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*CORRESPONDENCE Cees Dekker ⊠ c.dekker@tudelft.nl

[†]These authors have contributed equally to this work

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MukBEF-dependent chromosomal organization in widened *Escherichia coli*

Aleksandre Japaridze[†], Raman van Wee[†], Christos Gogou, Jacob W. J. Kerssemakers, Daan F. van den Berg and Cees Dekker*

Department of Bionanoscience, Kavli Institute of Nanoscience Delft, Delft University of Technology, Delft, Netherlands

The bacterial chromosome is spatially organized through protein-mediated compaction, supercoiling, and cell-boundary confinement. Structural Maintenance of Chromosomes (SMC) complexes are a major class of chromosomeorganizing proteins present throughout all domains of life. Here, we study the role of the Escherichia coli SMC complex MukBEF in chromosome architecture and segregation. Using quantitative live-cell imaging of shape-manipulated cells, we show that MukBEF is crucial to preserve the toroidal topology of the Escherichia coli chromosome and that it is non-uniformly distributed along the chromosome: it prefers locations toward the origin and away from the terminus of replication, and it is unevenly distributed over the origin of replication along the two chromosome arms. Using an ATP hydrolysis-deficient MukB mutant, we confirm that MukBEF translocation along the chromosome is ATP-dependent, in contrast to its loading onto DNA. MukBEF and MatP are furthermore found to be essential for sister chromosome decatenation. We propose a model that explains how MukBEF, MatP, and their interacting partners organize the chromosome and contribute to sister segregation. The combination of bacterial cell-shape modification and quantitative fluorescence microscopy paves way to investigating chromosome-organization factors in vivo.

KEYWORDS

E. coli, MukBEF, SMC, chromosome segregation, nucleoid architecture, MatP

Introduction

The intricate organization of genetic material in chromosomes remains incompletely understood, even in thoroughly studied bacteria such as *Escherichia coli* (*E. coli*). In order to fit into the volume of a single cell, the nucleoid needs to be strongly compacted (Travers and Muskhelishvili, 2005), while preserving a complex spatial and dynamic organization that facilitates vital cellular processes. In *E. coli*, compaction is achieved by a combined interplay of DNA supercoiling (Blot et al., 2006), nucleoid associated proteins (NAPs) (Luijsterburg et al., 2006), and a Structural Maintenance of Chromosomes (SMC) complex called MukBEF (Niki et al., 1991; Thanbichler et al., 2005; Petrushenko et al., 2006; Reyes-Lamothe et al., 2008; Nolivos and Sherratt, 2014; Rybenkov et al., 2014). MukBEF is a pentamer consisting of double copies of MukB and MukE subunits and a single kleisin unit called MukF (Rybenkov et al., 2014). DNA can be bound and reshaped *in vitro* by the hinge-like configuration of the MukB subunits in an ATP-dependent manner (Chen et al., 2008). Recent studies reported that a 6-fold increase in the

number of MukBEF copies led to the formation of a ring-like structure of SMCs along the toroidal nucleoid (Mäkelä and Sherratt, 2020; Mäkelä et al., 2021). This ring was hypothesized to function as a chromosomal backbone from which peripheral DNA loops protrude.

MukBEF also mediates chromosomal interactions with other proteins that organize and disentangle sister chromosomes during replication and segregation (Nolivos et al., 2016). One such protein is the so-called Macrodomain ter Protein (MatP). MatP's organizational role is commonly associated with its active displacement of MukBEF from the ter region (Nolivos et al., 2016). MatP binds matS sites near the terminus of replication and localizes the ter macrodomain to the midcell through a direct interaction with the divisome (Mercier et al., 2008; Espéli et al., 2012). There is growing in vivo evidence that in the absence of MatP, MukBEF is unable to be displaced from the ter region which then results in severe condensation of the ter region (Nolivos et al., 2016; Lioy et al., 2018; Mäkelä and Sherratt, 2020). Recently, the molecular structure of the MukBEF-MatP-matS nucleoprotein complex was resolved using cryo-EM, revealing how the subunits of MukBEF and MatP directly interact (Bürmann et al., 2021). matSbound MatP was found to sit at the center of the MukBEF ring, potentially blocking MukBEF translocation in the ter domain and promoting ATP-dependent un-loading of the SMC in vivo. Furthermore, deletion of MukBEF was shown to result in anucleation and defects in chromosome segregation(Niki et al., 1991; Yamazoe et al., 1999; Danilova et al., 2007; Nicolas et al., 2014), whereas deletion of matP led to premature segregation of sister foci in the ter macrodomain and their mis-localization relative to the divisome before cell division (Mercier et al., 2008; Nolivos et al., 2016).

Another important direct interaction partner of MukBEF (and possibly an indirect one for MatP as well) is the topoisomerase IV (Kumar et al., 2017, 2022; Fisher et al., 2021). Topo IV influences the linking number of the nucleoid primarily in the ori region, where MukBEF is predominantly localized. Furthermore, it has been found to mediate the timely segregation throughout replication as well as the decatenation of sister chromosomes after replication (Goto and Wang, 1982; Peng and Marians, 1993; Seol et al., 2013). Although these findings form a foundation for the understanding of chromosomal organization, the high degree of nucleoid compaction in combination with simultaneous ongoing replication cycles have so far impeded direct *in vivo* visualization of their actions.

To map spatiotemporal localizations and investigate the interactions of MukBEF and MatP with the chromosome, we employed a method to synchronize chromosome replication in a population of E. coli cells [by using a temperature-sensitive dnaC allele (Saifi and Ferat, 2012)] and simultaneously increased their size through cell-shape manipulation. For the latter, treatment of cells with low doses of the A22 inhibited the polymerization of the MreB filaments, thereby disrupting the typical rod shape of E. coli (Varma and Young, 2009). These cells gradually expanded in size and typically reached at least 2-fold larger width and length. Concomitantly, the spatial constraint that the cell wall imposed on the nucleoid was thus reduced. In previous studies, we showed how the chromosome in such expanded shapes exhibited a toroidal topology and remained physiologically active in the cell, (Karczmarek et al., 2007), preserving its capability to replicate and segregate its chromosomes and re-adopting a rod shape upon A22 depletion, suggesting that treatment with A22 did not impact cell viability (Wu et al., 2019a; Japaridze et al., 2020). Interestingly, similar widened cell wall-deficient bacteria (Mickiewicz et al., 2019) were also observed in patients with recurring infections, suggesting that bacterial cells are capable of naturally reshaping their size and cell wall composition.

Here, we use quantitative fluorescence imaging to study the distribution of MukBEF along the chromosome and we characterize structural changes that result from the mutation or deletion of MukBEF subunits in living cells. We reaffirm that MukBEF positions along the chromosome with a strong ori-proximal and ter-distal spatial bias in the presence of MatP near the terminus of replication. The preferential localization of MukBEF away from ter is strongly dependent on its ability to hydrolyze ATP, as is its ability to compact the nucleoid. Additionally, MukB is found to spread asymmetrically over the origin of replication along the chromosome arms. We corroborate that the deletion of *matP* does not alter the capacity of MukBEF to bind and compact the nucleoid, but that its localization along the genome is directed by MatP. Upon deletion of matP, MukBEF displays a 3-fold increased presence near the ter region which leads to a local compaction of this domain, and which results in severe segregation defects. Deletion of either MukBEF or MatP was found to impair sister decatenation, resulting in the formation of dimer chromosomes. Our quantitative fluorescence analysis in combination with increased spatial resolution in live shape-modified cells offers new means for investigating chromosome organization in vivo.

