, and the OD600 was set at ~0.2 in M2G medium (2 ml). DnaA was depleted for 1, 2, and 3 h in separate cultures at 28°C, and the CFU assay was performed. Control sample (c) were not depleted of DnaA and were incubated with vanillate (250 μM) for 3 h. PYE plates supplemented with vanillate (250 μM) were incubated at 28°C for 2 days prior to obtaining the images. The data shown are representative of three independent replicates.
have no effect on the periodicity of DnaA activity, suggesting that dnaA transcriptional regulation is not the principal modulator of DnaA periodicity (34). In Caulobacter, CtrA regulates the spatial activity of DnaA so that chromosome replication initiates only in the stalked cells (24, 25, 34). However, CtrA is not involved in the periodicity of DnaA’s activity (34). Thus, the molecular mechanism that triggers DnaA to initiate chromosome replication with such precise periodicity remains unclear. A hypothetical scenario is that some type of regulator that facilitates this periodicity process is found within the microenvironment at ori’s location at the time of replication initiation. Our data suggest that there is no regulator/modulator of DnaA that is fixed at the stalked pole’s microdomain. We cannot however exclude the possibility that a potential modulator does exist and that this modulator could migrate along with ori because either it binds directly to ori or it is recruited by DnaA. Biochemical characterization of DnaA’s activity at ori is required to identify the mechanism that DnaA uses to regulate its temporal activity with such remarkable accuracy.

**Localization of the centromere dictates the orientation of ParA’s activity.** The partitioning protein ParA is another example of a bacterial protein organized in a gradient to establish cellular polarity. In Caulobacter, ParA forms a gradient with concentrations gradually decreasing from the new pole to the stalked pole (10–12, 20). Interestingly, this stable gradient of ParA is established well before chromosome replication and segregation are initiated (10–12). Thus, the question remains as to what activates this asymmetric organization of ParA. Our data suggest that the organization of the ParA gradient can be reconstructed in the opposite orientation by rearranging the location of the parS-ParB complex (Fig. 6). Cells with parS-ParB at the new pole displayed higher levels of ParA at the stalked pole than at the new pole.

We propose that the localization of the centromere complex directs the arrangement of the ParA gradient. This model is consistent with how ParA partitioning systems segregate low-copy-number plasmids to maintain inheritance. Using in vitro reconstitution assays, the ParB-parS complexes of these plasmids were shown to chase and rearrange the ParA gradient (35–37). For chromosome segregation, the specific component of the centromere complex that triggers the organization of ParA remains to be determined. There are several proteins that bind parS and/or interact with parS-ParB that could serve as potential regulators of ParA’s subcellular organization. One possibility is that ParB itself triggers ParA’s organization by inducing ParA to hydrolyze ATP (9, 10, 14), consistent with what has been observed with partitioning of plasmids (35–37). Another possibility is MlpZ, the inhibitor of FtsZ polymerization, which colocalizes with the parS-ParB complex (38). The replication initiator DnaA is another candidate because it also binds parS and has been proposed to be involved in the onset of centromere segregation (16). Last, the anchoring protein PopZ, which interacts with ParB, has been proposed to be involved in ParA activity and directionality (6–8, 19, 22). Our data suggest that the bipolar localization of PopZ is elicited primarily by centromere segregation, as suggested previously (39). About 90% of cells displayed PopZ bipolarly localized in cells with an unreplicated centromere translocated to the new pole (Fig. 7). However, only upon the onset of chromosome replication did we observe 100% of cells with bipolarly localized PopZ, suggesting that replication initiation may also influence PopZ localization. It remains to be determined whether PopZ plays a role in the organization of ParA.

We have demonstrated that ParA can successfully segregate the parS centromere from the new pole to the stalked pole, which is the reverse direction to that observed in wild-type cells. We propose that once a stable gradient of ParA is formed in cells with translocated unreplicated centromeres, the ParA-DNA interaction relay previously shown to provide the force necessary for centromere segregation (20) can initiate and segregate one centromere in the reverse direction.

**Robustness of cells to reorganize.** Establishment of cellular polarity is required for asymmetric cell division. Notably, the signaling factors involved in establishing polarity in C. crescentus are conserved among bacteria from diverse environmental niches (22,
40–44), like Brucella abortus (causative agent for brucellosis in mammals) (45, 46), Sinorhizobium meliloti (plant symbiont) (47), and Agrobacterium tumefaciens (plant pathogen) (48). However, little is known about how the gradients of these signaling polarity factors are formed or how they function in bacteria with diverse life styles. In this work, we asked what happens to the ability of cells to grow when the organization of the two highly conserved chromosomal loci (ori and parS centromere) are flipped in orientation. We showed that the regulators (DnaA and ParA) can easily adapt to the new locations of these sites and proceed with their activities, and in the case of ParA proceed to orient segregation in the reverse direction. Remarkably, cells were able to recover the “forced” rearrangement of these chromosomal loci and continue to grow with no measurable delays. Our results revealed the robustness and flexibility that cells have to rearrange their signaling polarity factors.

