

## ARTICLE

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# Bacterial partition complexes segregate within the volume of the nucleoid

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Precise and rapid DNA segregation is required for proper inheritance of genetic material. In most bacteria and archaea, this process is assured by a broadly conserved mitotic-like apparatus in which a NTPase (ParA) displaces the partition complex. Competing observations and models imply starkly different 3D localization patterns of the components of the partition machinery during segregation. Here we use super-resolution microscopies to localize in 3D each component of the segregation apparatus with respect to the bacterial chromosome. We show that Par proteins locate within the nucleoid volume and reveal that proper volumetric localization and segregation of partition complexes requires ATPase and DNA-binding activities of ParA. Finally, we find that the localization patterns of the different components of the partition system highly correlate with dense chromosomal regions. We propose a new mechanism in which the nucleoid provides a scaffold to guide the proper segregation of partition complexes.

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he preservation of all living forms relies on the faithful segregation of their duplicated genetic material upon cell division. To assure the relocalization of replicated DNA, prokaryotic organisms have evolved dedicated apparatuses, namely partition systems (Par), that are classified according to the nature of the ATPase providing the energy to displace DNA<sup>1,2</sup>. In particular, Type I Par systems are the most prevalent in low copy-number plasmids and the only type known to be present on bacterial chromosomes<sup>3</sup>. It contains three essential elements: a repeat of centromere-like DNA sequences (*parS*), a DNA-binding protein (ParB) and a deviant Walker A-type ATPase (ParA)<sup>2,4</sup>. ParB specifically recognizes *parS* and, upon binding, forms the partition complex (ParBS)<sup>5–8</sup> that will be displaced by the action of ParA. The partition systems of plasmids F and P1 in *Escherichia coli*, and of the chromosome of *Bacillus subtilis*, are among the best characterized<sup>9</sup>.

Different models have been proposed to account for the assembly of partition complexes and for the role of ParA in their segregation. At physiological levels, ParB binds over a large genomic region flanking *parS* (10–12 kb). This extended binding pattern has been interpreted as resulting from the one-dimensional spreading of ParB from *parS*<sup>10–12</sup>, and more recently because of the dynamic stochastic binding and unbinding of ParB around *parS*<sup>13</sup>. Partition complexes are initially positioned at midcell and segregate to their new subcellular locations after or concomitantly with replication<sup>1</sup>. Importantly, proper segregation of partition complexes requires the ParB-stimulated ATPase activity of ParA as well as ParA's ability to bind nonspecific DNA (ns-DNA). Altering these activities leads to different degrees of plasmid instability and chromosome segregation defects<sup>14–18</sup>.

Two families of models have been developed to explain the mechanism by which ParA segregates partition complexes. The first class of models ('filament' models) proposes that dynamic ParA filaments pull partition complexes to their resting positions, either around or straight through the nucleoid (Fig. 1a, upper panel)<sup>19,20</sup>. The second class of models ('non-filament' models) involves a diffusion-ratchet mechanism ('brownian-ratchet' model) in which ParA dimers or short filaments push-pull partition complexes on the surface of the nucleoid (Fig. 1a, lower panel)<sup>21-24</sup>. A variation of this mechanism ('DNA relay' model) was recently proposed in which chromosome elasticity also plays a role by partly powering movement of the partition complex<sup>25</sup> (Fig. 1a, lower panel). All these models were largely based on biochemical assays (for example, polymerization, DNA binding and ATPase activity assays), two-dimensional (2D) imaging and in vitro microscopy assays monitoring the dynamics of ParBbound plasmids on a 2D ParA-bound DNA carpet, and proposed that partition components segregate through the space between the nucleoid surface and the membrane. In brief, these various models proposed distinct localization patterns of Par components within or around the nucleoid. Importantly, these patterns constitute essential ingredients of the proposed mechanisms. Discerning between these models requires the knowledge, in vivo, of the precise three-dimensional (3D) localization of the machinery components, yet so far this has been limited by the optical resolution of conventional microscopy methods.

Here we combine whole-chromosome labelling strategies with multicolour 3D super-resolution technologies to investigate the segregation machineries of plasmid F and the *B. subtilis* chromosome. Our strategy allowed us to reveal that partition complexes as well as the ParA ATPase reside within the nucleoid interior. By functional studies, we determine that both Walker-type ATPase and dynamic DNA-binding activities of ParA are necessary for proper volumetric relocalization of partition complexes during segregation. Finally, we unveil that both partition complexes and

ParA tend to colocalize with high-density chromosomal regions (HDRs), suggesting a new mechanism by which chromosomal DNA may play a role by locally enriching components of the segregation apparatus to promote proper segregation.

#### Results

Partition complexes reside inside the bacterial nucleoid. We first implemented an experimental strategy to directly detect the 3D localization of partition complexes within the volume of the nucleoid. To this end, we combined high-density whole-chromosome labelling with high-throughput, 3D structured illumination microscopy (3D-SIM)<sup>26</sup>, a super-resolution method that provides a twofold increase in both lateral and axial resolution as compared with diffraction-limited microscopies. DNA was either fluorescently labelled by the ns-DNA-binding protein histone-like U93 (HU) tagged with monomeric fluorescent protein mCherry (nucleoids in E. coli strains) or by the intercalating agent 4.6-diamidino-2phenylindole (nucleoids in B. subtilis strains), while localization of partition complexes was detected by tagging ParB with a fluorescent tag. ParB from the F-plasmid (ParB<sub>F</sub> or SopB) was tagged with monomeric Venus (mVenus) at the locus under the control of the endogenous autoregulated promoter, while B. subtilis ParB (ParB<sub>Bsu</sub> or Spo0J) was tagged using monomeric green fluorescent protein (GFP) also at its natural locus (see Supplementary Methods).