Results

MukBEF spreads non-uniformly along the chromosome

Throughout this study we performed simultaneous four-color imaging of the chromosome (DAPI or HU-mYpet) and MukB (mYpet) together with the origin and the terminus of replication [via Fluorescence Repressor Operator Systems in living cells (Wang et al., 2006; Figure 1A)]. Rod-shaped cells grown in minimal media typically display one or two origins of replication and a single terminus, indicating that cells are in the process of replication (Khan et al., 2016; Figure 1B). In order to circumvent the optical limitations due to high degree of chromosome compaction and cell-to-cell variability due to ongoing replication cycles, we performed experiments with A22-widened temperature-sensitive (dnaCts) E. coli cells (Saifi and Ferat, 2012). Cells were first synchronized by growing above permissive temperature [and hence cells maintained only a single chromosome (Saifi and Ferat, 2012; Wu et al., 2019b)] in the presence of low doses of A22 to grow larger in size before imaging. While the cell width is expanded laterally two-fold within the plane, the cell height is still limited to ~1 micron out of the plane, due to the agarose pad used for imaging (Wu et al., 2019a; Japaridze et al., 2020).

In these widened cells, the chromosomes organize into a toroidal configuration, with the origin and terminus of replication positioned at opposite halves of the chromosome ring (Wu et al., 2019a; Japaridze et al., 2020). Surprisingly, in the widened cells the MukBEF complexes did not form a single tight cluster as they do in rod-shaped cells (Figure 1C). Rather, the MukBEF signal distributed over the toroidal chromosome to adopt a significantly extended cluster with a Ferret diameter of $0.6 \pm 0.3 \,\mu\text{m}$ (mean \pm SD, N = 235 cells) compared to a diffraction-limited diameter of $0.4 \pm 0.1 \,\mu\text{m}$



Quantitative localization of MukB complexes along the chromosome. (A) Schematic representation of a circular *Escherichia coli* genome with FROS arrays at *ori1* and *ter3* location and MukB-mYpet labeling. (B) Representative image of rod-shaped and (C) widened *E. coli* cells in phase contrast and the fluorescence channels of *ori1, ter3* and MukB. Cell outline is indicated in white. (D) Typical example of the thresholding process to calculate fluorescence signal overlap. (E) Percentage of MukB intensity that overlaps with the DNA mask for wildtype (N=260 cells), *AmatP* (N=121 cells), and MukB^{EO}EF mutants (N=131 cells). (F) Signal overlap between MukB and *ori1* and MukB and *ter3* foci in wildtype (N=260 cells), *AmatP* cells (N=121 cells) and MukB^{EO}EF cells (N=131 cells). Error bars report on the standard deviation. (G) Schematic representation of MukBEF positioning along the circular *E. coli* chromosome and its regulation by MatP. Statistical significance was determined by performing a single factor ANOVA test. The following conventions are used: ns: 0.05 < p, *: 0.01 . **: <math>0.001 . We report a significant difference in results if <math>p < 0.05.

(mean \pm SD, N = 118 cells) for the clusters in rod-shaped cells (Supplementary Figure S1). In the widened cells, the signal spread along the left and right arms of the chromosome and the number of MukBEF clusters was typically one or two, as was the number of ori foci in a cell (Figure 1D). The total MukB signal was slightly higher in the widened cells (mean value 30% higher relative to rod-shaped cells, Supplementary Figure S1). Next, we measured the ability of MukBEF to bind the nucleoid. The distribution of MukB across the nucleoid reflects not only the nucleoid-binding ability of MukBEF but also nucleoid condensation and association between MukBEF and other interacting partners. We determined the percentage of the nucleoid that was covered by MukBEF clusters (Supplementary Figure S2). In wildtype cells, we found that 8.6 ± 0.5% (mean ± SD, N= 260 cells) of the total DNA

material was colocalized with the MukBEF (Figure 1E). We compared this with two mutants: $\Delta matP$ cells, in which the MatP protein was deleted, and MukB^{EQ}EF cells, whose MukB subunits were impaired in hydrolyzing ATP (Woo et al., 2009). For the $\Delta matP$ and MukB^{EQ}EF mutants, this overlap increased significantly to $10.8 \pm 0.8\%$ (mean \pm SD, N = 121 cells) and $10.8 \pm 0.9\%$ (mean \pm SD, N = 131 cells), respectively. We hypothesize that this enhanced coverage of the DNA by MukBEF is because MukBEF also partly occupies the ter region in the mutants, indicating that both MatP and MukB ATPase activity are required for MukBEF displacement.

Furthermore, we quantitatively evaluated MukBEF positioning along the chromosome, by investigating the colocalization of MukBEF with the origin and terminus of replication as both positions were marked with loci (Figure 1F). In $93 \pm 2\%$ of all cells (mean \pm SD, N = 260 cells), MukB and ori signal overlapped, whereas we only observed a $3.8 \pm 2.0\%$ (mean \pm SD, N = 260 cells), overlap between MukB and the ter signal (Figure 1F). Since MukB is spread out in clusters, MukB overlap with ori was not mutually exclusive with overlap with ter. In $\Delta matP$ cells, we found that MukBEF colocalization with the ori was preserved, but to a clearly lesser extent than in wildtype cells in only 70 ± 5% (mean ± SD, N = 121 cells) of $\Delta matP$ cells, the MukBEF signal was overlapping with ori (Figure 1F). This reduction in colocalization could be explained by a redistribution of MukBEF toward the MatP-depleted ter region. Indeed, we observed a 3-fold increase in colocalization of MukB with the ter in $\Delta matP$ cells to $11 \pm 1\%$ (4% in wildtype cells). Furthermore, the fraction of cells in which MukB did not overlap with either focus was 10-fold higher than in wildtype (20% versus 2%). MatP thus is crucial to displace MukBEF from the ter macrodomain. In wildtype cells, MatP is observed as spots flanking the stretched-out ter region (Supplementary Figure S3). Deletion of *matP* leads to compaction of the ter region, and subsequent reorganization of the chromosome with a lower DNA cluster number. Since the total amount of DNA in the chromosome is constant, a lower cluster number implies a stronger compaction of the chromosome (Supplementary Figure S4). The MukB^{EQ} distribution along the nucleoid of MukB^{EQ}EF mutants was similar to the $\Delta matP$ mutants, as MukB^{EQ} overlapped with the ori focus in 72% of MukB^{EQ}EF cells (N=131 cells), 20% of the cells showed an overlap with the ter focus, and the remaining 8% showed no overlap with either. The 5-fold increase (20% versus 4%) in colocalization of the MukBEQ signal with the ter region for the MukB^{EQ}EF cells relative to the wildtype is remarkable as it shows that MukBEF can bind to the ter region but apparently is unable to get displaced if its capability to hydrolyze ATP is impaired, in agreement with earlier work (Nolivos et al., 2016).

