

Chromosome Organization and Replisome Dynamics in *Mycobacterium smegmatis*

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ABSTRACT Subcellular organization of the bacterial nucleoid and spatiotemporal dynamics of DNA replication and segregation have been studied intensively, but the functional link between these processes remains poorly understood. Here we use quantitative time-lapse fluorescence microscopy for single-cell analysis of chromosome organization and DNA replisome dynamics in *Mycobacterium smegmatis*. We report that DNA replication takes place near midcell, where, following assembly of the replisome on the replication origin, the left and right replication forks colocalize throughout the replication cycle. From its initial position near the cell pole, a fluorescently tagged chromosomal locus (*attB*, 245° from the origin) moves rapidly to the replisome complex just before it is replicated. The newly duplicated *attB* loci then segregate to mirror-symmetric positions relative to midcell. Genetic ablation of ParB, a component of the ParABS chromosome segregation system, causes marked defects in chromosome organization, condensation, and segregation. ParB deficiency also results in mislocalization of the DNA replication machinery and SMC (structural maintenance of chromosome) protein. These observations suggest that ParB and SMC play important and overlapping roles in chromosome organization and replisome dynamics in mycobacteria.

IMPORTANCE We studied the spatiotemporal organization of the chromosome and DNA replication machinery in *Mycobacterium smegmatis*, a fast-growing relative of the human pathogen *Mycobacterium tuberculosis*. We show that genetic ablation of the DNA-binding proteins ParB and SMC disturbs the organization of the chromosome and causes a severe defect in subcellular localization and movement of the DNA replication complexes. These observations suggest that ParB and SMC provide a functional link between chromosome organization and DNA replication dynamics. This work also reveals important differences in the biological roles of the ParABS and SMC systems in mycobacteria versus better-characterized model organisms, such as *Escherichia coli* and *Bacillus subtilis*.

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During the bacterial cell cycle, chromosome replication and segregation must be coordinated with cell division. Bacteria typically contain a single circular chromosome. DNA replication initiates at a unique origin (*ori*) (1), proceeds bidirectionally around the chromosome, and terminates in the terminus region (*ter*) opposite to *ori* (2). Different models of DNA replication have been described in bacteria based on the subcellular localization and dynamics of the sister replisomes that replicate the left and right chromosome arms. In *Bacillus subtilis* and *Pseudomonas aeruginosa*, the sister replisomes colocalize around midcell, forming a “replication factory,” and the chromosome is spooled through the replisome complex during the process of replication (3, 4). In sharp contrast, in *Escherichia coli*, the replisomes first assemble at *ori* in the midcell region and then separate and move to opposite cell halves before returning to midcell for termination (5). In *Caulobacter crescentus* and *Helicobacter pylori*, the replisomes assemble at *ori* near the old cell pole before moving jointly toward midcell (6, 7); *Myxococcus xanthus* displays similar replisome dynamics, although the two replication forks often split and merge during the DNA replication cycle (8). In these species, the

nonrandom movement of sister replisomes during the process of replication, which has been conserved in distantly related species (9–12), has led to a model in which the DNA polymerases that replicate the left and right replicores function independently of each other and use the chromosome as both a template and a track.

Visualization of fluorescently tagged chromosomal loci in live cells reveals that bacterial chromosomes are spatially organized with individual chromosomal loci occupying specific subcellular addresses along the cell length (13, 14). This spatial chromosome arrangement is restored in the daughter cells after completion of chromosome replication and segregation. Proteins involved in the organization of the chromosome arms along the cell length include the chromosome partitioning system (ParABS) and the structural maintenance of chromosome (SMC) protein (14, 15).

The ParABS system, which is present in diverse bacteria (16), comprises a DNA binding protein (ParB), a Walker-type ATPase (ParA), and centromere-like *parS* sequences located in the *ori*-proximal region of the chromosome. Upon binding to *parS* sites, ParB oligomerizes to form large nucleoprotein complexes called

segrosomes. Segrosomes organize the *ori*-proximal regions of newly replicated chromosomes and facilitate their proper positioning in the future daughter cells, either at the cell center (*B. subtilis*) or close to the cell poles (*C. crescentus*) (14). This process is followed by segregation and condensation of other regions of the chromosome (14). The formation and positioning of ParB complexes depend on ParA. Filaments of ParA undergo ParB-dependent polymerization and depolymerization and may provide motive force for segrosomes within bacterial cells (14, 17), although more recently this interpretation has been contested (18). With the exception of *C. crescentus* (19) and *M. xanthus* (8), deletion of *parA* or *parB* is not lethal. In some species, ParA/ParB deficiency causes only minor defects in chromosome organization and segregation, which might imply redundancy in the systems that control positioning of the chromosome. In other species, such as *Pseudomonas putida* (20) and *Mycobacterium smegmatis* (21, 22), lack of ParA or ParB results in a severe defect in chromosome inheritance, as indicated by the large fraction of anucleate cells (up to 10%). In *M. smegmatis*, ParB binds two *ori*-proximal *parS* sites (20) and ParA and ParB together may localize *ori* around the quarter-cell positions (21).

SMC protein complexes, as well as SMC-like MukBEF proteins in *E. coli* and related bacteria, have been shown to play an important role in chromosome organization, condensation, and segregation (14, 15). SMC deficiency usually leads to global chromosome decondensation and increased formation of anucleate cells (15). In all of the bacteria studied so far, SMC forms variable numbers of discrete subcellular foci (15). In *B. subtilis* (23, 24) and *Streptococcus pneumoniae* (25), SMC is enriched in chromosomal regions surrounding *ori*. ParB may assist the loading of SMC at *ori*, possibly through direct protein-protein interactions (23–25). In *P. aeruginosa* (26) and *M. smegmatis* (27), SMC deficiency results in a weak or no discernible phenotype, respectively.

The spatial organization of DNA replication in bacteria mirrors the organization of the chromosome within the nucleoid, and the replication process itself could play a role in establishing the organization of the chromosome (5, 28). In *E. coli*, the subcellular localization of chromosome replication initiation is dependent on MukB, which mediates the positioning of *ori* and replication forks through an unknown mechanism (6, 29, 30). In *B. subtilis*, Soj (ParA homolog) and Spo0J (ParB homolog) are also involved in the initiation of DNA replication (31, 32). To date, however, the relationship between subcellular chromosome organization and spatiotemporal localization of the replication forks has not been extensively investigated. In a previous study, we measured the single-cell dynamics of chromosome replication, cell growth, and cell division in *M. smegmatis* with a dual-reporter strain expressing fluorescent markers of the cell division septum (Wag31-GFP) and the DNA replisome (mCherry-DnaN) (33). Here, we use fluorescent reporter strains, microfluidics, and quantitative time-lapse microscopy to investigate the spatiotemporal localization of DNA replication forks relative to specific chromosomal loci in single cells of *M. smegmatis*. We find that DNA replication takes place at midcell, where the two replication forks colocalize. A fluorescently tagged chromosomal locus (*attB*, situated 245° from *ori*) is actively pulled to the replisome just prior to its duplication, and the newly duplicated loci segregate to mirror-symmetric positions relative to midcell. We also find that genetic ablation of proteins involved in the maintenance of chromosome organization (ParB and SMC) results in a severe defect in subcellular localization and

movement of DNA replisomes. These observations reveal a functional link between subcellular chromosome organization and spatiotemporal localization of DNA replication in mycobacteria.

RESULTS

Subcellular localization of DNA replisome complexes. We tracked the spatiotemporal dynamics of DNA replication and cytokinesis in *M. smegmatis* by using a dual-reporter strain expressing fluorescent markers of the DNA replisome (mCherry-DnaN) and cell division septum (Wag31-green fluorescent protein [GFP]) (Fig. 1A) (33). The *mCherry-dnaN* gene replaces the *dnaN* gene at the native chromosomal locus and encodes a functional protein, as evidenced by normal (wild-type) growth kinetics and cell morphology (Fig. 1A; see Fig. S1A in the supplemental material). The mCherry-DnaN fusion protein forms bright diffraction-limited foci that dynamically assemble and disassemble inside the cells, signaling the onset and completion of DNA replication, respectively (Fig. 1A) (33). We compared the localization of mCherry-DnaN with other replisome components by GFP-tagging SSB (single-strand DNA-binding protein) (Fig. 1B; see Fig. S1B in the supplemental material) or DnaX (DNA replisome *tau* subunit) (Fig. 1C; see Fig. S1B). In both cases, we found that mCherry-DnaN foci localize close to the SSB-GFP foci (Fig. 1B and D) or DnaX-GFP foci (Fig. 1C; see Fig. S1C). Microscopic analysis of wild-type cultures reveals cells with zero (13%), one (57%), two (27%), or three (4%) mCherry-DnaN foci (Fig. 1A and E). The cell fraction with zero foci is enriched for short cells ($\leq 3 \mu\text{m}$) and long cells (6 to 7 μm), presumably corresponding to the prereplication B period and postreplication D period of the cell cycle, respectively. Cells containing one or more foci, corresponding to the C period of the cell cycle, are predominantly of intermediate length (between 3 and 6 μm). In the small number of cells with three foci, which are most abundant in the fraction of very large cells ($>7 \mu\text{m}$), one is bright and the other two are relatively dim. mCherry-DnaN foci are located near midcell in cells with one focus ($43\% \pm 6\%$ of the cell length) or two foci ($39\% \pm 12\%$ and $56\% \pm 11\%$ of the cell length), and in the latter case, the spatial distribution is broad (Fig. 1F).