We simultaneously imaged the bacterial chromosome and plasmidic/chromosomal ParB by 3D-SIM. *E. coli* nucleoids displayed a range of sizes between 800 and 4,000 nm in length and 530 ± 50 nm in width (full-width at half-maximum (FWHM); Supplementary Fig. 1A,B). ParB foci were observed in a large proportion of cells (~98%; Fig. 1b,c, and pie charts therein), consistent with the fluorescent tag not affecting plasmid stability<sup>13</sup> or chromosome segregation<sup>26</sup>. The number of partition complexes per nucleoid ( $2.2 \pm 0.8$ , mean  $\pm$  s.d., Supplementary Fig. 1C) agreed well with the distributions observed using widefield microscopy or photo-activated localization microscopy (PALM)<sup>13</sup>. Partition complexes had a typical size of  $140 \pm 35$  nm (Supplementary Fig. 2), consistent with the sizes of partition complexes recently reported using PALM (~150 nm)<sup>13</sup>.

Importantly, partition complexes were confined within the bacterial nucleoid for both plasmidic and chromosomal systems (Fig. 1b,c), consistent with preliminary observations from B. subtilis ParB<sup>26</sup>. Single projections of isolevel representations of 3D densities can often be misleading. Thus, we developed an algorithm to simultaneously visualize both colours (that is, nucleoid and ParB) in 3D as well as the three orthogonal 2D projections at the single-cell level. This visualization clearly reveals that ParB is localized within the bacterial nucleoid, and is particularly close to the centre of the nucleoid long axis (Fig. 1d,e, Supplementary Movies 1 and 2 and Supplementary Fig. 3). To determine whether this localization pattern was observed at the population level, we built localization density histograms of the positions of partition complexes within nucleoids from thousands of cells (see Methods for details). Figure 1f,g displays the projection of the cumulated localization of partition complexes for nucleoids of similar sizes (<1,500 nm in length) in 2D representations (xy and yz planes). Lateral histograms (xy plane) revealed that both plasmidic and chromosomal partition complexes are positioned along the longitudinal axis (Fig. 1f,g). ParBS<sub>F</sub> partition complexes localized to the quarter nucleoid positions, while  $ParBS_{Bsu}$  complexes were positioned close to the nucleoid edges, as recently described<sup>26,27</sup>. Importantly, axial histograms (yz plane) revealed that  $ParB_{F}$  and  $ParB_{Bsu}$  complexes were also radially confined to the centre of the nucleoid axis (Fig. 1f,g, lower panels) but with a larger spread with respect to the nucleoid volume because of the lower axial resolution of 3D-SIM.

Acquisition of a single two-colour volume in 3D-SIM requires  $\sim 20-50$  s and involves complex image-reconstruction algorithms. Thus, we used a second advanced microscopy method to discard any potential image-blurring effects caused by the



movement of the nucleoid and partition complexes during the acquisition time or any image-reconstruction artefact. Multifocus microscopy (MFM)<sup>28</sup> allows for multicolour, near-instantaneous volumetric acquisitions ( $\sim 25 \text{ ms}$  per channel) of biological specimens. Particularly, MFM permits the simultaneous acquisition of nine imaging planes and thus the volumetric acquisitions of entire nucleoids and partition complexes in a single snapshot without resorting to complex image-reconstruction algorithms (Fig. 1h, left and middle panels and Supplementary Fig. 4). Simultaneous MFM imaging of nucleoids and ParB<sub>F</sub> resulted in equivalent results to those obtained by 3D-SIM (Fig. 1h, right panel).

Overall, these results demonstrate that the localization of partition complexes does not require interactions with the bacterial nucleoid surface. However, in that case, how can the ParA ATPase interact with ParB nucleoprotein complexes positioned inside the nucleoid?

ParA displaces ParB within the bacterial nucleoid. Previous in vitro and 2D microscopy studies have led to a model in which ParA forms filaments (Fig. 1a)<sup>19,20,29</sup>. To test this model, we imaged ParA<sub>F</sub>-mVenus using 3D-SIM. We observed that ParA<sub>F</sub> spreads into gradient-like, asymmetric structures with higher density of protein at one nucleoid pole (Fig. 2 and Supplementary Movie 3). The lateral and axial projections of the 3D volume of ParA<sub>F</sub> revealed smaller and dimmer fluorescent patches distributed along the ParA<sub>F</sub> density (Fig. 2a). To determine whether these asymmetric, discontinuous ParA structures distributed over the membrane/nucleoid surface or rather penetrated the nucleoid volume, we imaged ParA<sub>F</sub>-mVenus and nucleoid simultaneously by multicolour 3D-SIM (Fig. 2b and Supplementary Movie 4). Strikingly, ParA<sub>F</sub> was distributed within the interior of the nucleoid extending asymmetrically along the long cell axis. 2D lateral histograms of ParA localization (xy plane) were constructed as previously described but aligning