MATERIALS AND METHODS

Bacterial strains and growth conditions. Lists of strains, plasmids, and primers used in this work are provided in Tables S1 and S2 in the supplemental material. Plasmids constructed in this study were created by cloning PCR products amplified using wild-type CB15N (NA1000) or Yersinia pestis KIM5 pMT1 genomic DNA into pNPTS138, pXCHYC-2, or pXCFPN-2 vectors (49). The constructs were transformed into E. coli DH5α cells and grown at 37°C in Luria-Bertani (LB) medium. All primers used for cloning are listed in Table S2. Plasmid carrying cfp-parB(pMT1) was done by the parB(pMT1) gene sequence isolation with KpnI and NheI restriction from PM395 (LSS269) and ligation to the equally treated xylose-inducible integrating plasmid pXCFPN-2 (Kanr) (9). The parA gene was cloned into integrating pXCHYC-2 (Kanr) plasmid under a xylose-inducible promoter to express mCherry-tagged C-terminal protein fusions (49). The Gibson cloning method (50) was used to construct the plasmids used to delete or insert a gene into the Caulobacter genome.

Construction of indicator strain PM500 with fluorescently labeled origin of replication. To track the cellular localization of ori, we engineered a fluorescent tag to be inserted near ori using the Y. pestis parS(pMT1) chromosomal sequence and its corresponding gene encoding ParB(pMT1) (17). To insert the parS(pMT1) site approximately 1 kb far away from the ori, the cloned parS(pMT1) sequence from PM395 (LSS270) and around 600 bp of CCNA0001 C-terminal and CCNA0002 N-terminal sequences were assembled into the pNPTS138 plasmid (9). The parS-ParB(pMT1) system from Yersinia has been previously used in Caulobacter and shown not to interfere with the activity of the native Caulobacter ParABS partitioning system (9). To control the expression of dnaA, we first engineered an additional copy of dnaA to replace the vanA gene, resulting in the expression of dnaA regulated by the VanA promoter (vanA::dnaA) (49). Using this merodiploid strain, the native gene encoding DnaA was deleted, leaving no scars on the genome. The final indicator strain PM500 has the genotype xyLex::cfp-parB(pMT1) parS(pMT1) at nucleotide 1108, dnaA::ΔvanA::dnaA. We refer here to the parS(pMT1) localized near ori simply as ori.

Growth assays. Overnight cultures grown from Caulobacter frozen stocks in M2G liquid medium were diluted to an optical density at 600 nm (OD₆₀₀) of 0.2 (2 ml) in 13-mm glass tubes. Cultures were incubated at 28°C, and the optical density at 600 nm was monitored every hour to monitor the growth rates of bacteria.

Synchronization. A culture of Caulobacter in M2G (15 ml) was inoculated with a saturated overnight M2G culture and grown to an OD₆₀₀ of ~0.3. The medium was supplemented with vanillate (250 μM) and antibiotics as noted. Cells were pelleted using centrifugation at 6,000 rpm for 10 min at 4°C. The cell pellet was resuspended in about 800 μl of 1× M2 salts and mixed well with Percoll (900 μl; Sigma-Aldrich) to generate a density gradient. Swarmer cells (bottom layer) were separated out from the stalked/predvisional cells (top layer) by centrifuging at 11,000 rpm for 20 min at 4°C. Collected swarmer cells were washed twice with cold 1× M2 salts by spinning at 8,000 rpm for 3 min at 4°C and resuspended in M2G medium to the appropriate OD₆₀₀. When cells were not synchronized, the cultures grown to an OD₆₀₀ of ~0.3 were pelleted and washed with 1× M2 salts three times.

Fluorescence microscopy. The cells (1 to 3 μl) were spotted on agar pads (1% agarose in M2G) and imaged using phase-contrast and fluorescence microscopy in Zeiss Axio Observer 2.1 inverted microscope, set up with a Plan-Apochromat 100×/1.40 Oil Ph3 M27 (WD = 0.17 mm) objective, AxioCam 506 mono camera and ZEN lite software. Agar pads supplemented with vanillate (250 μM) were used in time-lapse assays when needed. Images were analyzed using Fiji software (51), and localization of fluorescent foci was counted using the Cell Counter plugin.

CFU assay. The cultures were serially diluted (10-fold) by mixing 10 μl of culture with 90 μl of PYE medium in a sterile 96-well plate. Five microliters of each sample was spotted onto PYE agar (1.5%) plates supplemented with vanillate (250 μM) if needed. CFU counts were obtained from the plates incubated at 28°C for 2 days.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.01002-19.

MOVIE S1, MOV file, 2.7 MB.

MOVIE S2, MOV file, 3.2 MB.
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