The *M. smegmatis* ParB protein binds specifically to two *parS* sequences located near *ori*, multimerizes, and spreads out over the adjacent DNA (21). We tracked the subcellular localization of *ori* in space and time by using a fluorescent reporter strain expressing a ParB-mCherry fusion protein, which forms diffraction-limited foci in the vicinity of *ori*. The *parB-mCherry* gene replaces the *parB* gene at the native chromosomal locus and encodes a functional protein, as evidenced by normal (wild-type) growth kinetics and cell morphology (data not shown). In newborn cells with a single ParB-mCherry focus located around midcell, duplication of the focus is followed by rapid segregation of the two foci toward opposite cell poles until they reach the quarter-cell positions, where they remain for the duration of the cell cycle (see Fig. S1D in the supplemental material). In dual-reporter cells expressing SSB-GFP (replisome marker) and ParB-mCherry (*ori* marker), the first appearance of the SSB-GFP focus, indicating assembly of the DNA replisome complex, is strongly correlated with the position of the ParB-mCherry focus (Fig. 1G and H). Two physically separated ParB-mCherry foci become visible, on average, 5 min after the first appearance of the SSB-GFP focus (Fig. 1I). Immediately thereafter, the newly separated ParB-mCherry foci move rapidly away from the SSB-GFP focus (Fig. 1G). Similar results were ob-

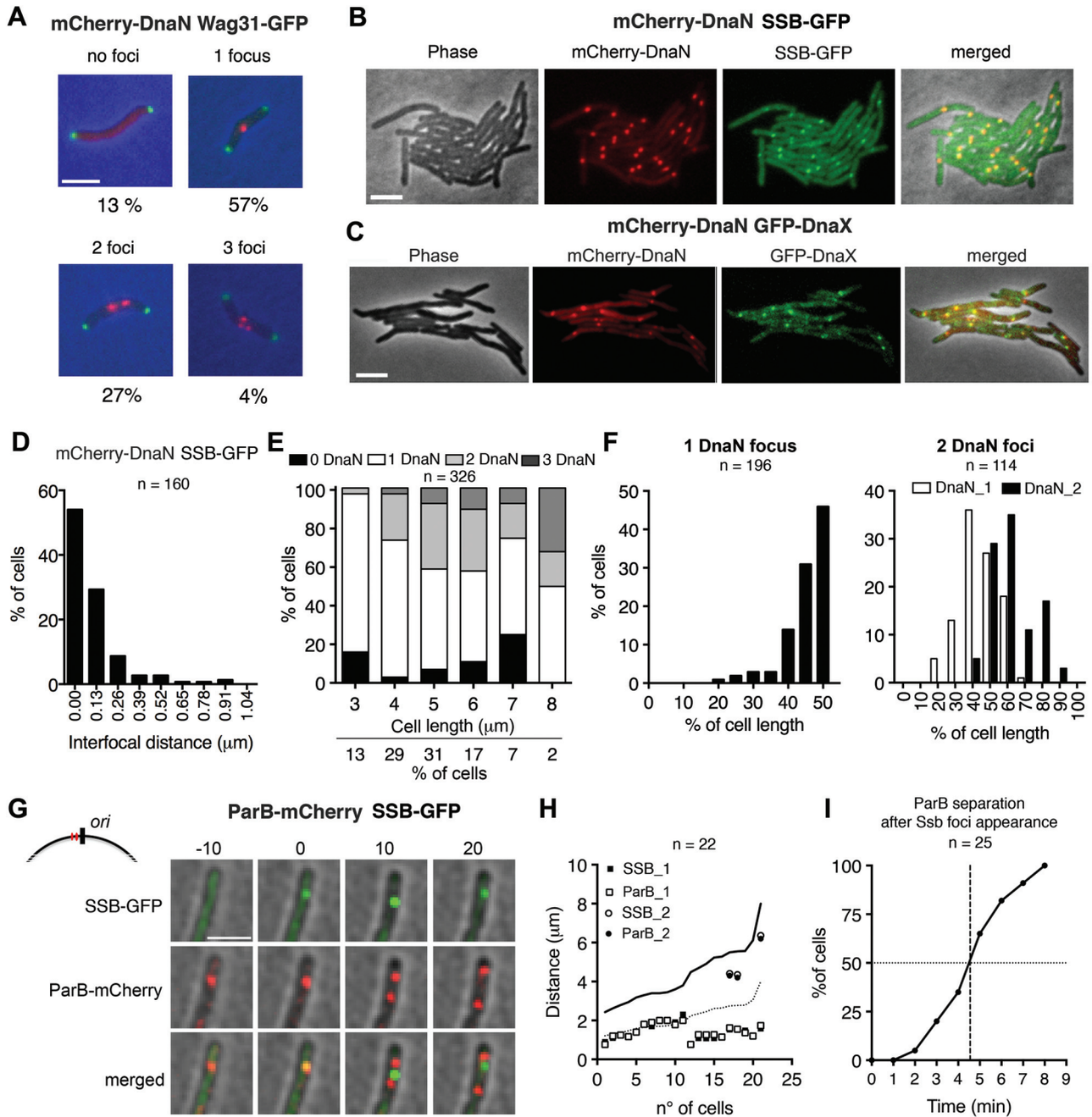


FIG 1 Subcellular localization of DNA replisome complexes. (A) Representative snapshots of wild-type cells with zero, one, two, or three mCherry-DnaN foci (red) and their frequencies ($n = 645$). Green foci, Wag31-GFP. Scale bar, $3 \mu\text{m}$. (B) Colocalization of mCherry-DnaN foci with SSB-GFP foci. Scale bar, $3 \mu\text{m}$. (C) Colocalization of mCherry-DnaN foci with DnaX-GFP foci. Scale bar, $3 \mu\text{m}$. (D) Distribution of interfocal distances between mCherry-DnaN and SSB-GFP foci ($n = 160$). (E) Number of mCherry-DnaN foci relative to cell length ($n = 326$). (F) Position of mCherry-DnaN foci in cells with one ($n = 196$) or two ($n = 114$) foci. (G) Time-lapse analysis of cells expressing ParB-mCherry and SSB-GFP. Images were recorded at 10-min intervals. As shown in the schematic, ParB binds two *parS* sequences (red lines) located in the origin-proximal region (black line). Newly formed SSB-GFP foci colocalize with ParB-mCherry (0 min). Duplication and segregation of ParB-mCherry foci occur shortly (10 min) after replisome assembly, and segregation continues thereafter (20 min). Scale bar, $3 \mu\text{m}$. (H) Positional correlation between ParB-mCherry and SSB-GFP foci at the first appearance of SSB-GFP foci ($n = 22$). DNA replication initiates prior to division in some cells. Solid line, cell length; dotted line, midcell. (I) Time-lapse analysis of cells expressing ParB-mCherry and SSB-GFP. Images were recorded at 1-min intervals. Time intervals between the first appearance of SSB-GFP foci and the separation of ParB-mCherry foci are plotted ($n = 25$).

tained with a dual-fluorescent-reporter strain expressing ParB-mCherry (*ori* marker) and GFP-DnaN (replisome marker) fusion proteins (see Fig. S1E). These observations suggest that *ori* is replicated within 5 min after the first appearance of the replisome

focus; therefore, replication initiation must occur within 0 to 5 min after replisome assembly. These data are also consistent with the notion that the newly duplicated *ori* sequences remain cohered for no more than a few minutes after *ori* is replicated.