#### Figure 1 | Partition complexes segregate through the interior of the

bacterial nucleoid. (a) Schematic representation of filament (upper panel) and Brownian-ratchet (lower panel) models for active bacterial DNA segregation. In Brownian-ratchet models, partition complexes are segregated by a pure ParA diffusion-ratchet (lower panel left) or combined with the propulsion provided by the elastic properties of the chromosome (lower panel right). (b,c) Typical multicolour 3D-SIM images of (a) HUmCherry-labelled nucleoids (red) and plasmidic ParB<sub>F</sub>-mVenus (green) in E. coli (DLT3053/pJYB234), or (b) 4,6-diamidino-2-phenylindole-stained nucleoids (red) and chromosomal ParB<sub>Bsu</sub>-GFP (green) in B. subtilis (HM671). (d,e) Characteristic 3D volume of single nucleoids (solid red) and ParB foci (green) with orthogonal 2D projections of nucleoid (red contour) and ParB (green spot) densities for (d) ParB<sub>F</sub>-mVenus and (e) ParB<sub>Bsu</sub>-GFP. (f,g) 2D density distributions of ParB intensity signal along the xy (lateral) and yz (axial) planes for (f) E. coli and (g) B. subtilis strains. Only nucleoids of equal length were used for calculation of ParB densities (see Methods for details). Nucleoid contours are represented with solid black lines with capsid shapes. Dimensions of nucleoid contours were measured as the fullwidth at half-maximum of the intensity signal along each nucleoid dimension. N depicts the number of nucleoids analysed. All panels of the figure represent one representative experiment of at least six biological replicates. (h) Instantaneous 3D MFM of ParB<sub>F</sub> and nucleoids labelled with HU-mCherry. Left: MFM scheme representing the simultaneously acquired nine z planes for fast volumetric imaging. Middle: single processed image in which the nine simultaneously imaged z planes can be visualized at once after correction of chromatic aberrations (see Supplementary Fig. 4 and Supplementary Methods). Right: volumetric reconstruction of bacterial nucleoid (red) and ParB<sub>F</sub> (green) from image in the left panel (only two nucleoids are shown for clarity).



**Figure 2 | ParA is localized within the nucleoid volume.** (a) 3D-SIM image from DLT3053/pJYB243 showing ParA<sub>F</sub> (solid blue) with side projections of ParA<sub>F</sub> densities (colour-coded scale for projections (a.u.) represents fluorescence intensity increasing from blue to red). (b) Typical two-colour, 3D-SIM images of a single nucleoid labelled with HU-mCherry and the ParA<sub>F</sub>-mVenus distribution at its interior. Left: nucleoid periphery is shown in light grey, while higher-density regions at the centre of the nucleoid are shown in orange-red. Right: ParA<sub>F</sub> is represented in blue-green. Middle: overlay of the two colours showing the positioning of ParA<sub>F</sub> relative to the nucleoid. (c) 2D distribution of ParA along the lateral (*xy*) and axial (*yz*) planes of the nucleoids. Nucleoid orientation is aligned from the brightest ParA<sub>F</sub> patch. *N* depicts the number of cells analysed. All panels of the figure depict one representative experiment of at least five biological replicates.

ParA images from different nucleoids to orient the highest density of ParA specifically on the right side of the histogram (Fig. 2c). The region with highest ParA density resided well within the nucleoid volume, and localized close to the quarter nucleoid positions displayed by ParB complexes (Fig. 2c, left panel, dotted lines). Interestingly, a second weaker ParA<sub>F</sub> density maximum emerged near the opposite ParB<sub>F</sub> quarter nucleoid position (Fig. 2c, left panel, compare with Fig. 1f). Axial histograms (*yz* plane) recapitulated well the volumetric distribution of ParA<sub>F</sub>. This distribution was radially confined to the longitudinal axis (as that of ParB<sub>F</sub>), but was more broadly distributed (Fig. 2c, right panel).

**ParA regulates the volumetric localization of ParBS**. These results suggest that the nucleoid scaffold itself or the interaction between partition complexes and ParA could be responsible for the intranucleoid localization of the components of the segregation machinery. To functionally test these hypotheses, we

evaluated the subnucleoid localization of partition complexes in absence of ParA<sub>F</sub> and for specific ParA<sub>F</sub> mutants known to affect DNA binding and ATP hydrolysis activities. First, E. coli cells carrying the ParBS<sub>F</sub> system but not ParA<sub>F</sub> were grown with antibiotics to maintain the F-plasmid. In absence of ParA<sub>F</sub>, partition complexes displayed a very dynamic behaviour that could not be captured using 3D-SIM. As the localization of partition complexes in wild-type cells is equivalent when imaged either by 3D-SIM or widefield deconvolution microscopy (Supplementary Fig. 5), we turned to the latter technique to facilitate the detection of partition complexes in mutant strains. About 55% of cells showed distinct bright  $ParB_F$  foci compared with >98% of cells carrying the wild-type ParABS partition system (Figs 1b and 3a). This finding is expected from the plasmid loss rate observed without ParA<sub>F</sub> (Supplementary Table 1). Volumetric views of single nucleoids reveal that most ParB<sub>E</sub> foci were now located outside the nucleoid (Fig. 3b and Supplementary Movies 5-6). These single-cell observations were recapitulated at the population level by calculating the lateral and axial histograms of ParB localization. ParB<sub>F</sub> was preferentially located at the nucleoid periphery, while the nucleoid interior was mostly devoid of ParB density (Fig. 3c). These results indicate a key role of  $ParA_F$  in the localization of the partition complex within the nucleoid volume.