MukBEF is distributed asymmetrically over the left and right chromosome arms

Visualizing the toroidal structure of the chromosome and separately observing the two arms of the chromosome enabled us to zoom in further on the spreading of the MukB in wildtype cells, and $\Delta matP$ and MukB^{EQ}EF mutants (Figures 2A,B). We first measured the relative local spread of the MukB near the origin of replication along the two arms of the chromosome. As the origin of replication lies between both arms, a central axis can be drawn from the chromosome's center of mass to ori to divide the chromosome in two arms. Then we checked if the distribution over both arms was symmetric, by defining the local asymmetricity as the absolute difference between the fluorescent signal along the two chromosome halves divided by their sum (see Materials and methods). If the fluorescent intensity was equally spread over the two arms, the asymmetricity equals 0, while it equals 1 if all signal is on one of the chromosome arms (Figures 2C,D). We first determined the median asymmetricity of DNA in wildtype cells as an intrinsic control to account for errors introduced during data-acquisition (e.g., caused by the finite image resolution and the dynamic nature of the chromosome). In wildtype cells the median asymmetricity of the local DNA signal was 0.27 (Figures 2C,E), while for the MukB signal it was significantly higher at 0.33 (N=284) (Figures 2D,E). Thus, it was clear that locally one of the arms (irrespective of the DNA content) within one chromosome had significantly more MukB signal compared to the other. To identify which arm that was, we performed ChiP-Seq data analysis and checked the local spreading of MukBEF binding sites near the oriC (Supplementary Figure S5), based on available data from Nolivos et al. (2016). In the 0.5 Mbp vicinity to oriC, there were 20% relatively more upstream than downstream peaks, indicating that MukB predominantly occupies the right arm. Interestingly, when we measured the asymmetricity of the MukB in the $\Delta matP$ cells, the spreading was even more asymmetric (median asymmetricity for DNA was 0.32, but for MukB 0.49, N=108) (Figure 2E). However, in the MukB^{EQ}EF cells the asymmetricity of the DNA and MukB spreading along the chromosome was not significantly different (0.25 and 0.26, respectively N = 124).

MatP and ATP hydrolysis by MukBEF are needed for chromosome compaction and organization

To quantify chromosome compaction by MukBEF, we analyzed whether the DNA regions overlapping with MukBEF were associated with an increased DNA compaction compared to the rest of the chromosome. Interestingly, in wildtype cells the DNA density within the MukBEF regions was indeed higher, by a factor of 1.8 ± 0.05 (mean \pm sem, N = 260 cells), compared to the DNA signal elsewhere along the chromosome (Figure 3A), highlighting MukBEF's ability to locally compact the nucleoid. Despite the increased prevalence of MukBEF at the ter region in *AmatP* cells, the magnitude of local DNA compaction of MukBEF-occupied regions was not affected by the deletion of *matP*, as it was very similar to the wildtype [Figure 3A; 1.75 ± 0.05 times higher (mean \pm sem, N = 121 cells)]. Strikingly, the compaction of the DNA that overlapped with MukB in MukBEQEF mutants was not significantly increased and only slightly higher compared to any other chromosomal region where MukBEF was not present [Figure 3A; 1.2±0.03 times higher than elsewhere along the chromosome (mean \pm sem, N = 131 cells)]. This suggests that ATP hydrolysis is a requirement for compaction by MukBEF.

We next investigated the effects of altered MukBEF localization and activity on chromosome shape parameters. In both $\Delta matP$ cells and MukB^{EQ}EF mutants, we found that the chromosome contour length was significantly shorter than in wildtype (Figure 3B). While wildtype cells had an average length of $4.5 \pm 0.1 \,\mu\text{m}$ (mean \pm sem, N=222 cells), it was reduced to $3.5 \pm 0.1 \,\mu\text{m}$ (mean \pm sem, N=148cells) and $3.8 \pm 0.1 \,\mu\text{m}$ (mean \pm sem, N=226 cells) in $\Delta matP$ cells and MukB^{EQ}EF mutants, respectively. For the chromosome width,



FIGURE 2

Asymmetric spreading of MukB complexes along the right and left arms of the chromosome. Representative images of (A) $\Delta matP$, (B) MukB^{EO}EF *E. coli* cells in phase contrast and the fluorescence channels of *ori1* (red), *ter3* (cyan), MukB (yellow), DNA stained with DAPI (blue) and their overlay. Cell outline is indicated in white. Measuring the local spreading of (C) DNA, and (D) MukB near the origin of replication. Left: the central ridge of the fluorescent signal is determined. Middle: the central axis (orange line) is detected along the origin (red spot) to the chromosome center of mass (white spot). Only for the circular area (red circle) around the origin, asymmetricity is determined. Right: a symmetry value is defined as the difference in signal from anticlockwise area (blue) and clockwise area (red) (relative to the line connecting ori and the center of mass) divided by their sum signal. Orange and light blue colors denote high and low signal intensity, respectively. (E) Asymmetricity distributions for DNA and MukB channels as defined in panels c and d for wildtype (N=284 cells), *AmatP* (N=108 cells), and MukB^{EO}EF mutants (N=124 cells). Horizontal black lines represent median values. Statistical significance was determined by performing a single factor ANOVA test. The following conventions are used: ns: 0.05 < p, *: 0.01 , ****: <math>p < 0.0001. We report a significant difference in results if p < 0.05.

characterized by the average full-width-at-half-maximum along the chromosome (FWHM), a different trend was observed (Figure 3C). We found that chromosomes of wildtype and $\Delta matP$ cells had similar width of $0.44 \pm 0.08 \,\mu\text{m}$ (mean $\pm \,\text{sem}$, $N = 222 \,\text{cells}$) and $0.45 \pm 0.08 \,\mu\text{m}$ (mean $\pm \,\text{sem}$, $N = 148 \,\text{cells}$) respectively, while MukB^{EQ}EF mutants formed a chromosome with a slightly reduced width of $0.41 \pm 0.07 \,\mu\text{m}$ (mean $\pm \,\text{sem}$, $N = 226 \,\text{cells}$).

Since the DNA compaction in wildtype and $\Delta matP$ mutant cells was similar, we conclude that MukBEF can bind and compact the chromosome independently of MatP. Aside from impeded MukBEF displacement over the DNA, the MukB^{EQ}EF mutant also showed a clear impairment in its ability to compact the chromosome (Figure 3A). Hence our findings suggest that MukBEF needs to hydrolyze ATP in order to compact the DNA and redistribute its position along the nucleoid (Figure 3D), as similarly shown by others (Woo et al., 2009; Badrinarayanan et al., 2012; Nolivos et al., 2016). The altered chromosome length and width in the mutants show that MukBEF is an important factor in the global chromosome organization where both ATP hydrolysis and MukBEF's interaction with MatP are required for its faithful functioning.

mukB or *matP* deletion leads to chromosome decatenation defects and dimerization

Finally, to probe the roles of MukBEF and MatP in *E. coli* chromosome segregation, we visualized the chromosome structure after replication initiation in strains with either a *mukB* or *matP* deletion (Supplementary Figure S6). However, since the *mukB* deleted



cells did not grow at temperatures above 24°C (Niki et al., 1991), cells could not be synchronized for replication initiation. As a result, these cells typically contained more than one chromosome at any given time. In 74% of the $\Delta mukB$ cells (N=172 cells) there were 2 or more complete chromosomes, as recognized by a multitude of ori and ter foci, while this phenotype was not observed in wildtype cells (Wang and Sherratt, 2010; Japaridze et al., 2020). As shown in Figure 4A, chromosome dimers that were shaped like a figure-eight shape were observed in 91% of cells (N=172 cells) with two complete chromosomes (i.e., cells with 2 ori and 2 ter foci), indicating that the decatenation of sister chromosomes was impaired in the absence of MukB. Interestingly, in the remaining 9% of polyploid cells, the chromosome dimers were organized in a toroidal configuration (Figure 4B, more examples in Supplementary Figure S7).