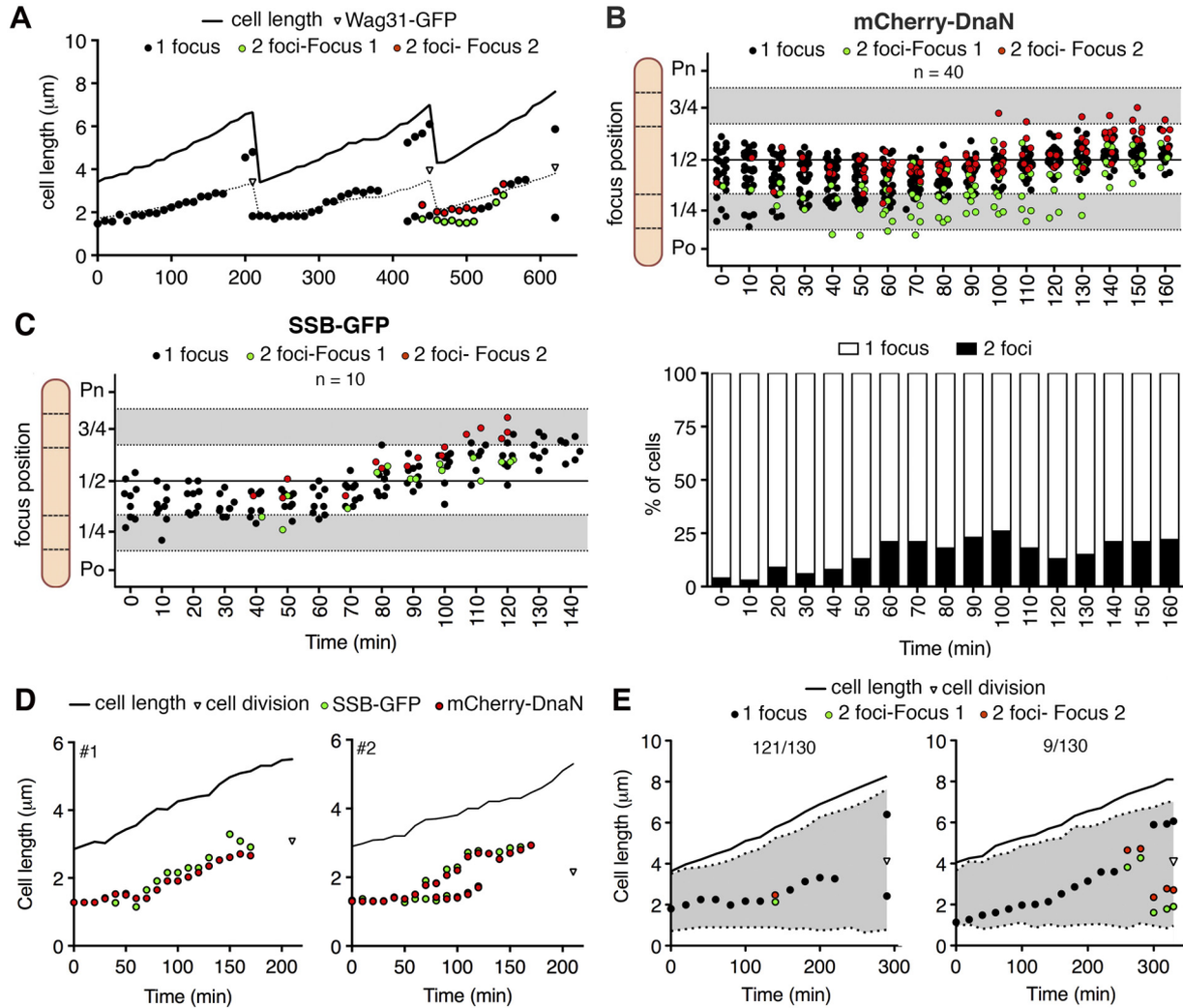


FIG 2 Spatiotemporal tracking of DNA replisomes in single cells. (A) Representative time traces of subcellular positions of overlapping (black circles) or split (green and red circles) mCherry-DnaN foci and Wag31-GFP foci (triangles) relative to the old cell pole. Images were recorded at 10-min intervals. Also see Fig. S2 in the supplemental material. (B) mCherry-DnaN foci during a single DNA replication cycle in 40 time-lapse series. Time zero corresponds to the first appearance of mCherry-DnaN foci. Upper panel, positions of overlapping (black circles) or split (green and red circles) mCherry-DnaN foci; lower panel, fraction of cells with one or two mCherry-DnaN foci at each time point. Po, old cell pole; Pn, new cell pole. (C) Positions of overlapping (black circles) or split (green and red circles) SSB-GFP foci during one DNA replication cycle in 10 time-lapse series. Time zero corresponds to the first appearance of SSB-GFP foci. (D) Two representative time traces of positions of mCherry-DnaN (red circles) and SSB-GFP (green circles) foci relative to the old cell pole. Images were recorded at 10-min intervals. Also see Fig. S2. (E) two representative time traces of the positions of overlapping (black circles) and split (green and red circles) mCherry-DnaN foci and nucleoid stained with SYTO green (grey-shaded area) relative to the old cell pole. Images were recorded at 20-min intervals. Data are representative of 130 time-lapse series in which 121/130 cells show pattern 1 (left panel) and 9/130 cells show pattern 2 (right panel).

Spatiotemporal tracking of DNA replisomes in single cells.

The spatiotemporal distribution of mCherry-DnaN foci suggests that the replication machinery preferentially localizes around midcell and the replication forks colocalize during a large fraction of the cell cycle. We tracked the dynamic behavior of mCherry-DnaN foci in 40 randomly selected cells at 10-min intervals from their appearance to their disappearance, corresponding to replication initiation and termination, respectively (Fig. 2A; see Fig. S2A in the supplemental material) (33). On average, mCherry-DnaN foci are present for 162 ± 26 min, compared with an average interdivision time of 198 ± 32 min, indicating that the replication forks progress with an average speed of about 400 bp/s. At the initiation of DNA replication, a single mCherry-DnaN focus appears around midcell ($44\% \pm 11\%$ of the cell length) and

the replication forks remain colocalized as a single focus for most of the C period, although dynamic splitting and rejoining of replisomes sometimes occur (Fig. 2A and B). When split, the two mCherry-DnaN foci usually remain close to each other around midcell. However, in a small number of cells (3/40), the two foci occupy different cell halves for much of the DNA replication cycle (see Fig. S2A). On average, the net distance traveled by the replisome from assembly to disassembly is about 1/10 of the cell length. For replisomes that exhibit splitting, movement toward the quarter-cell positions is more pronounced (Fig. 2A and B; see Fig. S2A and B). The replisome shows a similarly dynamic localization pattern when SSB-GFP foci are tracked separately in wild-type cells (Fig. 2C) or in parallel with mCherry-DnaN in a dual-reporter strain (Fig. 1B and 2D; see Fig. S2C).

Movement of the replisome foci coincides with replication of the chromosome. We therefore used time-lapse microscopy to track the spatiotemporal dynamics of mCherry-DnaN foci relative to the nucleoid, which can be visualized by staining with SYTO green. In ~92% of the cells, mCherry-DnaN foci track along the middle of the cells and the nucleoid is equally distributed around them (Fig. 2E, cell 1). In the remaining ~8% of the cells, the mCherry-DnaN foci first appear at the edge of the nucleoid closest to the old cell pole, and as the C period progresses, they move toward the middle of the cell (Fig. 2E, cell 2). The latter pattern is similar to the replisome dynamics described for *C. crescentus* (6) and *M. xanthus* (8).

Subcellular localization of *ori* requires ParB. The observation that ParB marks the position of the chromosomal origin and the location of replisome assembly suggests that ParB deficiency might affect the subcellular localization of the replication machinery. As reported previously (21), we found that a *parB* deletion mutant ($\Delta parB$) has a discernible growth defect (see Fig. S3A in the supplemental material) and 15% of the cells are anucleate (see Fig. S3B), suggesting a defect in chromosome segregation. The distributions of interdivision times are similar for wild-type and $\Delta parB$ strains (see Fig. S3C), suggesting that the mutant's growth defect is not due to delayed cell division but rather to the formation of nonviable anucleate cells.

In wild-type cells, the cell division septum is positioned with a narrow spatial distribution that is skewed toward the new cell pole (52 to 58% of the cell length) (see Fig. S3D) (33). In contrast, the division septum can be located anywhere along the cell length in $\Delta parB$ cells, except for 30% of the cell length closest to the old cell pole and 20% of the cell length closest to the new cell pole (see Fig. S3D), which accounts for the broader distribution of cell birth lengths for $\Delta parB$ cells than for wild-type cells (see Fig. S3E). Thus, ParB is important for subcellular septum positioning but has little impact on the timing of septum formation.

ParB deficiency also has an impact on nucleoid structure (see Fig. S3B) (21). The compact and condensed nucleoid bodies observed in wild-type cells appear to be more decondensed, stretched, or elongated in $\Delta parB$ cells. Moreover, in $\Delta parB$ cells, the nucleoid structure is highly irregular and cells with increased DNA content or guillotined chromosomes also occur (see Fig. S3B). Consistent with these observations, some $\Delta parB$ cells contain nucleoids that are longer than wild-type nucleoids (see Fig. S3F).

We investigated the impact of ParB deficiency on the localization of the chromosomal origin at the time of replication initiation. The subcellular positions of newly assembled mCherry-DnaN foci can be used to infer *ori* localization in wild-type and $\Delta parB$ cells because the replisome colocalizes with *ori* at the initiation of chromosome replication (Fig. 1G and H; see Fig. S1E in the supplemental material). The Wag31-GFP marker can be used to distinguish between initiation of DNA replication after (one *ori*) or before (two *ori*) cytokinesis (33). We found that newly assembled mCherry-DnaN foci are frequently mislocalized in ParB-deficient cells. In newborn cells with one *ori*, mCherry-DnaN foci appear at midcell in ~90% of wild-type cells compared to just ~40% of $\Delta parB$ cells, in which mCherry-DnaN foci more frequently appear at the quarter-cell (~50%) and polar (~10%) positions (Fig. 3A; see Fig. S4A). In newborn cells with two *ori*, mCherry-DnaN foci appear at the quarter-cell positions in ~80% of wild-type cells compared to just ~16% of $\Delta parB$ cells, in which

mCherry-DnaN foci more frequently appear at the polar positions (~50%) (Fig. 3A; see Fig. S4B). The aberrant localization of replisomes observed in $\Delta parB$ cells reflects a mispositioning within the nucleoid bodies. In ParB-deficient cells, mCherry-DnaN foci assemble close to the edge of the nucleoid rather than the center of the nucleoid, as observed in wild-type cells (Fig. 3B). Although elongated nucleoids are evident in ~22% of $\Delta parB$ cells, average nucleoid length at initiation of chromosome replication is not significantly different for wild-type and $\Delta parB$ cells (see Fig. S4C). These observations indicate that ParB plays an important role in subcellular positioning of *ori* and replisome assembly, as well as chromosome organization and segregation.

Subcellular localization of *ori* does not require SMC. The SMC protein plays an important role in chromosome organization, condensation, and segregation in *B. subtilis* (23, 24, 34, 35) and *C. crescentus* (36). The *M. smegmatis* genome encodes a single SMC homolog, MSMEG_2423 (referred to as MSMEG_2422 in reference 27). A previous study demonstrated that SMC deficiency did not result in any discernible growth defects, but the effect of SMC deficiency on chromosome organization and segregation was not reported (27). We confirmed that deletion of the *smc* gene (Δsmc) does not significantly affect bacterial growth at the population level (data not shown), although single-cell imaging reveals the presence of small numbers of anucleate cells (~0.3%) and cells containing elongated nucleoids (see Fig. S3F in the supplemental material). We also found that Δsmc cells are slightly defective in septum localization, showing a broader distribution of septum positions than wild-type cells (see Fig. S3D). Subcellular localization of replisome assembly (first appearance of mCherry-DnaN foci) relative to cell length (Fig. 3A; see Fig. S4A and S4B) or nucleoid length (Fig. 3B; see Fig. S4C) is similar in wild-type and Δsmc cells, although the distributions are slightly broader in mutant cells. Similarly, we found that subcellular localization of ParB-mCherry foci (*ori* marker) is indistinguishable in wild-type and Δsmc cells (see Fig. S4D). These observations indicate that SMC plays only a minor role in subcellular positioning of *ori* and replisome assembly, as well as chromosome organization and segregation.

Subcellular localization of the chromosomal *attB* locus requires ParB and SMC. Fluorescently tagged chromosomal loci can be imaged in live cells by using fluorescent repressor-operator systems (FROS) comprising an integrated array of *tetO* operator sequences and a TetR-GFP fusion protein that binds specifically to the *tetO* array. We used a mycobacteriophage L5 integrase-dependent plasmid (see Fig. S4E in the supplemental material) to integrate the *tetO* array at the chromosomal *attB* locus, which is located on the left arm of the chromosome at 245° relative to *ori* (37) (see Fig. S4F). The FROS-*attB* reporter strain has no discernible growth defect compared to wild-type cells (data not shown).