We next investigated how two key activities of  $ParA_F$ —ATPase stimulation by  $ParB_F$  and ns-DNA binding—regulate the localization of partition complexes by employing  $ParA_F$  mutants.  $ParA_F^{K120Q}$  retains wild-type basal ATPase hydrolysis rate; however, its ATPase activity is no longer stimulated by  $ParB_F^{30}$ . In contrast,  $ParA_F^{K340A}$  retains its ability to interact with  $ParB_F$  but is partially impaired at ns-DNA-binding<sup>14,18</sup>. The percentage of cells displaying  $ParB_F$  foci was reduced from 98% (wild type) to 41% and 21% in the mutants ( $ParA_F^{K120Q}$  and  $ParA_F^{K340A}$ , respectively, pie charts in Fig. 3d–g, and Supplementary Table 1).

Partition complexes localized outside the nucleoid volume for both ParA<sub>F</sub> mutants (Fig. 3e-h and Supplementary Movies 7-9) and tended to localize to regions displaying less spatial confinement (that is, near cell poles and at the locations of future division septa), as previously shown for large protein complexes or plasmids without partitioning systems<sup>31,32</sup>. To validate these results at the population level, we calculated the mean localization of partition complexes within nucleoids (Fig. 3f-i). For  $ParA_{F}^{K120Q}$ , lateral (XY)  $ParB_{F}$  density histograms (Fig. 3f) displayed a similar pattern of localization than that of cells lacking  $ParA_F$  (Fig. 3c), with partition complexes clearly locating at the exterior of the nucleoid. ParAF<sup>K340A</sup> also exhibited abnormal distributions of ParB<sub>F</sub> but with a more heterogeneous distribution (Fig. 3i) likely because of residual interactions between  $ParA_F^{K340A}$  and  $ParB_F$  as well as with ns-DNA. Interestingly, axial  $ParB_F$  density histograms (yz) for both mutants showed a disturbed radial profile in which ParB<sub>F</sub> localized mostly to the edges of nucleoids (Fig. 3f-i, right panels). These results indicate that proper volumetric localization of partition complexes requires ns-DNA binding by ParA as well as the stimulation of ParA's ATPase activity by ParB.

**The partition machinery localizes to high-density regions.** We recently described the existence of a new structural feature of the bacterial nucleoid of *B. subtilis*: HDRs<sup>26</sup>. HDRs represent regions of the nucleoid with denser DNA content and only resolvable by super-resolution microscopies. As both ParB and ParA distributed within the nucleoid volume, we evaluated whether HDRs played a role in their localization.

Using two-colour 3D-SIM, we simultaneously imaged the nucleoid using HU-mCherry labelling and  $ParB_F$ -mVenus. Volumetric images of the nucleoid revealed that HDRs are also

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**Figure 3** | **ParA**<sub>F</sub> **determines the 3D** localization of partition complexes within bacterial nucleoids. (**a**-**g**) Widefield deconvolution imaging of *E. coli* cells harbouring F-plasmid (**a**) lacking ParA<sub>F</sub> (DLT3053/pJYB263; **d**) with ParA<sup>F120Q</sup><sub>F</sub> (DLT3053/pJYB277) and (**g**) with ParA<sup>F340A</sup><sub>F</sub>, an allele with reduced ns-DNA-binding capacity (DLT3053/pJYB279). Representative fields of view of two-colour widefield deconvolution imaging of HU-mCherry-labelled nucleoids (red) and ParB<sub>F</sub>-mVenus (green) are shown for each of the mutants. Pie charts represent the percentage of nucleoids with (green) and without (red) ParB<sub>F</sub> foci, reflecting the frequency of plasmid loss in these mutants. (**b-h**) Characteristic 3D volume of single nucleoids (solid red) and ParB<sub>F</sub> foci (green) with orthogonal 2D projections of nucleoid (red contours) and ParB<sub>F</sub> densities (green spots) for the three mutants shown in **a-g**. (**c-i**) 2D density distributions of ParB<sub>F</sub> intensity signal along the *xy* (lateral) and *yz* (axial) planes for mutants displayed in **a-g**. Solid black lines represent nucleoid periphery. See legend of Fig. 1f,g and Methods for more details. Residual ParB<sub>F</sub> density at midcell in h-i arises from ParB<sub>F</sub> at the centre of nucleoids about to split (see examples in **e,f**). *N* represents number of cells. All panels of the figure depict one representative experiment of at least two biological replicates.

a common feature of E. coli chromosomes (Fig. 4a and Supplementary Movie 10). HDR positions in E. coli were detected as previously described<sup>26</sup> (see Methods for details). Interestingly, HDRs and ParB<sub>F</sub> displayed a degree of colocalization ( $63 \pm 36\%$ , N = 4,744) larger than random (20%, Fig. 4b and Supplementary Figs 6A and 7A), suggesting that partition complexes may interact with high-density chromosomal DNA regions during segregation and persist throughout the cell cycle (Supplementary Fig. 7C). Next, we simultaneously imaged ParA<sub>F</sub>-GFP and HUlabelled nucleoids at super resolution. Interestingly, overlays of HDRs with ParA<sub>F</sub> exhibited a large degree of positional correlation, even higher than that observed for partition complexes ( $82 \pm 24\%$ , N = 9,553; Fig. 4c and Supplementary Figs 6B and 7B). ParA and ParBS form dynamic complexes; thus, slight differences in position between HDRs/ParA-ParBS may arise from movement of complexes between acquisitions (  $\sim 20$  s).