We related the number of ori foci to the total fluorescence intensity of the corresponding chromosome (normalized to cells with a single ori) and found that they scaled almost linearly (Figure 4C), suggesting that the number of ori sites (in combination with ter foci) indeed indicated the number of fully replicated chromosomes in a cell.

Dimer chromosomes had a normalized fluorescence intensity of 2.2 ± 0.7 (mean \pm SD, N=18 cells), while chromosomes with four oris had an intensity of 3.7 ± 1.2 (mean \pm SD, N=14 cells), compared to single chromosomes [1.0 ± 0.2 (mean \pm SD, N=14 cells)].

A closer inspection of the ori and ter foci in $\Delta mukB$ cells revealed a striking difference compared to wildtype. Wildtype cells grown in minimal medium typically exhibited 2 ori/ter ratios: either we observed a single ori and a single ter (1:1) in cells with a single non-replicating chromosome, or we observed 2 oris and a single ter (i.e., a ratio of 2:1) in cells that were replicating. In $\Delta mukB$ cells, however, only 17% of the cells showed either of these counts, whereas we found 15 other combinations of ori and ter counts (Supplementary Figure S7). We grouped these cells by the number of ter foci and plotted the ori counts in a histogram and found an increase in the average number of oris as the number of ters increases; from 2.4 ± 1.1 oris for cells with 1 ter, to 3.4 ± 1.1 for 2 ters and 4.3 ± 1.0 for 3 ters (all mean ± sd; Figure 4D). The occasional observation of single cells carrying chromosomes with three oris suggests that initiation of daughter chromosomes can occur asynchronously, while Japaridze et al.



overlay for a typical $\Delta mukB$ cell widened with A22 are shown. The cell outline is shown in white and the impaired decatenation is schematically depicted. (B) Idem as A, now for the donut-shaped chromosomes. (C) Normalized sum chromosome signal (integrated HU-mYpet signal) in $\Delta mukB$ cells. Horizontal black lines show the median values (1,02; 2,10; 3,82). (D) Distribution of the observed ori (x-axis) & ter (color) counts in $\Delta mukB$ mutants (N=167 cells). Although observed, cells with four ter foci were excluded in the plot due to their low statistics (N=5 cells). The distributions are fitted with a Gaussian function, whose mean is indicated by the dotted lines. (E) Overlay of the fluorescence signals for DNA, ori and ter for two representative $\Delta matP$ cells. The cell outline is shown in white, and the toroidal configuration is schematically depicted. (F) Proposed model for the interaction between chromosomal organizers. Our data (green box) show that MatP inhibits chromosome decatenation, while novobiocin inhibits the latter, leading to segregation defects. These notions fit within the larger picture in which XerCD and FtsK are also interacting with TopolV and MatP, respectively, as well as with each other. Double headed arrows indicate mutual regulation, whole the dotted arrows indicate putative regulatory levels.

four oris indicate that the next round of replication was initiated before proper sister chromosome decatenation (Khan et al., 2016).

Deletion of *matP* also causes a strong phenotypic change in chromosome segregation. In $\Delta matP$ cells, the ter region was not properly positioned relative to the mid cell (geometric midcell of the fitted ellipsoid) throughout the segregation, as seen in wildtype (Japaridze et al., 2020). Cells with 2 ter foci were often observed in a stage of early septation, even though the circular sister chromosomes

were still topologically catenated (Movie 1). Surprisingly, we could see the formation of toroidal dimer chromosomes in these cells as well. In 10.5% of the replicating $\Delta matP$ cells, the two sister chromosomes would form one larger ring-like chromosome. Two pairs of ori and ter foci were present in these dimer chromosomes, which were evenly spaced and with a distinct order of ori-ter-ori-ter along the ring contour of the chromosome (Figure 4E, more examples in Supplementary Figure S7). These cells displayed a similar phenotype as wildtype cells treated with the DNA gyrase inhibitor Novobiocin (Maxwell, 1993; Supplementary Figure S8), indicating that deletion of either *mukB* or *matP* impairs chromosome decatenation by Topo IV.

Discussion

In this study, we investigated the role of MukBEF and MatP in chromosome organization and segregation in A22-widened temperature-sensitive (*dnaCts*) *E. coli* cells. Because of the cell widening (typically twice larger in width and length compared to rod-shaped cells), the toroid-shape chromosome could be resolved and the positioning of the MukBEF along the nucleoid could be measured with increased spatial resolution without the need to overexpress MukBEF or cell fixation. Additionally, chromosome replication can be synchronized by first culturing the cells above the permissive temperature (40° C), and subsequently re-initiating replication by transferring them to room temperature (Wu et al., 2019a; Japaridze et al., 2020).

Through quantitative imaging, we corroborated the earlier findings in rod-shaped cells that the MukBEF SMC in *E. coli* localizes near the origins of replication and away from the ter region (Nolivos et al., 2016). In the absence of either ATP-hydrolysis by MukB (MukB^{EQ} mutant) or MatP, we showed that MukBEF has a 3 to 5-fold higher association with ter, indicating that MukBEF normally binds to this macrodomain prior to being actively expelled toward ori by MatP (Figures 1E,G; Supplementary Figure S5). Combined, the data elucidate that MatP-mediated MukBEF expulsion needs MukBEF's ATP-dependent dissociation from the chromosome rather than preventing DNA binding altogether. This finding is in good agreement with earlier works, which showed that MukBEF is displaced from the ter region by MatP (Mercier et al., 2008).

Further, we observed an asymmetry of MukBEF occupancy on the left versus right arm of the chromosome (Figures 2D,E), which is consistent with similar reports using 3C/HiC (Lieberman-Aiden et al., 2009) or ChiP-qPCR (Kim and Dekker, 2018), in other organisms such as Corynebacterium glutamicum (Wang et al., 2015; Tran et al., 2017) and Bacillus subtilis (Antar et al., 2021). There, however, SMCs were loaded at specific parS genomic sequences near the origin of replication and spread preferentially toward the left arm of the chromosome. The observation that the asymmetric spreading is lost in the MukBEQEF cells points to the ATP-dependence of MukB spreading along the chromosome. Similar to reports for other bacteria (Böhm et al., 2020; Antar et al., 2021), we observed a preferential spreading toward the right arm of the chromosome (Supplementary Figure S5; Nolivos et al., 2016). Considering that the rDNA genes located on this arm have been shown to load SMC proteins in these other bacterial species (Niki and Yano, 2016; Yano and Niki, 2017), future efforts could investigate a potential direct link between MukB spreading and rDNA genes in E. coli.

Next, we showed that the local chromosomal regions that colocalize with MukBEF are on average almost twice as compacted as the rest of the chromosome (Figure 3A). Again, MukBEF required its ATPase activity to carry out its DNA-compacting function, as the MukB^{EQ}EF mutant showed no such increased compaction in MukBEF-occupied regions and had an altered chromosome width and length compared to wildtype (Figures 3B,C). This is consistent with the

hypothesis that the local compaction is due to loop extrusion by MukBEF.