Wild-type cells containing one (~57%) or two (~43%) FROS-*attB* foci (see Fig. S4G) average $4.4 \pm 1.0 \mu\text{m}$ and $5.7 \pm 0.8 \mu\text{m}$ in length, respectively (see Fig. S4H). A large fraction (~15%) of $\Delta parB$ cells exhibit no visible FROS-*attB* foci, consistent with the formation of anucleate cells, and most of these cells are shorter than average (see Fig. S4H). Excluding anucleate cells, the fractions of $\Delta parB$ cells with one FROS-*attB* focus (~55%) or two FROS-*attB* foci (~43%) are similar to those of wild-type cells, but the mutant cells are longer, on average, at 4.7 ± 1.1 and $6.4 \pm 1.2 \mu\text{m}$, respectively (see Fig. S4H). Very long cells (average, $8.7 \pm 2.0 \mu\text{m}$) with three or even four FROS-*attB* foci comprise a small

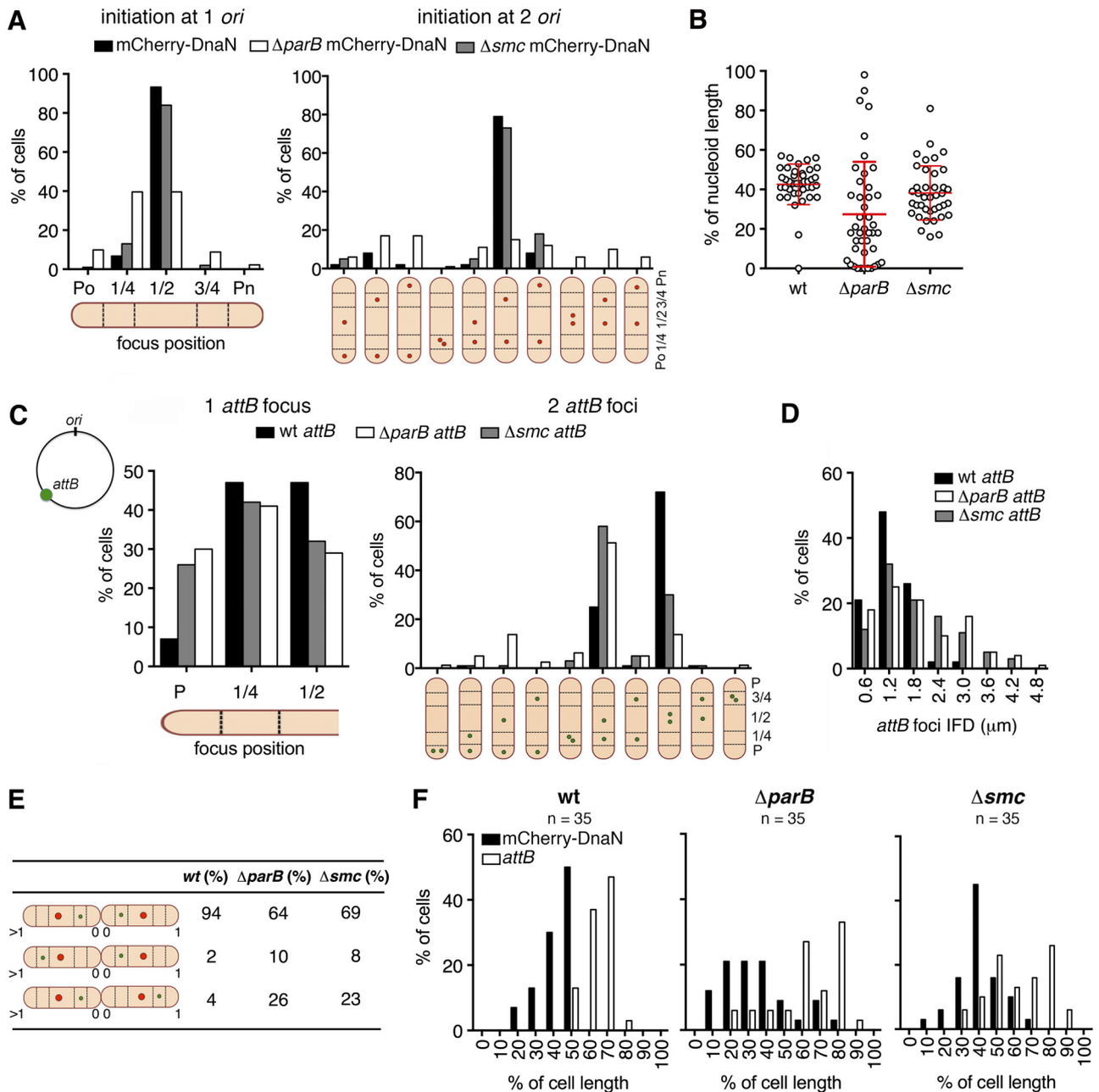


FIG 3 Subcellular localization of *ori* and replisome assembly require ParB but not SMC. (A) Distribution of positions of mCherry-DnaN foci relative to the old cell pole (Po) at initiation of DNA replication in wild-type (black bars), $\Delta parB$ (white bars), and Δsmc (grey bars) cells ($n = 100$ per strain) that initiate replication after cytokinesis (one *ori*, left panel) or before cytokinesis (two *ori*, right panel). Timing of cytokinesis is defined by the first appearance of Wag31-GFP at midcell. Cells that initiate DNA replication before cytokinesis are classified on the basis of the positions of the two mCherry-DnaN foci (schematic). (B) Positions of mCherry-DnaN foci at initiation of DNA replication relative to the SYTO green-stained nucleoid length in wild-type (wt), $\Delta parB$, and Δsmc cells ($n = 40$ per strain). (C) Distribution of positions of FROS-*attB* foci in wild-type (wt; black bars, $n = 135$), $\Delta parB$ (white bars, $n = 112$), and Δsmc (grey bars, $n = 196$) cells containing one FROS-*attB* focus (left panel) or two FROS-*attB* foci (right panel). Cells with two FROS-*attB* foci are classified on the basis of the positions of the two foci (schematic). (D) Distribution of IFDs of FROS-*attB* foci in wild-type (wt; black bars, $n = 305$), $\Delta parB$ (white bars, $n = 351$), and Δsmc (grey bars, $n = 261$) cells with two FROS-*attB* foci. (E) Positions of mCherry-DnaN foci (red circle) and FROS-*attB* foci (green circle) at the time of replisome assembly in sibling cells. Three scenarios and their relative frequencies are shown for wild-type (wt), $\Delta parB$, and Δsmc cells. (F) Distribution of positions of mCherry-DnaN foci (black bars) and FROS-*attB* foci (white bars) at the time of replisome assembly relative to the old cell pole (pole age, ≥ 1) in wild-type (wt), $\Delta parB$, and Δsmc cells ($n = 35$ per strain).

fraction ($\sim 2\%$) of $\Delta parB$ cells but are never observed among wild-type cells (see Fig. S4H).

The subcellular localization of FROS-*attB* foci is also impaired in ParB-deficient cells. In wild-type cells containing a single

FROS-*attB* focus, the focus is usually localized at the midcell or quarter-cell position or less frequently ($\sim 6\%$) at a polar position (Fig. 3C). In $\Delta parB$ cells containing a single FROS-*attB* focus, the focus is more frequently ($\sim 30\%$) localized at a polar position

(Fig. 3C). In wild-type cells containing two FROS-*attB* foci, the foci are usually distributed symmetrically relative to midcell, with average positions at $37\% \pm 5\%$ and $54\% \pm 6\%$ of the cell length (Fig. 3C; see Fig. S4I). In $\Delta parB$ cells containing two FROS-*attB* foci, the foci are usually distributed asymmetrically relative to midcell, with average positions at $23\% \pm 9\%$ and $49\% \pm 13\%$ of the cell length (Fig. 3C; see Fig. S4I). The distribution of interfocal distances (IFDs) between the two FROS-*attB* foci is also broader in $\Delta parB$ cells than in wild-type cells (Fig. 3D). Similar localization patterns are reflected in the position of FROS-*attB* foci relative to the nucleoid length in cells with one focus (see Fig. S4J) or two foci (see Fig. S4K). In $\Delta parB$ cells, the FROS-*attB* foci are located close to the edge of the nucleoid (average position, $21\% \pm 13\%$ relative to the closest cell pole) rather than farther inside the nucleoid, as observed in wild-type cells (average position, $38\% \pm 9\%$ relative to the closest pole) (see Fig. S4K).

Although SMC deficiency has only minor effects on subcellular positioning of *ori* and replisome assembly (see above), we found that subcellular localization of FROS-*attB* foci is markedly altered in Δsmc cells. Indeed, the localization of FROS-*attB* foci in Δsmc cells with one focus (average position, $27\% \pm 12\%$ of cell length) or two foci (average positions, $28\% \pm 9\%$ and $53\% \pm 9\%$ of cell length) is similar to that in $\Delta parB$ cells (Fig. 3C). However, the distribution of subcellular positioning of FROS-*attB* foci is less heterogeneous in Δsmc cells than in $\Delta parB$ cells and no cells with more than two foci occur among Δsmc cells (Fig. 3C and D; see Fig. S4I).

We examined the subcellular localization of the FROS-*attB* locus relative to the replisome at the time of replisome assembly (first appearance of mCherry-DnaN foci) in sibling cell pairs. Three distinct scenarios were observed (Fig. 3E). In wild-type cells at the time of replisome assembly, ~94% of sibling cells show a mirror-symmetric distribution of FROS-*attB* foci relative to the division septum DnaN-*attB*-septum-*attB*-DnaN. The fraction of cells exhibiting an asymmetric distribution of FROS-*attB* foci relative to the division septum DnaN-*attB*-septum-DnaN-*attB* is greater in $\Delta parB$ cells (~36%) and Δsmc cells (~31%) than that in wild-type cells (~6%). In all three strains, asymmetric distribution of FROS-*attB* foci is more common among new-pole siblings (pole age, 1-0) compared to old-pole siblings (pole age, >1-0). On average, the IFD between mCherry-DnaN and FROS-*attB* foci is greater in $\Delta parB$ cells than in wild-type cells, whereas Δsmc cells show an intermediate phenotype due to mislocalization of FROS-*attB* foci (Fig. 3F).