To investigate whether localization of  $ParA_F$  to HDRs influenced the process of segregation, we followed the dynamic movement of single  $ParA_F$  molecules in live cells using single-particle tracking PALM (spt-PALM)<sup>33,34</sup>. Wild-type and  $ParA_F$  mutants were labelled with the photo-activatable fluorescent protein mEos2. Histograms of the mean-squared displacement from individual trajectories of wild-type  $ParA_F$  revealed two distinct species: a dynamic population that explored large regions of the nucleoid and a static species that localized to discrete patches (Fig. 4d,e). The diffusion coefficients of these species were consistent with previous observations of DNA-associated proteins and free proteins<sup>35</sup>. The number of static ParA patches was similar to the number of ParA patches colocalizing with HDRs observed by 3D-SIM (compare panels c and e of Fig. 4). Strikingly, for the ParA mutant with reduced ns-DNA-binding capacity (ParA<sub>F</sub><sup>X340A</sup>), the static population became negligible and the localization of ParA was homogeneous over the entire nucleoid (Fig. 4f,g). Importantly, this ParA mutation leads to abnormal positioning of partition complexes and to segregation defects (Fig. 3g–i), indicating ParA localization to HDRs is important for the volumetric localization and proper segregation of partition complexes.

Our results show that there is an apparent correlation between the 3D positions of HDRs and ParA/ParBS complexes. To determine whether this positional correlation is conserved during the cell cycle, we analysed the localizations of ParA and ParBS complexes together with those of HDRs as a function of nucleoid size, which directly correlate with the cell cycle stage<sup>36</sup>. First, we computed the probability density of localization of HDR as a function of nucleoid length and longitudinal position. Maxima in the HDR probability density represent regions where HDRs display a conserved or enriched subcellular localization (Fig. 4h). HDR maxima were localized at midcell for nascent nucleoids, and



**Figure 4 | Partition complexes segregate through dense chromosomal DNA regions. (a)** Typical 3D volume of chromosomal DNA stained with HU-mCherry. Nucleoid DNA density in the volumetric representation and side projections is colour-coded from blue to red. HDRs stretch from pole to pole and are joined by a semicontinuous filamentous density. (b) ParB<sub>F</sub> localization correlates with positions occupied by HDRs. Left: maximum intensity projections of individual nucleoids labelled by HU-mCherry (DLT3053/pJYB234). Right: maximum intensity projections of the same nucleoids but for ParB<sub>F</sub>-mVenus. Middle: 2D projection of HDRs (red) and ParB<sub>F</sub> foci (green) positions represented as 2D Gaussian distributions with sizes corresponding to the resolution of 3D-SIM for each emission channel. Note that this image represents a 2D projection, not a *z* section. HDRs colocalizing with ParB foci are shown in yellow. (c) ParA<sub>F</sub> localization correlates with positions occupied by HDRs. Same representations as in **b** but for HDRs and ParA<sub>F</sub> (DLT3053/pJYB243). A strong correlation between HDR and ParA is observed despite the nucleoid and ParAB dynamics between sequential acquisitions for each colour. (**d-f**) Distributions of the apparent diffusion coefficient, *D*, of single-molecule trajectories of wild-type ParA (DLT3284/pJYB286, *N* = 63 cells) and ParAF<sup>K340A</sup> (DLT3286/pJYB288, *N* = 17 cells). Trajectories were classified as a static species (red) and a fast diffusing dynamic species (blue). (**e-g**) Representative examples of static and dynamic trajectories are illustrated using the same colour code of **d-f** (red: static; blue: dynamic). Both panels depict one representative experiment of at least two biological replicates. (**h-j**) ParB<sub>F</sub> and ParA<sub>F</sub> choreography is concerted with HDR positions throughout the cell cycle. 2D localization density of (**h**) HDRs in *E. coli*, (**i**) ParB-mVenus and (**j**) ParA<sub>F</sub>-mVenus as a function of the cell cycle. Dashed purple lines in **i**, j provide a guide to the eye for the localization p

moved to quarter positions where they remained until the next cycle of cell division. By two-colour 3D-SIM, we simultaneously localized the positions of ParB<sub>F</sub> complexes and calculated their probability density of localization as a function of cell cycle progression. Strikingly, we observed that the segregation pattern of ParBS complexes followed closely that of HDR maxima (Fig. 4i, see overlayed dashed line for density of HDR maxima), consistent with our previous studies of B. subtilis ParB<sup>26</sup>. Both HDR and ParBS maxima in the localization density maps forked at a nucleoid size of  $\sim 2,300$  nm (Fig. 4h,i, horizontal grey lines). Finally, we performed a similar experiment to detect the mean cell cycle positioning of ParA<sub>F</sub> (Fig. 4j). ParA<sub>F</sub> localized predominantly to the edges of the nucleoid for the majority of the cell cycle, consistent with the polar localization of ParA<sub>F</sub> observed at fixed nucleoid sizes (Fig. 2c). In the localization density map, ParA<sub>F</sub> maxima forked earlier than HDRs/ParBS  $(\sim 1,800 \text{ nm}, \text{ horizontal grey line in Fig. 4j})$  at a nucleoid size matching the dwindling of chromosomal DNA density at the centre of the nucleoid (Fig. 4h). These findings are not inconsistent with the colocalization between HDRs and ParA patches at the single cell level, as kymograph maxima highlight a preferential positioning of HDRs/Par proteins within the nucleoid volume in a population. Interestingly, perturbation of HDRs using rifampicin treatment, an antibiotic known to block