MukBEF and MatP are not merely factors that set the density of the chromosome, but they also play a major role in DNA segregation. For instance, MukBEF was recently shown to direct newly replicated origins toward the daughter cells (Mäkelä et al., 2021), and as a result, deletion of MukB resulted in anucleated cells and in cells that have unsegregated oriC at the older cell pole. MatP was conversely shown to be necessary for the proper localization of the ter domain after segregation and subsequent cell division (Galli et al., 2017) and MatP was found to be involved in sister cohesion during the chromosomal movement right before cell division (Crozat et al., 2020). Here, we confirm that segregation defects result from the disruption of the MatP or MukBEF function, after which E. coli chromosomes does adopt a toroidal dimer configuration (Figures 4A,B,E). In some cases, toroidal chromosomes consisting of two or more chromosomes formed, recognized by the multiple ori and ter foci (Supplementary Figure S7) and a higher total DNA sum signal (Figure 4C). The variability in relative positioning of ori and ter foci along the chromosome in these cells might be the result of a disruption of recombination, where sister chromosomes remain concatenated and can move relative to each other. In the case of $\Delta mukB$, the fixed ori-ter-ori-ter order along the chromosome indicates the conjoinment of sisters into a single chromosome -possibly through improper recombination between sister dif sites. These data are consistent with recent HiC data showing that both deletion of mukB or matP (Lioy et al., 2018), as well as extended sister-chromosome catenation through impairment of topo activity (Conin et al., 2022) can lead to massive reorganization of the E. coli genome and to emergence of novel chromosome contact points.

As MukBEF and MatP themselves are not known to directly regulate sister chromosome decatenation, questions arise about other molecular players involved in the pathway to the observed segregation defects. The existing literature allows identification of potential interaction partners that may explain our observations (Figure 4F). For example, loss of the ability to fully segregate chromosomes in $\Delta mukB$ can at least in part be understood through the inherent loss of MukBEF's ability to recruit chromosomal TopoIV, leading to reduced sister untangling throughout segregation (Fisher et al., 2021). Further, MatP was found to be essential for DNA translocation by the FtsK protein (Stouf et al., 2013), which is another component that processes DNA in the late stages of cell division (Wang et al., 2020). The FtsK-mediated translocation at dif ultimately ceases upon contact with the XerCD recombinase system (Bonné et al., 2009; Graham et al., 2010), and XerCD and TopoIV decatenate the sister chromosomes in the final phase of segregation (Sciochetti and Piggot, 2000; el Sayyed et al., 2016; Farrokhi et al., 2019). So, there are multiple observations that exemplify MatP's involvement in the tight regulation of TopoIVmediated sister-resolution at ter: MatP's competition with TopoIV to bind MukB (Nicolas et al., 2014), its function to actively displace MukBEF from ter, and the XerCD-TopoIV interaction. XerCD necessitates and specifically guides TopoIV to dif to resolve concatenated sisters (Gogou et al., 2021). matP deletion could therefore indirectly disrupt chromosome recombination, leading to a single toroidal dimer chromosome instead of properly decatenated sisters. In the absence of MatP, the MukBEF-TopoIV complex spreads to the ter region, where XerCD normally orchestrates sister

decatenation by guiding TopoIV to a specific site. Predominant MatP binding to the hinge of TopoIV in the *matS*-MatP-enriched ter could minimize random action of TopoIV in wildtype cells.

All in all, applying our approach to resolve protein-chromosome spatial interactions in widened live *E. coli* cells provided new insights in how MukBEF and MatP shape the chromosome. By performing quantitative fluorescence microscopy on artificially enlarged cells, we effectively gained spatial resolution of the interactions between multiple simultaneously tagged targets. This approach allowed us to acquire new information on how MukBEF and MatP localize relative to each other and along the genome. The presented platform holds good potential to further resolve the spatial interactome that governs chromosome homeostasis, replication, and segregation in prokaryotes.

Materials and methods

Strain construction

All strains were derivatives of E. coli K12 AB1157 strain and were constructed through P1 transduction (Thomason et al., 2014). To construct strain AJ2820, which has the wildtype MukB replaced by mukB-mYPet at the original position, strain SN192 (Badrinarayanan et al., 2012; Nolivos et al., 2016) (lacO240::hyg at ori1, tetO240::gen at ter3, Plac-lacImCherry frt at leuB, Plac-tetR-mCerulean frt at galK, mukB-mYPet frt), was transduced with P1 phage derived from FW1957 (Danilova et al., 2007) ($dnaC2(ts) \Delta mdoB::aph:: frt$) to introduce the temperature sensitive DnaC variant.. To construct strain AJ2822(*AmatP*), strain SN302 (Nolivos et al., 2016) (lacO240::hyg at ori1, tetO240::gen at ter3, Plac-lacImCherry frt at leuB, Plac-tetR-mCerulean frt at galK, *mukB-mYPet frt,* Δ *matP::cat CM*^{*R*}), a kind gift from David Sherratt, was transduced with P1 phage derived from FW1957 (Danilova et al., 2007) $(dnaC2(ts) \Delta m doB::aph:: frt)$ to introduce the temperature sensitive DnaC variant. *AmukB* (strain Ab243) and MukB^{EQ}EF (strain SN311) were generated elsewhere (Nolivos et al., 2016). To construct strain AJ2843 (MatP-mCherry), BN2830 cells (MG1655, hupA-mYPet:: frt, dnaC2 (ts):: aph frt kanR) (Japaridze et al., 2020) was transduced with P1 phage derived from MG1655 matP-mcherry::kan (Mercier et al., 2008) a kind gift from Olivier Espeli.

Growth conditions

For obtaining cells with circular chromosomes, we grew cells in liquid M9 minimal medium (Fluka Analytical) supplemented with 2 mM MgSO₄, 0.1 mM CaCl₂, 0.4% glycerol (Sigma-Aldrich), and 0.01% PHA (Fluka Analytical) overnight at 30°C to reach late exponential phase. On the day of the experiment, the overnight culture was refreshed (1:100 vol) by growing for 2 h on fresh M9 minimal medium at 30°C. We then pipetted 1 μ l culture onto a cover glass and immediately covered the cells with a flat agarose pad, containing the above composition of M9 medium, A22 (final 4 μ g/mL), as well as 3% agarose (Wu et al., 2019a). The cover glass was then placed onto a baseplate and sealed with parafilm to prevent evaporation. The baseplate was placed onto the microscope inside a heated chamber set at 40°C for 2.5 h to stop the cells from replicating and to let them grow into round shapes. In $\Delta mukB$ cells incubation

and imaging were performed at room temperature (22 $^{\circ}$ C) as these cells are unable to grow at higher temperatures.

For experiments with cells lacking HU-mYpet labeling (AB243, AJ2820, and AJ2822) we used DAPI to stain the nucleoids. On the day of the experiment, the overnight culture was refreshed (1:100 vol) by growing for 2h on fresh M9 minimal medium at 30°C. We then added A22 (final 4µg/mL) to the cell culture and transferred the sample to a 40°C incubator for 2.5 h. To minimize any possible artifacts arising from the incubation of cells with the DAPI (Japaridze et al., 2015; Supplementary Figure S9), we added DAPI (final concentration 1 µg/mL) last, just before imaging. 1 mL of the cell sample was incubated with DAPI for less than 1 min and then centrifuged at 4000 rpm for 2 min. 900 µL of the supernatant was removed and the pellet was resuspended in the remaining ~100 µl medium. 1 µL of the remaining culture was deposited onto the cover glass immediately covered with a flat agarose pad, containing M9 medium, A22 (final 4µg/mL), as well as 3% agarose. The sample was placed onto the microscope-stage and imaged immediately.