ParB/SMC deficiency affects the number of replisomes per cell. Because ParB and SMC seem to play important roles in chromosome organization, we investigated whether these factors might also affect the number and subcellular localization of replisomes. Microscopic analysis of $\Delta parB$ cultures revealed cells with zero, one, two, three, or four mCherry-DnaN foci (Fig. 4A; cf. Fig. 1A for wild-type cells). The fraction of cells with zero mCherry-DnaN foci is greater among $\Delta parB$ cells (28%) than wild-type cells (13%) and comprises mainly short cells (2 to 3 μm) and long cells ($\geq 7 \mu\text{m}$), presumably corresponding to the prereplication B period and postreplication D period of the cell cycle, respectively (see Fig. S5A in the supplemental material). The fraction of cells with one mCherry-DnaN focus is smaller among $\Delta parB$ cells (21%) than among wild-type cells (57%) and comprises mainly cells of intermediate length (between 3 and 6 μm) (see Fig. S5A). The fraction of cells containing two mCherry-

DnaN foci is similar in $\Delta parB$ cells (30%) and wild-type cells (27%), whereas the fraction of cells containing three mCherry-DnaN foci is higher among $\Delta parB$ cells (17%) than among wild-type cells (4%). Also, $\Delta parB$ cells include a small fraction of long cells containing four usually rather dim mCherry-DnaN foci (3%), which are never observed among wild-type cells. The spatial distribution of mCherry-DnaN foci is broader in $\Delta parB$ cells containing one, two, or three foci (Fig. 4C to E) than in wild-type cells (Fig. 1F), suggesting that ParB plays an important role in replisome positioning. Similar perturbations of the distribution of replisome numbers (Fig. 4B) and their subcellular localization (Fig. 4C to E) occur among Δsmc cells, except that the fraction of cells with zero mCherry-DnaN foci is smaller (11%) and the fraction of cells with three mCherry-DnaN foci is greater (30%) than in $\Delta parB$ cells.

ParB/SMC deficiency affects replisome dynamics. Previously we reported that wild-type *M. smegmatis* cells do not undergo multifork replication, in contrast to cells of better-characterized species such as *B. subtilis* and *E. coli* (33). At the population level, the *ori-ter* ratios are similar in wild-type and $\Delta parB$ strains (D. Jakimovic, personal communication), suggesting that ParB deficiency probably does not result in replication reinitiation and multifork replication. We therefore used time-lapse microscopy to identify the reason for the increased number of mCherry-DnaN foci observed in $\Delta parB$ cells.

Lineage analysis of time-lapse series revealed that ~21% of newborn $\Delta parB$ cells contain two chromosomes as a consequence of missegregation prior to division. In many of these cells, only one replisome assembles and three FROS-*attB* foci are visible at the time of replisome disassembly, indicating that only one of the two chromosomes undergoes replication. Subsequently, these cells divide into daughter cells that inherit one or two chromosomes. Eventually, cells containing two chromosomes at birth divide without undergoing another round of replication, thereby generating two daughter cells with normal ploidy. ParB-deficient cells include a small fraction (~2%) of elongated cells with three or even four chromosomes (see Fig. S5B in the supplemental material). These polyploid cells are characterized by simultaneous rounds of DNA replication on different chromosomes, resulting in the appearance of several mCherry-DnaN foci that often split and converge.

Although chromosome missegregation events may contribute to the increased number of replisomes observed in $\Delta parB$ cells, they cannot account for the large fraction of $\Delta parB$ cells (~80%) with normal ploidy but increased replisome numbers (see Fig. S5C). Nor can missegregation events account for the increased number of mCherry-DnaN foci in Δsmc cells, in which anucleate and polyploidy cells are uncommon (~0.3%). Time-lapse microscopy of $\Delta parB$ cells containing one chromosome at the time of replisome assembly reveals that the fraction of cells with two or three mCherry-DnaN foci increases during the elongation phase of DNA replication (Fig. 4F; see Fig. S6A and Movie S1). In ~30% of $\Delta parB$ cells undergoing DNA replication, the mCherry-DnaN focus splits into two separate foci around 20 to 30 min after initiation. The separated foci are mobile and frequently localize to different cell halves, and many of them split and converge repeatedly, generating cells with three or four foci (Fig. 4F). Similar replisome dynamics occur in Δsmc cells, although the distribution of subcellular positions of mCherry-DnaN foci tends to be narrower (Fig. 4G; see Fig. S6B and Movie S2). On average, the

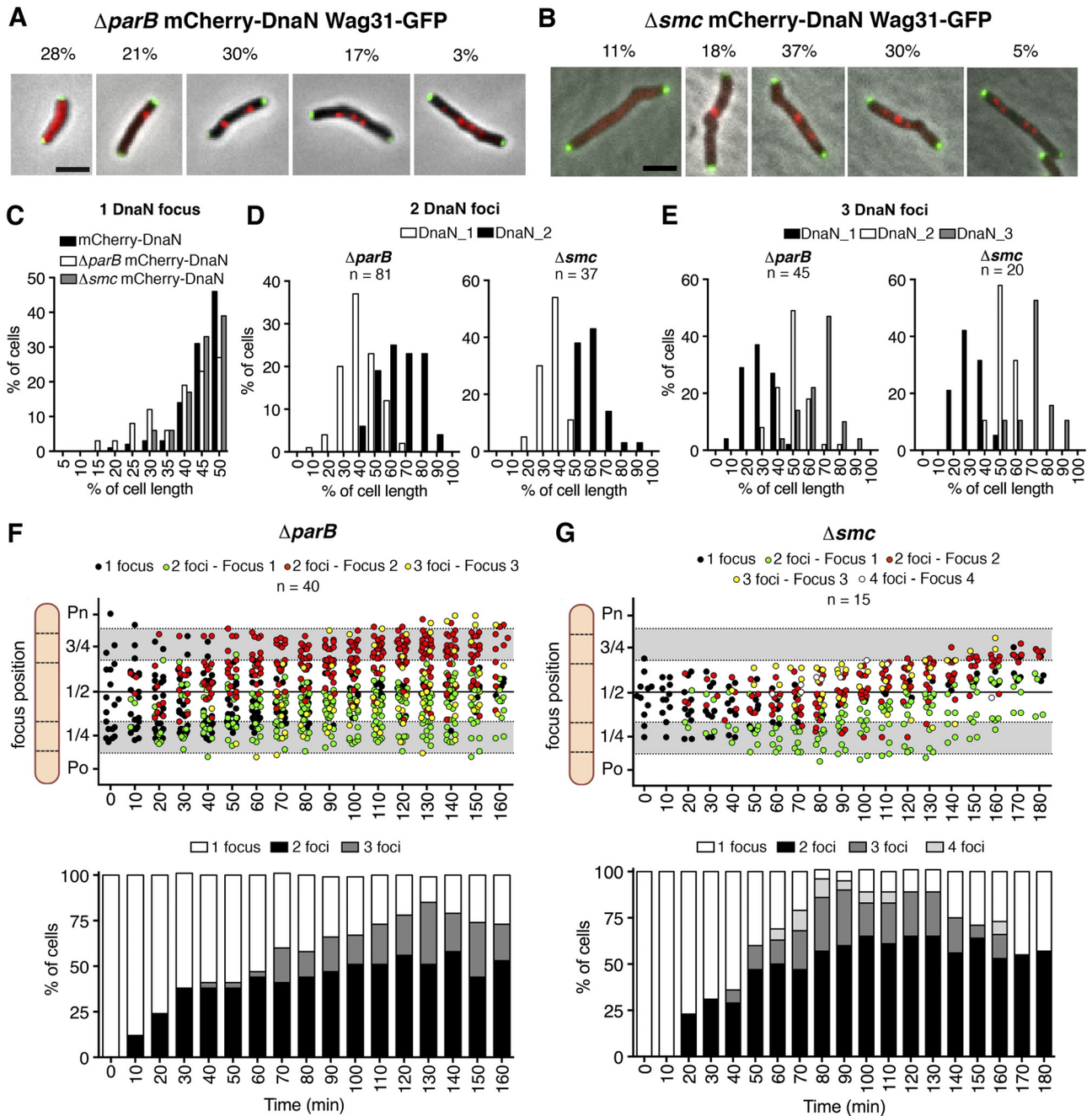


FIG 4 ParB and SMC deficiency affects replisome dynamics. (A and B) Representative snapshots of $\Delta parB$ (A) and Δsmc (B) cells with zero, one, two, three, or four mCherry-DnaN foci and their frequencies ($n = 494 \Delta parB$ cells, $n = 150 \Delta smc$ cells). Green foci, Wag31-GFP. Scale bar, 3 μm . (C) Distribution of positions of mCherry-DnaN foci in wild-type ($n = 196$), $\Delta parB$ ($n = 130$), and Δsmc ($n = 140$) cells with one focus. (D) Distribution of positions of mCherry-DnaN foci in $\Delta parB$ ($n = 81$) and Δsmc ($n = 37$) cells with two foci. (E) Distribution of positions of mCherry-DnaN foci in $\Delta parB$ ($n = 45$) and Δsmc ($n = 20$) cells with three foci. (F and G) Subcellular positions of mCherry-DnaN foci in 20 time-lapse series of $\Delta parB$ cells (F) and 15 time-lapse series of Δsmc cells (G). Images were recorded at 10-min intervals. Time zero corresponds to the first appearance of mCherry-DnaN foci. Upper panels, overlapping replication forks (black circles) and split replication forks (green and red circles); lower panels, fractions of cells with one, two, three, or four mCherry-DnaN foci at each time point. Po, old cell pole; Pn, new cell pole.