transcription and to affect chromosome structure, led to a spreading of HDRs positioning throughout the nucleoid volume as well as the cell cycle, and affected the positioning of both ParA and ParB in a very similar manner as that of HDRs (Supplementary Fig. 8).

#### Discussion

In this study, we investigated the mechanism of segregation of two model partition systems in bacteria by combining 3D-SIM, MFM, spt-PALM and widefield deconvolution microscopy with quantitative high-throughput analysis. Our experiments demonstrate that all components of the partition machinery are localized within the nucleoid interior during segregation, and that the DNA-binding and ATPase activity of the ParA transport protein are essential for this localization. These results call for a re-evaluation of several aspects of previously proposed models.

Two main mechanisms were proposed to account for the segregation of bacterial partition complexes: filament and brownian-ratchet-like models<sup>37</sup> (Fig. 1a). Filament models proposed that ParA functions as a nucleoid-spanning filament that either pushes or pulls partition complexes<sup>19,20</sup>. In contrast, we failed to observe extended ParA filaments using either 3D-SIM or spt-PALM. These results are compatible with recent



Figure 5 | Hitch-hiking model for bacterial chromosome segregation. The Hitch-hiking model for bacterial chromosome segregation. See text for description. Cell outline is shown as a grey mesh, nucleoid as a red cylinder, ParBS partition complexes in green, static DNA-bound ParA in cyan, free ParA-ADP in blue and HDRs as red diffuse circles.

observations in which it was shown by 2D sub-diffraction imaging that *Caulobacter crescentus* ParA does not form continuous filaments<sup>25</sup>. Overall, these results are inconsistent with extended ParA filaments or helical structures driving segregation of partition complexes<sup>1</sup>. Importantly, our ability to detect the 3D localization of ParA within the nucleoid allowed us to determine that ParA localizes specifically within the volume of the nucleoid. This volumetric localization of ParA is consistent with previous reports in other model systems<sup>20,29</sup>, suggesting that it may constitute a universal property of partition complexes. In addition, we observed by both 3D-SIM and spt-PALM that ParA forms small patches extending between nucleoid poles through the interior of the nucleoid (see model below).

On the basis of the observation of 2D diffusion of partition complexes on DNA-coated surfaces in vitro, most recent models postulated that partition complexes are segregated over the nucleoid surface via interactions with ns-DNA-bound ParA dimers or oligomers<sup>21-24</sup>. This model attributed a function to the cytosolic space between nucleoid and cell membrane because the high spatial confinement at this interface promotes frequent associations between plasmid-bound ParB and nucleoid-bound ParA<sup>21</sup>. It is worth noting, however, that general Brownianratchet-like mechanisms do not make specific statements about the spatial distribution of partition components. A variation of this model (DNA relay) proposed that, in addition to the brownian-ratchet, chromosomal DNA dynamics may also power partition complex propagation over the nucleoid surface<sup>25</sup>. These conclusions were mainly reached by simulations of diffusion of partition complexes in 2D. Here we show by several microscopy methods and for plasmid and chromosomal model systems that in vivo both partition complexes and the ParA ATPase localize predominantly within the nucleoid volume and in average to the longitudinal cell axis. Our findings are consistent with previous observations of the localization of B. subtilis ParB<sup>26</sup>, and are incompatible with partition complex segregation via surfaceactivated mechanisms requiring an enrichment of ParBS and ParA at the interface between nucleoid and cell membrane<sup>24,25</sup>.

Interestingly, our studies also reveal that the volumetric localization of ParA patches and partition complexes is highly correlated with the positions of HDRs. HDRs are present in both *B. subtilis*<sup>26</sup> and *E. coli* (Fig. 4) and represent regions where DNA is more highly compacted, probably because of the stochastic, dynamic association of chromosomal interaction domains<sup>26</sup>. The localization pattern of ParB<sub>F</sub> maxima seems to closely follow that

of HDR maxima throughout the cell cycle (Fig. 4h,i), and even upon perturbation of the nucleoid structure ParA and ParB mirror the altered localization of HDRs (Supplementary Fig. 8), consistent with ParBS complexes being constitutively located close to regions of high chromosomal DNA density. Remarkably, segregation patterns of HDR maxima and ParB are also highly correlated in B. subtilis<sup>26</sup>, despite the different cell cycle localization patterns of ParB in these species (Supplementary Fig. 9). However, in B. subtilis, ParB patterns do not follow HDR maxima as closely as those reported in E. coli, which could be explained by important differences in the genomic and spatial sizes of partition complexes of Par systems from these two species. In E. coli, replication origins are located at the centre of new cells and segregate to quarter positions concomitantly with replication $^{38-42}$ . This precise choreography closely resembles that of HDR maxima, and suggests that they may correspond to replication origin regions as in B. subtilis<sup>26</sup>. Overall, these observations suggest that the subcellular localization of partition complexes may also be influenced by the host replication/segregation machinery.