For treatment of replicating cells with Novobiocin (Maxwell, 1993), we first grew the cells in the presence of A22 as described above for 2.5 h to ensure they reach desired size and shape. Then we moved the baseplate to room-temperature for 10 min to re-initiate replication and afterwards added 10 μ l of Novobiocin (~50 μ g/mL final) to the agarose pad during replication initiation phase. Finally, the cells were moved back to 40°C chamber and imaged.

Fluorescence imaging

Wide-field Z scans were carried out using a Nikon Ti-E microscope with a 100X CFI Plan Apo Lambda Oil objective with an NA of 1.45. The field of view corresponded to 2048 × 2048 pixels with a pixel size of 0.065 µm x 0.065 µm. The microscope was enclosed by a custom-made chamber that was pre-heated overnight and kept at 40°C (except when imaging the *ΔmukB* cells). mCerulean was excited by SpectraX LED (Lumencor) λ_{ex} = 430–450 through a CFP filter cube ($\lambda_{ex} / \lambda_{bs} / \lambda_{em}$ = 426–446 / 455 / 460–500 nm). mYPet signal was excited by SpectraX LED λ_{ex} = 510/25 nm through a triple band-pass filter λ_{em} = 465/25–545/30–630/60 nm. mCherry signal was excited by SpectraX LED λ_{ex} = 575/25 through the same triple band-pass filter. Fluorescent signals were captured by Andor Zyla USB3.0 CMOS Camera. For each channel, between 3–19 slices were taken with a vertical step size of 227 nm (up to 2.3 µm in total).

Image deconvolution

Image stacks of 3–19 slices of Z stack in wide-field imaging were deconvolved using the Huygens Professional deconvolution software (Scientific Volume Imaging, Hilversum, The Netherlands), using an iterative Classic Maximum Likelihood Estimate (CMLE) algorithm with a point spread function (PSF) experimentally measured using 200 nm multicolor Tetrabeads (Invitrogen). The PSF of the single-frame non-deconvolved widefield images had a FWHM of 350 nm horizontally and 800 nm vertically. Deconvolution reduced the out-of-focus signal in the images, which also led to an improvement in lateral resolution.

Automated cell identification

Phase contrast images were fed into a customized Matlab program described earlier (Wu et al., 2019a), to produce masks of cell boundaries, which then were used to allocate chromosomes and foci in other fluorescence channels. A manual correction and rejection process was carried out as a final step of quality control, to correct or reject cells when neighboring cells were too close to allow the automated program to be distinguished.

Compaction, colocalization, foci counting and asymmetricity

First, a customized Matlab program was used to threshold all fluorescence signals with a Gaussian filter and remove background signal. The fluorescence intensity of the DNA signal was used as a direct measure for its density and relative compaction. The chromosome compaction was determined by measuring the average DNA fluorescence intensity of the regions where MukB was present compared to the rest of the chromosome per cell. To account for the size difference between the different cluster sizes, the average DNA intensity was normalised per area. If the average DNA intensity in the clusters colocalizing with MukB was similar to the intensity of the clusters that did not colocalize with MukB, the compaction ratio would be 1 (Supplementary Figure S10).

We defined MukB-DNA colocalization as the percentage of the total DNA intensity that overlaps with the MukB mask. As both ori and ter are single spots in approximation, we made colocalization of MukB with the foci binary. Either (part of) a locus overlapped with MukB, which we classified as colocalization or there was no overlap and thus no colocalization. The ori and ter foci were counted both visually as well as based on the sum fluorescent intensity for each cell and cells with more ter than ori foci were discarded.

To measure the asymmetricity of DNA and MukBEF signal, first the fluorescence intensities and the central ridge of the chromosome toroid were measured. Next, the geometric orientation was set by measuring a central axis through the position of the origin of replication, which we define as the border between the right and left chromosome arm, and the center-of-mass of the chromosome (not to be confused with the genomic left and right arms). Only a circular area around the origin was evaluated for local symmetry. The diameter of the masking circle was 20 pixels = 1.3μ m, in order to encompass the typical chromosome width and the typical local chromosome cluster sizes (Supplementary Figure S4). The local asymmetricity *A* for the DNA and MukB channels was defined by the following formula:

$$A = \frac{\left|\sum I_R - \sum I_L\right|}{\sum I_R + \sum I_L}$$

Where ΣI_R and ΣI_L are the sum fluorescent signals (DNA and MukB) on the right and left sides of the chromosome, respectively. If the fluorescent intensity is equally spread, the asymmetricity is equal to 0, if all signal is on one of the chromosome arms, then it is equal to1. Due to the unknown orientation of the cell, the colors (red and blue on Figure 2D) cannot be attributed to the genomic orientation of left and right chromosome arms.

Statistics

We used MATLAB's built-in functions for statistical analysis. On all figures, the following conventions are used: not significant (NS) 0.05 < p, *0.01 , *<math>0.001 , ***<math>p < 0.001.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary material, further inquiries can be directed to the corresponding author.

Author contributions

AJ and CD conceived the idea. AJ, RvW, and CG collected and analyzed the data. JK wrote the image analysis software. DvB performed Chip-seq analysis. AJ and RvW performed the bacterial strain cloning. CD supervised the project. All authors designed the experiments and contributed to the interpretation of the results and writing of the manuscript.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2023.1107093/ full#supplementary-material

References

Antar, H., Soh, Y. M., Zamuner, S., Bock, F. P., Anchimiuk, A., De Los Rios, P., et al. (2021). Relief of ParB autoinhibition by parS DNA catalysis and recycling of ParB by CTP hydrolysis promote bacterial centromere assembly. *Sci. Adv.* 7:eabj2854. doi: 10.1126/sciadv.abj2854

Badrinarayanan, A., Reyes-Lamothe, R., Uphoff, S., Leake, M. C., and Sherratt, D. J. (2012). In vivo architecture and action of bacterial structural maintenance of chromosome proteins. *Sci.* 338, 528–531. doi: 10.1126/science.1227126

Blot, N., Mavathur, R., Geertz, M., Travers, A., and Muskhelishvili, G. (2006). Homeostatic regulation of supercoiling sensitivity coordinates transcription of the bacterial genome. *EMBO Rep.* 7, 710–715. doi: 10.1038/SJ.EMBOR.7400729

Böhm, K., Giacomelli, G., Schmidt, A., Imhof, A., Koszul, R., Marbouty, M., et al. (2020). Chromosome organization by a conserved condensin-ParB system in the actinobacterium Corynebacterium glutamicum. *Nat. Commun.* 11, 1485–1417. doi: 10.1038/s41467-020-15238-4

Bonné, L., Bigot, S., Chevalier, F., Allemand, J. F., and Barre, F. X. (2009). Asymmetric DNA requirements in Xer recombination activation by FtsK. *Nucleic Acids Res.* 37, 2371–2380. doi: 10.1093/NAR/GKP104

Bürmann, F., Funke, L. F. H., Chin, J. W., and Löwe, J. (2021). Cryo-EM structure of MukBEF reveals DNA loop entrapment at chromosomal unloading sites. *Mol. Cell* 81, 4891–4906.e8. doi: 10.1016/j.molcel.2021.10.011

Chen, N., Zinchenko, A. A., Yoshikawa, Y., Araki, S., Adachi, S., Yamazoe, M., et al. (2008). ATP-induced shrinkage of DNA with MukB protein and the MukBEF complex of *Escherichia coli*. *J. Bacteriol.* 190, 3731–3737. doi: 10.1128/JB.01863-07