IFD of mCherry-DnaN foci during one DNA replication cycle is greater in $\Delta parB$ cells ($1.4 \pm 0.8 \mu\text{m}$; $P < 0.0001$ by Fisher's test) and Δsmc cells ($1.1 \pm 0.8 \mu\text{m}$; $P < 0.01$ by Fisher's test) than in wild-type cells ($0.8 \pm 0.4 \mu\text{m}$) (see Fig. S5D). In a small fraction of mutant cells ($\Delta parB$ and Δsmc), the replisomes do not split, or split and converge only rarely, during the DNA replication cycle

(see Fig. S5E). These observations suggest that ParB and SMC might be involved in the maintenance of cohesion between twin replication forks (and possibly leading- and lagging-strand replisomes) and restriction of the replication machinery to a specific subcellular address, thus generating the "factory-like" replisome organization observed in wild-type cells.

We conclude that the increased numbers of mCherry-DnaN foci observed in $\Delta parB$ and Δsmc cells are due not to multifork replication but rather to polyploidy ($\Delta parB$ cells) and splitting of sister replisomes ($\Delta parB$ and Δsmc cells).

Evidence for a “replication factory” near midcell. Two models of bacterial DNA replication have been proposed (38). In the first model, the replisomes move by tracking along the DNA, which is stationary; in the second model, the replisomes form a localized “replication factory” through which the DNA is spooled as it is replicated. We attempted to distinguish between these models by tracking the movement of replisome (mCherry-DnaN) foci relative to FROS-*attB* foci in wild-type cells (Fig. 5A and B). At the beginning of the C period, the FROS-*attB* focus and mCherry-DnaN focus localize close to the new cell pole and around midcell, respectively. Duplication of the FROS-*attB* focus later in the C period is preceded by a sudden inward movement of the FROS-*attB* focus toward the mCherry-DnaN focus until the two foci colocalize (Fig. 5A). The two newly duplicated FROS-*attB* foci then separate and move from midcell to ~40% and ~60% of the cell length, similar to the positions observed in snapshots of wild-type cells with two FROS-*attB* foci (Fig. 3C). Colocalization of FROS-*attB* and mCherry-DnaN foci occurs, on average, 109 ± 12 min after replication initiation at *ori*. Assuming a constant replication velocity, this value is in good agreement with the expected time of FROS-*attB* replication (~104 min postinitiation), based on the average C-period duration of 164 ± 16 min measured by time-lapse microscopy, the total length of the chromosome (6,988,192 bp), and the distance between *attB* and *ori* (2,236,678 bp). Separation of newly duplicated FROS-*attB* foci occurs, on average, 12 ± 8 min after colocalization of FROS-*attB* and mCherry-DnaN foci, and termination (replisome disassembly) occurs, on average, 52 ± 10 min after colocalization of FROS-*attB* and mCherry-DnaN foci, which is also in good agreement with the expected time to replicate the stretch of DNA between *attB* and *ter* (~57 min). During the early C period, FROS-*attB* foci move randomly with a speed of ~0.1 $\mu\text{m}/\text{min}$, which increases up to ~0.8 $\mu\text{m}/\text{min}$ during the inward movement of FROS-*attB* toward the replisome (Fig. 5B). These observations are consistent with a “replication factory” model, in which the replisome remains positioned near midcell while DNA is pulled toward the replisome, spooled through it, duplicated, and extruded (3).

Increased replisome movement in ParB- and SMC-deficient cells. Increased replisome mobility and splitting in ParB- and SMC-deficient cells suggests that, in the absence of constraints that restrict the replisome to a specific subcellular position, the split replisomes might move independently along the DNA. We addressed this possibility by tracking the movement of replisome (mCherry-DnaN) foci relative to FROS-*attB* foci in $\Delta parB$ cells. We found that in some cells the FROS-*attB* locus localizes near the old cell pole prior to duplication, and newly duplicated FROS-*attB* loci sometimes segregate asymmetrically with respect to midcell (for example, cells 3 and 4 in Fig. 5C). In these cases, one locus moves rapidly toward the new cell pole while the other locus shows little change in its subcellular position over time. In contrast to wild-type cells (Fig. 5B), in ~50% of $\Delta parB$ cells, the FROS-*attB* focus does not show any sudden movement toward the replisome before its duplication (Fig. 5C). Instead, the FROS-*attB* focus remains localized around the quarter-cell position while one of the two replisome foci (presumably the one replicating the left arm of the chromosome) moves toward the FROS-*attB* focus and

colocalizes with it just prior to its duplication (for example, cell 2 in Fig. 5C).

These observations suggest that in $\Delta parB$ cells the normal positioning of the replisome at midcell is lost and the replisome tracks along the DNA (6). Consistent with this interpretation, the net average movement of the mCherry-DnaN and FROS-*attB* foci during the 90-min period after replication initiation is $0.08\% \pm 0.06\%$ and $0.05\% \pm 0.05\%$ of the cell length in wild-type cells (Fig. 5D), compared to $0.24\% \pm 0.14\%$ and $0.13\% \pm 0.09\%$ of the cell length in $\Delta parB$ cells where the replisome does not undergo splitting (Fig. 5E). Movement of the mCherry-DnaN and FROS-*attB* foci is similarly increased in the fraction of $\Delta parB$ cells where the replisome does undergo splitting and the split foci move independently (Fig. 5E). In these cells, net movement of the split mCherry-DnaN foci is $0.47\% \pm 0.05\%$ of the cell length, whereas movement of the FROS-*attB* foci is $0.09\% \pm 0.13\%$ of the cell length, similar to that in wild-type cells (Fig. 5E).

The Δsmc strain behaves somewhat differently. In the fraction of Δsmc cells with split replisome foci, movement of the split foci relative to each other is $0.47\% \pm 0.07\%$ of the cell length, whereas movement of the FROS-*attB* foci is $0.08\% \pm 0.06\%$ of the cell length, similar to that in $\Delta parB$ cells (Fig. 5E). However, in the fraction of Δsmc cells with a single (unsplit) replisome focus, the movement of mCherry-DnaN and FROS-*attB* foci is $0.08\% \pm 0.06\%$ and $0.05\% \pm 0.05\%$ of the cell length, respectively, similar to that in wild-type cells (Fig. 5D).

Despite the striking heterogeneity of single-cell behavior, these observations suggest that in a large fraction of ParB- and SMC-deficient cells the replisome undergoes splitting and the split replisomes move through the cytoplasm as if tracking along the DNA. We have never observed this dynamic behavior in wild-type cells, where the replisome remains unsplit and positioned around midcell, appearing to spool the DNA through it rather than tracking along the DNA.

ParB is required for the normal subcellular localization of SMC. Phenotypic similarities between the *M. smegmatis* $\Delta parB$ and Δsmc mutants suggested that ParB and SMC might have overlapping roles in chromosome organization and replisome positioning. This interpretation is consistent with the observation that SMC foci colocalize with ParB foci in *B. subtilis* (24). We tracked the spatiotemporal dynamics of SMC localization in *M. smegmatis* by using reporter strains expressing an *smc-gfp* fusion gene, which replaces the wild-type *smc* gene at the native chromosomal locus. The fusion protein appears to be functional, as evidenced by the normal (wild-type) growth kinetics of the SMC-GFP reporter strain (see Fig. S3G in the supplemental material).

In wild-type cells, a variable number of discrete SMC-GFP foci are distributed throughout the cell, except at the polar positions (Fig. 6A). In ParB-deficient cells, the SMC-GFP fusion protein does not form bright fluorescent foci but rather appears to be dispersed in patches throughout the cytoplasm (Fig. 6A). The more diffuse pattern of SMC-GFP fluorescence in $\Delta parB$ cells is not due to cleavage of the SMC-GFP fusion protein and release of free GFP, as confirmed by immunoblot analysis (Fig. 6B). Although we found that SMC-GFP and ParB-mCherry foci seldom colocalize, they are frequently observed in close proximity to each other (Fig. 6C). Similarly, although SMC-GFP foci are frequently observed in close proximity to mCherry-DnaN foci, colocalization is uncommon (Fig. 6D).