In E. coli, HDRs occupy the nucleoid volume and are more likely found close to the longitudinal nucleoid axis (Fig. 4a-c and Supplementary Fig. 9)<sup>26</sup>. ATP-bound ParA interacts with ns-DNA and with itself<sup>43,44</sup>. Thus, these two activities and the volumetric localization of HDRs are consistent with the observation that ParA forms patches localizing at a discrete number of positions within the nucleoid volume. This may serve for the spatial concentration of the cellular pool of ParA into a small number of locations distributed inside the nucleoid, as suggested by the high degree of  $ParA_F$  colocalization with HDRs (>80%; Fig. 4c and Supplementary Fig. 7B) and the absence of ParA patches when its nonspecific interaction with DNA is impaired (Fig. 4f,g). In contrast to ParB<sub>F</sub>, ParA<sub>F</sub> maxima are located at the edges of the nucleoid, and split when chromosomal DNA density at the centre of the nucleoid begins to dwindle (Fig. 4j), consistent with the oscillatory behaviour of ParA<sub>F</sub> (ref. 45). Thus, the segregation pattern of ParA maxima seems in part to define the subcellular localization of ParBS complexes, and conversely ParA appears depleted in regions of high ParBS density. These observations are consistent with partition complexes dynamically following ParA concentration gradients<sup>29</sup>, triggering ParA dissociation by direct interaction with ParB22.

In absence of  $ParA_F$ , partition complexes displayed a very dynamic behaviour and lost their volumetric localization pattern. These results indicate that ParA directly participates in the

tethering of partition complexes to the nucleoid interior. Importantly, the ability of  $ParA_F$  to bind ns-DNA and to hydrolyse ATP through  $ParB_F$  stimulation were the key for the maintenance of partition complexes within the nucleoid volume (Fig. 3). In these mutants, partition complexes were excluded from the nuclear volume, likely because of impaired formation of ParA concentration gradients<sup>46</sup> and localized to the space between nucleoid and cell membrane mostly at cell poles. These findings are in good agreement with previous observations of higher intracellular mobility of *par* minus plasmids<sup>47</sup> and with nucleotide-bound state of ParA and ns-DNA binding playing a role in the longitudinal positioning of partition complexes<sup>1</sup>.

Thus, we favour a mechanism ('hitch-hiking' model) in which partition complexes are segregated by being recruited to high-density regions within the nucleoid by interactions mediated by the ParA ATPase. In this model (Fig. 5),

- (1) ParA assembles in small patches at regions of high DNA density. This could arise by the ns-DNA-binding activity of ParA and the local high concentration of DNA within HDRs. Dissociation of ParA from ns-DNA requires either ATP hydrolysis<sup>44</sup> or direct interaction with ParB<sup>22</sup>; thus, dissociation of short ParA oligomers is prevented when ParA is at HDRs. Without the presence in the close proximity of ParB or partition complexes, these ParA oligomers are relatively stable.
- (2) Partition complexes are most likely formed by nucleation and caging of hundreds of ParB dimers around a cluster of *parS* sites into small complexes (<150 nm, Supplementary Fig. 2)<sup>13</sup>. ParB and *parS*-bound ParB strongly stimulate the ATPase activity of ns-DNA-bound ParA complexes (40- and 120 fold, respectively<sup>18</sup>), leading to their dissociation from DNA<sup>24,37</sup>. Thus, interactions of ParB with HDR-bound ParA patches may trigger their progressive dissociation from HDRs, and subsequently release partition complexes from HDRs (Fig. 5, left panel). Unbound/ADP-bound ParA dimers can then rapidly diffuse, reload ATP to become competent for ns-DNA binding (after a time delay<sup>24</sup>) and preferentially oligomerize at new HDRs (Fig. 5, right panel).
- (3) This 'scanning' allows ParA oligomers to be up to date with changes in the conformation of the nucleoid, and to hitchhike on HDRs.
- (4) Interactions of ParB with ns-DNA-bound ParA are dynamic and could lead to a bias in the diffusion of partition complexes towards HDR-bound ParA patches. The radial concentration of ParA from the cytoplasm towards the longitudinal axis of the nucleoid, where DNA/ParA is more concentrated, could drive the partition complex within the nucleoid volume, while local gradients in ParA along the longitudinal axis of the nucleoid temporarily hold it in place. Once ParB has depleted local ParA in its vicinity, it can diffuse to the most proximal HDR enriched in ParA, hence moving along the heterogeneous distribution of ParA. The confinement diameter of  $\sim 200 \text{ nm}$  of ParB foci obtained from tracking experiments indicates that the area explored by partition complexes is comparable to the distance between ParA patches  $(248 \pm 74 \text{ nm}, \text{ see Supplementary})$ Fig. 10). Thus, diffusion should allow partition complexes to bridge two ParA patches without stalling. In brief, partition complex movement is composed of a brownian diffusion component and a directional bias, the essential constituents of a brownian-ratchet (Fig. 5).