Conin, B., Billault-Chaumartin, I., el Sayyed, H., Quenech'du, N., Cockram, C., Koszul, R., et al. (2022). Extended sister-chromosome catenation leads to massive reorganization of the *E. coli* genome. *Nucleic Acids Res.* 50, 2635–2650. doi: 10.1093/NAR/GKAC105

Crozat, E., Tardin, C., Salhi, M., Rousseau, P., Lablaine, A., Bertoni, T., et al. (2020). Post-replicative pairing of sister ter regions in *Escherichia coli* involves multiple activities of MatP. *Nat. Commun.* 11, 3796–3712. doi: 10.1038/s41467-020-17606-6

Danilova, O., Reyes-Lamothe, R., Pinskaya, M., Sherratt, D., and Possoz, C. (2007). MukB colocalizes with the oriC region and is required for organization of the two *Escherichia coli* chromosome arms into separate cell halves. *Mol. Microbiol.* 65, 1485–1492. doi: 10.1111/j.1365-2958.2007.05881.x

el Sayyed, H., le Chat, L., Lebailly, E., Vickridge, E., Pages, C., Cornet, F., et al. (2016). Mapping topoisomerase IV binding and activity sites on the *E. coli* genome. *PLoS Genet.* 12, e1006025–e1006022. doi: 10.1371/journal.pgen.1006025

Espéli, O., Borne, R., Dupaigne, P., Thiel, A., Gigant, E., Mercier, R., et al. (2012). A MatP-divisome interaction coordinates chromosome segregation with cell division in *E. coli. EMBO J.* 31, 3198–3211. doi: 10.1038/EMBOJ.2012.128

Farrokhi, A., Liu, H., and Szatmari, G. (2019). Characterization of the chromosome dimer resolution site in *Caulobacter crescentus*. J. Bacteriol. 201, e00391–e00319. doi: 10.1128/JB.00391-19

Fisher, G. L., Bolla, J. R., Rajasekar, K. V., Mäkelä, J., Baker, R., Zhou, M., et al. (2021). Competitive binding of MatP and topoisomerase IV to the MukB hinge domain. *elife* 10, 1–25. doi: 10.7554/eLife.70444

Galli, E., Midonet, C., Paly, E., and Barre, F. X. (2017). Fast growth conditions uncouple the final stages of chromosome segregation and cell division in *Escherichia coli*. *PLoS Genet*. 13:e1006702. doi: 10.1371/JOURNAL.PGEN.1006702

Gogou, C., Japaridze, A., and Dekker, C. (2021). Mechanisms for chromosome segregation in bacteria. *Front. Microbiol.* 12:685687. doi: 10.3389/fmicb.2021.685687

Goto, T., and Wang, J. C. (1982). Yeast DNA topoisomerase II. An ATP-dependent type II topoisomerase that catalyzes the catenation, decatenation, unknotting, and relaxation of double-stranded DNA rings. *J. Biol. Chem.* 257, 5866–5872. doi: 10.1016/S0021-9258(19)83859-0

Graham, J. E., Sivanathan, V., Sherratt, D. J., and Arciszewska, L. K. (2010). FtsK translocation on DNA stops at XerCD-dif. *Nucleic Acids Res.* 38, 72–81. doi: 10.1093/NAR/GKP843

Japaridze, A., Benke, A., Renevey, S., Benadiba, C., and Dietler, G. (2015). Influence of DNA binding dyes on bare DNA structure studied with atomic force microscopy. *Macromol.* 48, 1860–1865. doi: 10.1021/ma502537g

Japaridze, A., Gogou, C., Kerssemakers, J. W. J., Nguyen, H. M., and Dekker, C. (2020). Direct observation of independently moving replisomes in *Escherichia coli. Nat. Commun.* 11, 3109–3110. doi: 10.1038/s41467-020-16946-7

Karczmarek, A., Baselga, R. M. A., Alexeeva, S., Hansen, F. G., Vicente, M., Nanninga, N., et al. (2007). DNA and origin region segregation are not affected by the transition from rod to sphere after inhibition of *Escherichia coli* MreB by A22. *Mol. Microbiol.* 65, 51–63. doi: 10.1111/J.1365-2958.2007.05777.X

Khan, S. R., Mahaseth, T., Kouzminova, E. A., Cronan, G. E., and Kuzminov, A. (2016). Static and dynamic factors limit chromosomal replication complexity in *Escherichia coli*, avoiding dangers of runaway overreplication. *Genet.* 202, 945–960. doi: 10.1534/genetics.115.184697

Kim, T. H., and Dekker, J. (2018). ChIP-quantitative polymerase chain reaction (ChIPqPCR). *Cold Spring Harb. Protoc.* 2018:pdb.prot082628. doi: 10.1101/pdb.prot082628

Kumar, R., Bahng, S., and Marians, K. J. (2022). The MukB-topoisomerase IV interaction mutually suppresses their catalytic activities. *Nucleic Acids Res.* 50, 2621–2634. doi: 10.1093/nar/gkab1027

Kumar, R., Nurse, P., Bahng, S., Lee, C. M., and Marians, K. J. (2017). The MukB-topoisomerase IV interaction is required for proper chromosome compaction. *J. Biol. Chem.* 292, 16921–16932. doi: 10.1074/jbc.M117.803346

Lieberman-Aiden, E., van Berkum, N. L., Williams, L., Imakaev, M., Ragoczy, T., Telling, A., et al. (2009). Comprehensive mapping of long-range interactions reveals folding principles of the human genome. *Sci.* 326, 289–293. doi: 10.1126/science.1181369

Lioy, V. S., Cournac, A., Marbouty, M., Duigou, S., Mozziconacci, J., Espéli, O., et al. (2018). Multiscale structuring of the *E. coli* chromosome by nucleoid-associated and condensin proteins. *Cells* 172, 771–783.e18. doi: 10.1016/j.cell.2017.12.027

Luijsterburg, M. S., Noom, M. C., Wuite, G. J. L., and Dame, R. T. (2006). The architectural role of nucleoid-associated proteins in the organization of bacterial chromatin: a molecular perspective. *J. Struct. Biol.* 156, 262–272. doi: 10.1016/J. JSB.2006.05.006

Mäkelä, J., and Sherratt, D. J. (2020). Organization of the *Escherichia coli* chromosome by a MukBEF axial Core. *Mol. Cell* 78, 250–260.e5. doi: 10.1016/J.MOLCEL.2020.02.003

Mäkelä, J., Uphoff, S., and Sherratt, D. J. (2021). Nonrandom segregation of sister chromosomes by *Escherichia coli* MukBEF. *Proc. Natl. Acad. Sci. U. S. A.* 118:e2022078118. doi: 10.1073/pnas.2022078118

Maxwell, A. (1993). The interaction between coumarin drugs and DNA gyrase. *Mol. Microbiol.* 9, 681–686. doi: 10.1111/j.1365-2958.1993.tb01728.x

Mercier, R., Petit, M. A., Schbath, S., Robin, S., El Karoui, M., Boccard, F., et al. (2008). The MatP/matS site-specific system organizes the terminus region of the *E. coli* chromosome into a macrodomain. *Cells* 135, 475–485. doi: 10.1016/j.cell.2008.08.031

Mickiewicz, K. M., Kawai, Y., Drage, L., Gomes, M. C., Davison, F., Pickard, R., et al. (2019). Possible role of L-form switching in recurrent urinary tract infection. *Nat. Commun.* 10:4379. doi: 10.1038/s41467-019-12359-3