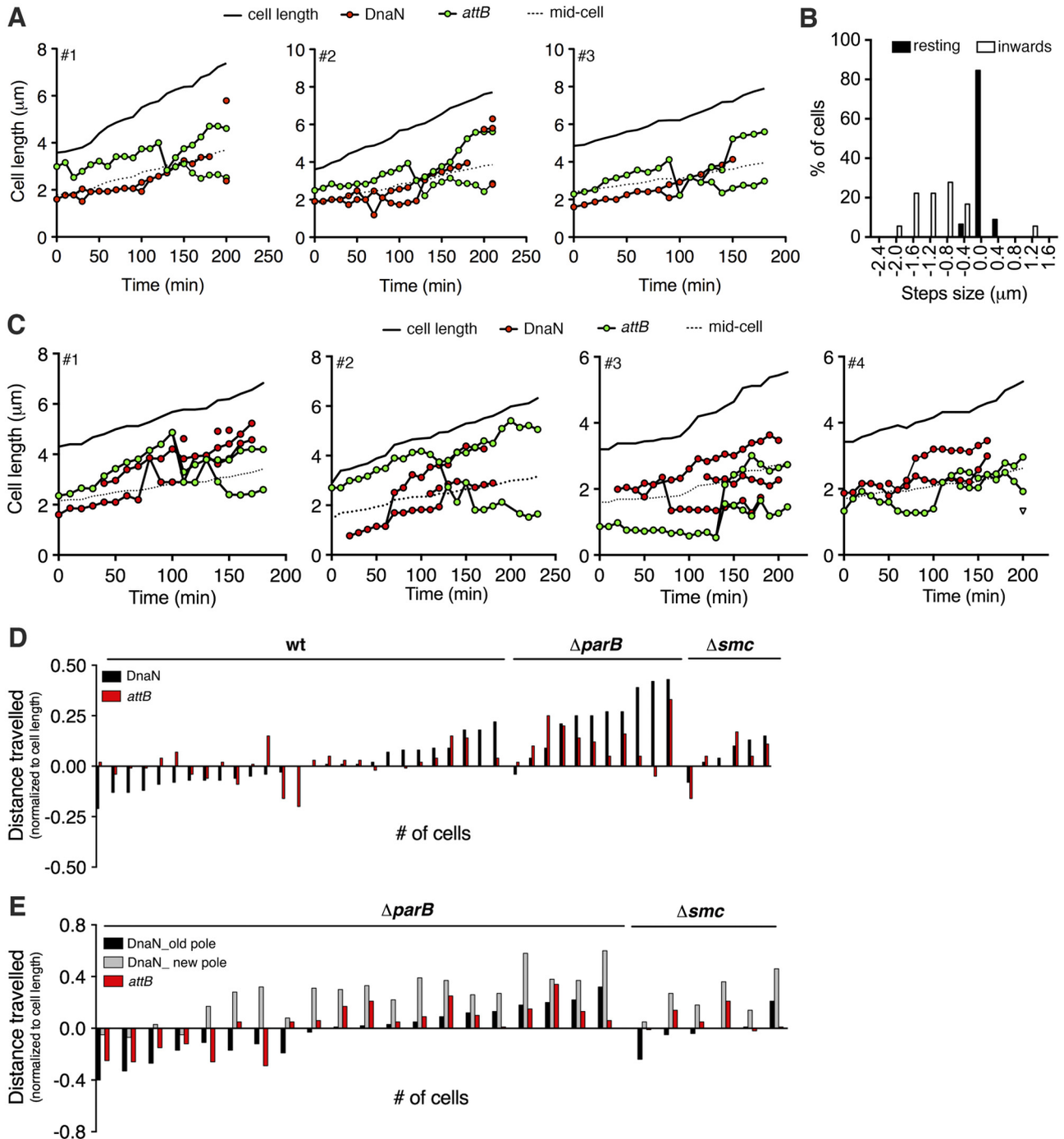


FIG 5 Altered movement of mCherry-DnaN and FROS-*attB* foci in ParB- and SMC-deficient cells. (A) Three representative time traces of positions of mCherry-DnaN (red circles) and FROS-*attB* (green circles) foci relative to the old cell pole. Images were recorded at 10-min intervals. Solid line, cell length; dotted line, midcell. See also Fig. S3B in the supplemental material. (B) Distribution of step sizes of FROS-*attB* foci. Images were recorded at 2-min intervals. Black bars, distribution of steps during the resting period prior to movement toward the replisome ($n = 168$). White bars, distribution of steps during movement toward replisome ($n = 18$). (C) Four representative time traces of positions of mCherry-DnaN (red circles) and FROS-*attB* (green circles) foci relative to the old cell pole in $\Delta parB$ cells. Images were recorded at 10-min intervals. Solid line, cell length; dotted line, midcell. See also Fig. S3B in the supplemental material. (D) Net movement of mCherry-DnaN foci (black bar) relative to FROS-*attB* foci (red bars) during the 90-min period after the first appearance of the replisome in wild-type, $\Delta parB$, and Δsmc cells with unsplit mCherry-DnaN foci. (E) Net movement of mCherry-DnaN foci (grey and black bars) relative to FROS-*attB* foci (red bars) during the 90-min period after the first appearance of the replisome in wild-type (wt), $\Delta parB$, and Δsmc cells with split mCherry-DnaN foci that move independently.

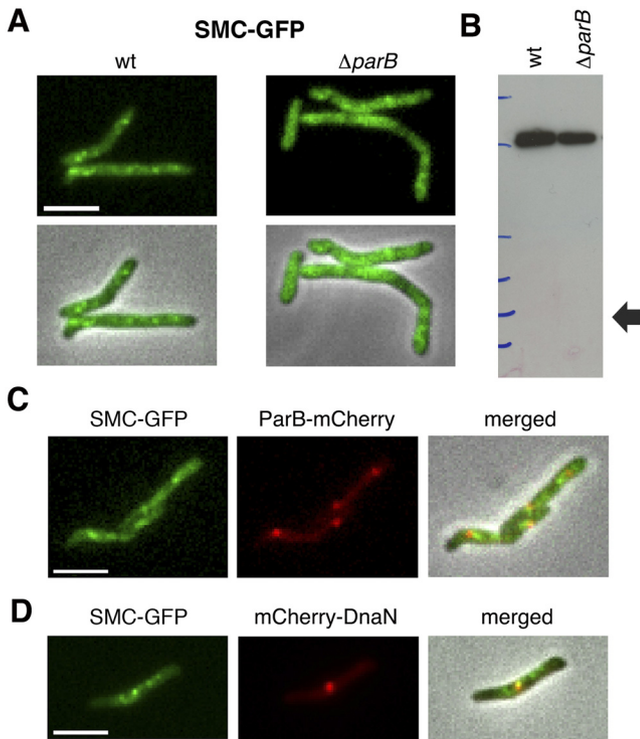


FIG 6 ParB deficiency affects subcellular localization of SMC-GFP. (A) Representative images of SMC-GFP foci in wild-type (wt) and $\Delta parB$ cells. Signal intensity is normalized to facilitate comparison. (B) Immunoblot assay of SMC-GFP in wild-type (wt) and $\Delta parB$ strains with anti-GFP antibody. The arrow shows the predicted position of GFP. (C) Representative images of SMC-GFP (green) and ParB-mCherry (red) foci in exponentially growing wild-type cells. Scale bar, 3 μm . (D) Representative images of SMC-GFP (green) and mCherry-DnaN (red) foci in exponentially growing wild-type cells. Scale bar, 3 μm .

DISCUSSION

In this study, we report that ParB and SMC coordinate chromosome organization and spatial localization of DNA replication in *M. smegmatis*. In wild-type cells, the replisome is positioned near midcell, the two replication forks colocalize throughout the DNA replication cycle, and the chromosomal *attB* locus moves rapidly from cell pole to midcell immediately before its duplication. These observations are consistent with the “replication factory” model proposed for *B. subtilis* and *P. aeruginosa* (3, 4). However, movement of the *attB* locus from cell pole to midcell is faster in *M. smegmatis* (~2 to 4 min) than the movement of a fluorescently tagged chromosomal locus in *B. subtilis* over a similar distance (~30 min) (3). We hypothesize that this movement could be due to the replisome functioning as a DNA-translocating molecular motor that “reels in” the *attB* locus once the slack in the intervening stretch of DNA has been taken up by the process of replication.

Prior to replication, the *M. smegmatis* chromosome is arranged in space such that *ori* is positioned near midcell while the *attB* locus on the left arm of the chromosome is in close proximity to the new cell pole, suggesting a speculative R-*ori*-L configuration (Fig. 7) (39). Confirmation of this configuration will require additional experiments with multiple fluorescently tagged loci on both the left and right arms of the chromosome. Maintenance of the wild-type chromosome configuration, which is recapitulated

with remarkable fidelity after each cell division, is somehow dependent on the concerted action of the ParB and SMC proteins. Although ParB-deficient cells exhibit enhanced cell-to-cell phenotypic variation, including a minority of cells with nearly wild-type behavior, in the majority of $\Delta parB$ cells, *ori* localization (and, consequently, replisome assembly) is biased toward cell pole or quarter-cell positions (Fig. 3A, initiation at two *ori*). Although *parB* is not essential for viability, the large fraction of anucleate cells formed by the $\Delta parB$ strain indicates that ParB is required for the high fidelity of *ori* positioning and segregation exhibited by wild-type cells.

Although the mechanisms responsible for *ori* localization and movement in *M. smegmatis* are unknown, ParB might play a role in the observed segregation of the *ori*-proximal region immediately after its duplication and subsequent maintenance of the duplicated *ori*-proximal regions around the quarter-cell positions, which will become the midcell positions in the daughter cells following cell division. Increased IFD between the *ori* and *attB* loci, as observed in ParB-deficient cells, suggests that in the absence of ParB-mediated constraints on *ori* positioning, the chromosome might adopt a longitudinal organization similar to that in *C. crescentus* (40, 41) or MukB-deficient *E. coli* (42). ParA has also been implicated in these processes, as indicated by the mislocalization of ParB in a ParA-deficient strain of *M. smegmatis* (22). However, the mechanism of action of the ParABS system in *M. smegmatis* is unclear. In *C. crescentus*, it has been proposed that ParA filaments extend across the cell, attach to the chromosome via interactions with the ParB-*parS* complex, and pull the *ori*-proximal region of the chromosome toward the new cell pole by depolymerization-mediated contraction, analogous to the role of the mitotic spindle in segregating eukaryotic chromosomes (43). However, an alternative “DNA relay” model has recently been proposed, in which an asymmetric spatial gradient of ParA molecules (with the highest concentration at the new cell pole) rectifies the otherwise symmetric elastic dynamics of the chromosome through transient interactions with the ParB-*parS* complex (18).