Recently, we observed that the *B. subtilis* chromosome dynamically condenses/de-condenses during its replication cycle. Decondensation of the origin domain (containing most

chromosomal *parS* sites) seems to occur concomitantly with replication initiation and with the relocalization of the origin from nucleoid pole to the middle of the new nucleoid. This relocalization is accompanied by a global change in replichore organization from a longitudinal to a transverse orientation<sup>26,27</sup>. Interestingly, a similar change in replichore organization was recently proposed for *E. coli*<sup>42</sup>. Thus, origin decondensation upon replication may lead to the preferential relocalization of ParA at HDRs other than the origin. We hypothesize that newly replicated origin regions may thus follow the bias in ParA localizations. Future experiments will need to be performed to test these hypotheses.

Members of the ParA/MinD family are involved in the subcellular positioning of a large number of protein machineries with a diverse variety of functions, including the conjugative transfer and type IV pili apparati, cellulose synthesis devices and cytoplasmic chemotaxis clusters<sup>2,48</sup>. Presumably, because of the nucleoid acting as a diffusion barrier and/or because of nucleoid exclusion forces, such large complexes have been assumed to occupy the cytosolic space or to use the inner bacterial membrane as a scaffold. Our findings reinforce the need of volumetric superresolution imaging of subcellular structures and machineries on the cellular context to determine the role of subcellular ultrastructures. such as the nucleoid or the cell membrane, in the process of transport. It remains to be demonstrated whether, as it occurs for the F-plasmid and B. subtilis chromosome, such transport systems employ the scaffold of the nucleoid and may thus co-opt other cellular machineries for the intracellular transport of large cargoes. We speculate that volumetric patterning of ParAs may have been evolutionarily advantageous in bacteria and archaea for a variety of biological functions to serve as molecular beacons for positioning and/or segregating all kinds of cargoes.

#### Methods

**Bacterial strain and plasmids and growth conditions.** Strains, plasmids and growth conditions are described in Supplementary Methods. For microscopy and plasmid stability assays, cultures were grown at 30 °C with aeration in a supplemented M9 minimal medium (MGC; Supplementary Methods). A full list of strains employed in this work is given in Supplementary Methods.

**Fluorescence microscopy**. 3D-SIM imaging was performed on an OMX V3 microscope (Applied Precision), using 405, 488 and 568 nm laser lines to excite DAPI, mVenus and mCherry respectively<sup>26,49</sup>. To perform dual colour volumetric imaging, 15-17 slices were acquired sequentially for each colour with a Z step of 125 nm between each slice. Reconstruction and alignment of 3D-SIM images was performed using softWoRx v5.0 (Applied Precision). Widefield deconvolution imaging was conducted on the same experimental set-up and used the same channel alignment procedure and algorithm. Refer to Supplementary Methods for more details.

Single-particle tracking PALM and MFM were conducted on home-built experimental set-ups based on Zeiss Axiovert 200 microscopes. Volumetric acquisitions for MFM imaging were permitted by inserting custom optical elements into the emission pathway between the microscope tube lens and the EM-CCD camera (DU-897; Andor). More details on the experimental set-ups can be found in the Supplementary Methods.

**Data analysis.** Data analysis was performed as described in Supplementary Methods and elsewhere<sup>26</sup>. Briefly, cells and nucleoids were automatically segmented using a 3D maximum entropy thresholding algorithm for binarization. The same methods were used to detect minimum and maximum surface/volume values for cells and nucleoids, respectively. Analysis of nucleoid length, width and height used the FWHM of the nucleoid long and short axes. Nucleoid dimensions were then used to draw the nucleoid contour in density histogram representations. ParB foci in each cell were detected as 3D local maxima, using a multidirectional derivation of the intensity scalar vector, and their positions were normalized by the nucleoid dimensions. Nucleoids depicted in figures were segmented by visual impression for 3D rendering of nucleoid surfaces (Figs 1d,e,2a,b,3b,e,h and 4a). 2D histograms of the positions of the ParB foci in the nucleoid were constructed with nucleoids of length <1,500 nm. The selected nucleoids were resampled 100 times using a bootstrapping method, and foci position orientations were randomized along the long, short and vertical axes of the nucleoids before computing 2D

histograms. 2D histograms were generated by computing the 2D spatial distribution of number of local maxima weighted by their corresponding local voxel intensity in the 3D-SIM stack. The mean histogram was then calculated from the 100 smoothed histograms built from the 100 bootstrapped samples. Note that for YZ histogram construction, solely ParB foci whose longitudinal coordinates lie within 60% of the nucleoid length were considered in order to avoid the contribution from ParB foci at the nucleoid poles. Histograms of ParA distribution within the nucleoid were computed the same way but ParA foci/patch orientation along the nucleoid long axis was set to keep the brightest patches on one side. spt-PALM data analysis is described in Supplementary Methods.

**Data availability**. The data that support the findings of this study are available from the corresponding author upon request.

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#### Author contributions

A.L.G., D.I.C., J.-Y.B. and M.N. conceived the idea, designed the research and discussed the data; A.L.G., D.I.C., J.-Y.B. and M.N. wrote the paper; A.L.G. and D.I.C. performed the experiments, A.L.G. wrote analysis routines and analysed the data; L.O., J.-B.F., S.A., A.L.G. and M.N. designed and performed MFM experiments; B.G., J.B.F. and M.N. designed and performed spt-PALM experiments and analysis; J.R., C.M.-D., H.M. and J.Y.B. constructed cell lines.

#### Additional information

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