Nicolas, E., Upton, A. L., Uphoff, S., Henry, O., Badrinarayanan, A., and Sherratt, D. (2014). The SMC complex MukBEF recruits topoisomerase IV to the origin of replication region in live *Escherichia coli*. *MBio* 5, e01001–e01013. doi: 10.1128/mBio.01001-13

Niki, H., Jaffé, A., Imamura, R., Ogura, T., and Hiraga, S. (1991). The new gene mukB codes for a 177 kd protein with coiled-coil domains involved in chromosome partitioning of *E. coli. EMBO J.* 10, 183–193. doi: 10.1002/j.1460-2075.1991.tb07935.x

Niki, H., and Yano, K. (2016). In vitro topological loading of bacterial condensin MukB on DNA, preferentially single-stranded DNA rather than double-stranded DNA. *Sci. Rep.* 6:29469. doi: 10.1038/srep29469

Nolivos, S., and Sherratt, D. (2014). The bacterial chromosome: architecture and action of bacterial SMC and SMC-like complexes. *FEMS Microbiol. Rev.* 38, 380–392. doi: 10.1111/1574-6976.12045

Nolivos, S., Upton, A. L., Badrinarayanan, A., Müller, J., Zawadzka, K., Wiktor, J., et al. (2016). MatP regulates the coordinated action of topoisomerase IV and MukBEF in chromosome segregation. *Nat. Commun.* 7:10466. doi: 10.1038/ncomms10466

Peng, H., and Marians, K. J. (1993). Decatenation activity of topoisomerase IV during oriC and pBR322 DNA replication in vitro. *Proc. Natl. Acad. Sci. U. S. A.* 90, 8571–8575. doi: 10.1073/pnas.90.18.8571

Petrushenko, Z. M., Lai, C. H., and Rybenkov, V. V. (2006). Antagonistic interactions of kleisins and DNA with bacterial condensin MukB. *J. Biol. Chem.* 281, 34208–34217. doi: 10.1074/jbc.M606723200

Reyes-Lamothe, R., Wang, X., and Sherratt, D. (2008). *Escherichia coli* and its chromosome. *Trends Microbiol.* 16, 238–245. doi: 10.1016/j.tim.2008.02.003

Rybenkov, V. V., Herrera, V., Petrushenko, Z. M., and Zhao, H. (2014). MukBEF, a chromosomal organizer. *J. Mol. Microbiol. Biotechnol.* 24, 371–383. doi: 10.1159/000369099

Saifi, B., and Ferat, J. L. (2012). Replication fork reactivation in a dnaC2 mutant at non-permissive temperature in *Escherichia coli*. *PLoS One* 7:e33613. doi: 10.1371/journal.pone.0033613

Sciochetti, S. A., and Piggot, P. J. (2000). A tale of two genomes: resolution of dimeric chromosomes in *Escherichia coli* and Bacillus subtilis. *Res. Microbiol.* 151, 503–511. doi: 10.1016/S0923-2508(00)00220-5

Seol, Y., Hardin, A. H., Strub, M. P., Charvin, G., and Neuman, K. C. (2013). Comparison of DNA decatenation by *Escherichia coli* topoisomerase IV and topoisomerase III: implications for non-equilibrium topology simplification. *Nucleic Acids Res.* 41, 4640–4649. doi: 10.1093/nar/gkt136

Stouf, M., Meile, J. C., and Cornet, F. (2013). FtsK actively segregates sister chromosomes in *Escherichia coli*. *Proc. Natl. Acad. Sci. U. S. A.* 110, 11157–11162. doi: 10.1073/pnas.1304080110

Thanbichler, M., Viollier, P. H., and Shapiro, L. (2005). The structure and function of the bacterial chromosome. *Curr. Opin. Genet. Dev.* 15, 153–162. doi: 10.1016/j.gde.2005.01.001

Thomason, L. C., Costantino, N., and Court, D. L. (2014). E. coli genome manipulation by P1 transduction. Curr. Protoc. Mol. Biol. 1, 1–17. doi: 10.1002/0471142727.mb0117s79

Tran, N. T., Laub, M. T., and Le, T. B. K. (2017). SMC progressively aligns chromosomal arms in *Caulobacter crescentus* but is antagonized by convergent transcription. *Cell Rep.* 20, 2057–2071. doi: 10.1016/j.celrep.2017.08.026

Travers, A., and Muskhelishvili, G. (2005). Bacterial chromatin. Curr. Opin. Genet. Dev. 15, 507–514. doi: 10.1016/J.GDE.2005.08.006

Varma, A., and Young, K. D. (2009). In *Escherichia coli*, MreB and FtsZ direct the synthesis of lateral cell wall via independent pathways that require PBP 2. *J. Bacteriol.* 191, 3526–3533. doi: 10.1128/JB.01812-08

Wang, M., Fang, C., Ma, B., Luo, X., and Hou, Z. (2020). Regulation of cytokinesis: FtsZ and its accessory proteins. *Curr. Genet.* 66, 43–49. doi: 10.1007/s00294-019-01005-6

Wang, X., Le, T. B. K., Lajoie, B. R., Dekker, J., Laub, M. T., and Rudner, D. Z. (2015). Condensin promotes the juxtaposition of DNA flanking its loading site in *Bacillus subtilis. Genes Dev.* 29, 1661–1675. doi: 10.1101/gad.265876.115

Wang, X., Liu, X., Possoz, C., and Sherratt, D. J. (2006). The two *Escherichia coli* chromosome arms locate to separate cell halves. *Genes Dev.* 20, 1727–1731. doi: 10.1101/GAD.388406

Wang, X., and Sherratt, D. J. (2010). Independent segregation of the two arms of the *Escherichia coli* ori region requires neither RNA synthesis nor MreB dynamics. *J. Bacteriol.* 192, 6143–6153. doi: 10.1128/JB.00861-10/SUPPL_FILE/WANG_AND_SHERRATT_SUPP.DOC

Woo, J. S., Lim, J. H., Shin, H. C., Suh, M. K., Ku, B., Lee, K. H., et al. (2009). Structural studies of a bacterial condensin complex reveal ATP-dependent disruption of intersubunit interactions. *Cells* 136, 85–96. doi: 10.1016/J. CELL.2008.10.050

Wu, F., Japaridze, A., Zheng, X., Wiktor, J., Kerssemakers, J. W. J., and Dekker, C. (2019a). Direct imaging of the circular chromosome in a live bacterium. *Nat. Commun.* 10, 2194–2199. doi: 10.1038/s41467-019-10221-0

Wu, F., Swain, P., Kuijpers, L., Zheng, X., Felter, K., Guurink, M., et al. (2019b). Cell boundary confinement sets the size and position of the *E. coli* chromosome. *Curr. Biol.* 29, 2131–2144.e4. doi: 10.1016/J.CUB.2019.05.015

Yamazoe, M., Onogi, T., Sunako, Y., Niki, H., Yamanaka, K., Ichimura, T., et al. (1999). Complex formation of MukB, MukE and MukF proteins involved in chromosome partitioning in *Escherichia coli. EMBO J.* 18, 5873–5884. doi: 10.1093/emboj/18.21.5873

Yano, K., and Niki, H. (2017). Multiple cis-acting rDNAs contribute to nucleoid separation and recruit the bacterial condensin Smc-ScpAB. *Cell Rep.* 21, 1347–1360. doi: 10.1016/j.celrep.2017.10.014