In contrast to the SMC protein of *B. subtilis* (34, 35), that of *M. smegmatis* seems to be involved in neither localization nor segregation of *ori*, although the aberrant subcellular positioning of the *attB* locus observed in SMC-deficient cells suggests that SMC does play a role in chromosome organization. In the absence of SMC, DNA synthesis continues and replicated foci are resolved and segregated, but in many cells, the subcellular chromosome organization differs from that in the wild type. We hypothesize that the SMC foci observed in cells expressing an SMC-GFP fusion protein serve as centers of SMC activity, as proposed in *B. subtilis* (24). ParB seems to play a role in the focal recruitment of SMC to the chromosome, as SMC foci are reduced or eliminated in ParB-deficient cells. In wild-type cells, SMC foci are often found in close proximity to but usually do not colocalize with ParB, suggesting that ParB might recruit SMC to the *ori*-proximal region and from there, SMC might diffuse along the chromosome, as proposed in *B. subtilis* and *S. pneumoniae* (23, 25). Moreover, it has been suggested that SMC might dictate the overall organization of the bacterial chromosome by condensing lengthwise along the two arms of the chromosome, thereby generating a stiffer structure into which newly synthesized DNA is folded (15, 44). Consistent with this idea, we found that the mirror-symmetrical localization of *attB* loci observed in wild-type sibling cells is partially lost in SMC-deficient cells, in which an asymmetric distribution of *attB* loci

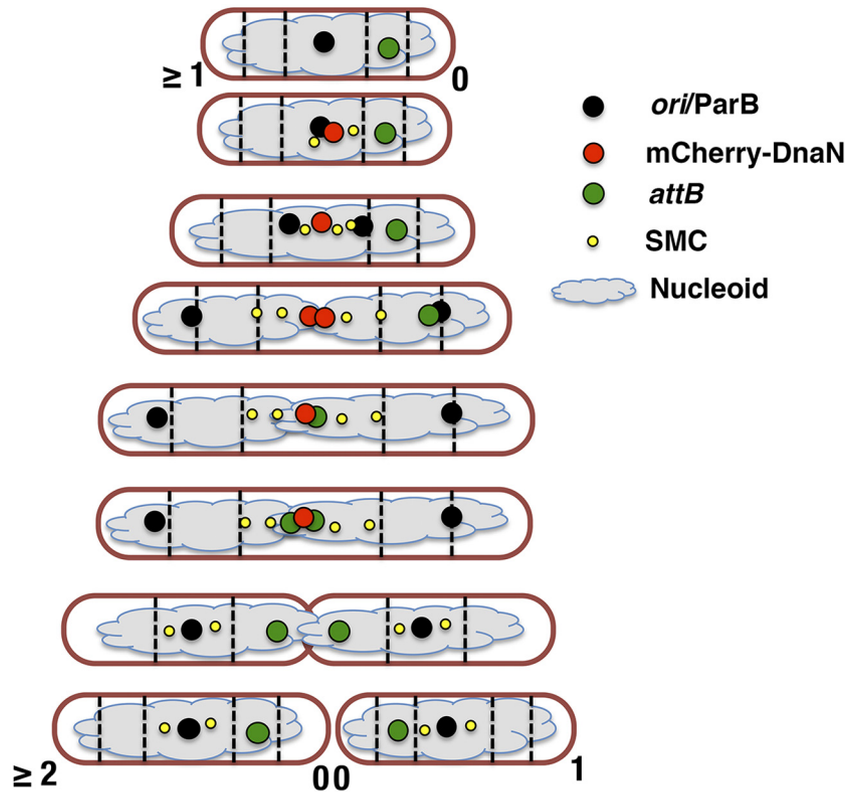


FIG 7 Schematic of chromosome organization and replisome dynamics in *M. smegmatis*.

relative to midcell is more frequently observed. We hypothesize that in wild-type sibling cells, the chromosomes adopt an R-L-R configuration, which may switch to an R-L-R-L configuration in the absence of SMC.

In a small fraction of wild-type cells the replisome undergoes splitting and the split replisomes localize to opposite cell halves for the remainder of the replication cycle. The frequency of cells containing split replisomes is strongly increased in ParB- and SMC-deficient strains, for two reasons: replisome splitting in a large fraction of $\Delta parB$ and Δsmc cells and transient polyploidy in a small fraction of $\Delta parB$ cells because of chromosome missegregation events during cell division. Both phenotypes might be attributable to impaired subcellular chromosome organization in the absence of ParB or SMC. Consistent with this interpretation, we found that the replisome can assemble at almost any subcellular position in ParB-deficient cells, whereas in wild-type and SMC-deficient cells, replisome assembly is confined to the midcell region. In ParB- and SMC-deficient cells, the split replisomes usually move toward the edge of the nucleoid, similar to those in *E. coli* (5), although in a small fraction of cells, the replisomes move longitudinally through the cell, similar to those in *C. crescentus* (6). In these species the pattern of replisome migration matches the orientation of the chromosome: L-ori-R for *E. coli* (39) and ori(out)-ter(in) for *C. crescentus* (40, 41). It is possible that the altered replisome dynamics observed in $\Delta parB$ and Δsmc cells might be linked to impaired subcellular chromosome organization in these mutants. Thus, chromosomal regions that are normally proximal to each other in wild-type cells might be spatially separated in ParB- and SMC-deficient cells, resulting in sep-

aration of replisomes engaged in replicating the left and right chromosome arms. Consistent with this interpretation, in ParB- and SMC-deficient cells, the split replisomes seem to track along the DNA following the subcellular disposition of the nucleoid.

Time-lapse analysis of ParB- and SMC-deficient strains sometimes revealed cells containing three or even four DnaN foci, which arise by transient splitting of the two main replisomes into smaller and fainter foci. We hypothesize that these additional foci might be engaged in the synthesis of Okazaki fragments on the lagging strand, consistent with the observation that these small and dim DnaN foci often colocalize with SSB foci. It is possible that the separation of leading- and lagging-strand replisomes might become transiently visible because of loss of local chromosome compaction in ParB- and SMC-deficient cells. An alternative explanation is suggested by the recent demonstration that the DnaN clamp zone provides a platform for spatiotemporal coupling of DNA replication and mismatch detection in *B. subtilis* (45). However, the latter interpretation is unlikely to be correct because SMC deficiency does not result in a mutator phenotype and SMC does not seem to play an important role in DNA repair (27).

Although localization of replisomes around midcell is apparently not essential for DNA replication and cell cycle progression, we hypothesize that replisome localization might play a role in positioning of the cell division septum. This idea arises from the observation that the division septum is often mislocalized in ParB- and SMC-deficient cells; this defect is particularly pronounced in ParB-deficient cells, which generate small anucleate progeny because of mispositioning of the division septum. Time-lapse mi-

scopy of dual-reporter strains expressing fluorescent markers of the replisome (mCherry-DnaN) and the division septum (Wag31-GFP) also supports the idea that the replisome might play an important role in division site selection (our unpublished data). Linking division site selection to the spatiotemporal dynamics of chromosome replication and segregation might serve as an alternative positioning mechanism in mycobacteria, which do not encode homologs of the minicell or nucleoid occlusion proteins responsible for division site selection in other bacteria (46).

MATERIALS AND METHODS

Bacterial strains and culture conditions. *M. smegmatis* mc²155 (wild type) and derivative strains were grown in Middlebrook 7H9 medium (Difco) supplemented with 0.5% albumin, 0.2% glucose, 0.085% NaCl, 0.5% glycerol, and 0.05% Tween 80. Cultures were grown at 37°C with aeration to mid-log phase (optical density at 600 nm [OD₆₀₀] of ~0.5). Aliquots were stored in 15% glycerol at -80°C and thawed at room temperature before use; individual aliquots were used once and discarded.

Oligonucleotides, plasmids, and bacterial strains. The oligonucleotides, plasmids, and bacterial strains used in this study are listed in Table S1.1-S1.3 in the supplemental material. Details of plasmid and strain construction are provided in Text S1.

Single-cell microscopy. For snapshot microscopy, bacteria were grown to mid-log phase (OD₆₀₀ of ~0.5) in 7H9 medium, collected by centrifugation (2,400 × g, 5 min), concentrated 10-fold in fresh 7H9 medium (37°C), and passed through a 5-μm-pore-size polyvinylidene difluoride syringe filter (Millipore) to remove clumps. The deplumped bacteria were transferred to a glass slide with a 1% agarose pad soaked in LB medium and covered with a coverslip. Nucleoid staining was done by incubating cells for 5 min with SYTO 17 red or SYTO 13 green (Invitrogen) diluted 1:2,000 prior to microscopy at room temperature.

For time-lapse microscopy, bacteria were prepared as described above, except that the deplumped cell suspension was spread on a glass coverslip, covered with a semipermeable membrane and cultured in a custom-made microfluidic device with a continuous flow of 7H9 medium at 37°C (flow rate, 25 μl/min), as previously described (47). Nucleoid staining was done by adding SYTO green (diluted 1:2,000) to the flow medium.

Bacteria were imaged with a DeltaVision personalDV microscope (Applied Precision) equipped with a 100× oil immersion objective and an environmental chamber maintained at 37°C. Images were recorded on phase-contrast and fluorescence channels (490/20-nm excitation filter and 528/38-nm emission filter for GFP, 575/25-nm excitation filter and 632/60-nm emission filter for mCherry) with a CoolSnap HQ2 camera. Images were processed with Softworx software (Applied Precision). Cell length was measured as the sum of short linear segments tracking along the center line of an individual cells in order to accommodate irregularities in cell shape. Interdivision time (*I_t*) was defined as the time interval between the first appearance of septal Wag31-GFP (birth) and the next appearance of septal Wag31-GFP (division), as previously described (33).

Immunoblotting. Bacteria were grown to mid-log phase (OD₆₀₀ of ~0.5), collected by centrifugation (2,400 × g, 5 min), and resuspended in lysis buffer (50 mM Tris-Cl, pH 7.5) containing protease inhibitors (Roche). Proteins were extracted in a FastPrep-24 instrument (MP Bio-medicals) with 0.1-mm zirconia beads, separated on NuPAGE Novex 3 to 5% Tris-acetate gels (Invitrogen), and electrotransferred to a nitrocellulose membrane. A mouse anti-GFP monoclonal antibody was used as the primary antibody (1:1,000). A rabbit anti-mouse antibody conjugated to horseradish peroxidase (Dako) was used as the secondary antibody (1:1,000).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <http://mbio.asm.org/lookup/suppl/doi:10.1128/mBio.01999-14/-/DCSupplemental>.

Text S1, DOCX file, 0.1 MB.

Table S1, DOCX file, 0.2 MB.

Figure S1, TIF file, 1.5 MB.

Figure S2, TIF file, 2.7 MB.

Figure S3, TIF file, 2.4 MB.

Figure S4, TIF file, 2.7 MB.

Figure S5, TIF file, 2.7 MB.

Figure S6, TIF file, 0.3 MB.

Movie S1, MOV file, 2.6 MB.

Movie S2, MOV file, 3.2 MB.

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