Robust and conserved stochastic self-assembly mechanism for dynamic ParB-parS partition complexes on bacterial chromosomes and plasmids

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Running head: Conserved stochastic self-assembly of partition complexes
Summary

Chromosome and plasmid segregation in bacteria are mostly driven by ParABS systems. These DNA partitioning machineries rely on large nucleoprotein complexes assembled on centromere sites (parS). However, the mechanism of how a few parS-bound ParB proteins nucleate the formation of highly concentrated ParB clusters remains unclear despite several proposed physico-mathematical models. We discriminated between these different models by varying some key parameters \textit{in vivo} using the plasmid F partition system. We found that ‘Nucleation & caging’ is the only coherent model recapitulating \textit{in vivo} data. We also showed that the stochastic self-assembly of partition complexes (i) does not directly involve ParA, (ii) results in a dynamic structure of discrete size independent of ParB concentration, and (iii) is not perturbed by active transcription but is by protein complexes. We refined the ‘Nucleation & Caging’ model and successfully applied it to the chromosomally-encoded Par system of \textit{Vibrio cholerae}, indicating that this stochastic self-assembly mechanism is widely conserved from plasmids to chromosomes.
Introduction

The segregation of DNA is an essential process for the faithful inheritance of genetic material. Minimalistic active partition systems, termed Par, ensure this key cell cycle step in bacteria (Baxter and Funnell, 2014) and archaea (Schumacher et al., 2015). Three main types of bacterial partition systems have been identified and classified by their NTPase signatures. Of these, the type I, also called ParABS, is the only one present on chromosomes and the most widespread on low-copy number plasmids (Gerdes et al., 2000). Each replicon encodes its own ParABS system and their proper intracellular positioning depends on the interactions of the three ParABS components: ParA, a Walker A cytoskeletal ATPase; ParB, a dimer DNA binding protein; and parS, a centromere-like DNA sequence that ParB binds specifically. The ParA-driven mechanism that ensures the proper location and the directed segregation of replicons relies on the positioning of ParBS partition complexes within the nucleoid volume (Le Gall et al., 2016) and on a reaction diffusion-based mechanism (Hu et al., 2017; Hwang et al., 2013; Lim et al., 2014; Walter et al., 2017).

The centromere-like parS sites are located close to the replication origin on chromosomes and plasmids, and are typically composed of 16-bp palindromic motifs (Lin and Grossman, 1998; Mori et al., 1986). ParB binds with high affinity to its cognate parS as dimers (Bouet et al., 2000; Hanai et al., 1996). This serves as a nucleation point for assembling high molecular weight ParB-parS partition complexes, as initially seen by the silencing of genes present in the vicinity of parS (Lobocka and Yarmolinsky, 1996; Lynch and Wang, 1995). ParB binds over 10-Kbp away from parS sites for all ParABS systems studied to date (Donczew et al., 2016; Lagage et al., 2016; Murray et al., 2006; Rodionov et al., 1999; Sanchez et al., 2015). This phenomenon, termed spreading, refers to the binding of ParB to centromere-flanking DNA regions in a non-specific manner. The propagation of ParB on DNA adjacent to parS is blocked by nucleoprotein complexes such as replication initiator complexes in the case of the
P1 and F plasmids (Rodionov et al., 1999; Sanchez et al., 2015), or repressor-operator complexes on the bacterial chromosome (Murray et al., 2006). These ‘roadblock’ effects led to the initial proposal that ParB propagates uni-dimensionally on both sides of the parS sites, in a so-called ‘1D-spreading’ model. However, this model was put into question as (i) the quantity of ParB dimers present in the cell was insufficient to continuously cover the observed spreading zone, and (ii) ParB binding to parS adjacent DNA resisted biochemical demonstration (reviewed in Funnell, 2016).

As an alternative to ‘1D-spreading’, two other models for partition complex assembly have been proposed, namely ‘Spreading & bridging’ (Broedersz et al., 2014) and ‘Nucleation & caging’ (Sanchez et al., 2015). Both models rely on strong ParB clustering with over 90% of ParB confined around parS (Sanchez et al., 2015). The ‘Spreading & bridging’ model proposes that nearest neighbour interactions (1D-spreading) initiated at parS and non-parS DNA sites in combination with their subsequent interactions in space (3D-bridging), lead in one of the conditions tested (strong spreading and bridging) to the condensation of the ParB-bound DNA into a large 3D complex over a contiguous 1D DNA domain (Broedersz et al., 2014; Graham et al., 2014). The ‘Nucleation & caging’ model rather proposes that the combination of dynamic but synergistic interactions, ParB-ParB and ParB-nsDNA (Fisher et al., 2017; Sanchez et al., 2015), clusters most of the ParB around parS nucleation sites where a few ParB dimers are stably bound (Fig. 1A). The in vivo ParB binding pattern from high resolution ChIP-sequencing data was described with an asymptotic decay as a characteristic power-law with an exponent b= -3/2, corresponding to the decreasing probability of the DNA to interact with the ParB cluster as a function of the genomic distance from parS (Sanchez et al., 2015). This model therefore proposes that the DNA surrounding the parS site interacts stochastically with the sphere of high ParB concentration. Interestingly, these three different
assembly mechanisms have been explicitly modelled (Broedersz et al., 2014; Sanchez et al., 2015), thus allowing their predictions to be experimentally tested.

To study the assembly mechanism of partition complexes, we used the archetypical type I partition system of the plasmid F from *E. coli*. By varying several key parameters, we evaluated ParB binding patterns *in vivo* in relation to predictions of each model. We also investigated the chromosomal ParABS system of the main chromosome of *Vibrio cholerae*. In all tested conditions, our data indicate that ParB binding profiles robustly correlate only with the predictions of the ‘Nucleation & caging’ model.

Results

ParB\textsubscript{F} distribution pattern around *parS*\textsubscript{F} is similar on chromosome and plasmid DNA

The plasmid F partition complex assembles on a centromere sequence, *parS*\textsubscript{F}, composed of twelve 43-bp tandem repeats (Helsberg and Eichenlaub, 1986), which contain ten 16-bp inverted repeat motifs to which ParB\textsubscript{F} binds specifically *in vitro* (Pillet et al., 2011) and *in vivo* (Sanchez et al., 2015). Partition complex assembly has been investigated using small versions of the plasmid F, either ~10- or ~60-Kbp. To discriminate between the different partition complex assembly models, we used two larger DNA molecules: the native 100-Kbp plasmid F (F1-10B; Table S1) and the 4.6-Mbp *E.coli* chromosome with *parS*\textsubscript{F} inserted at the *xylE* locus, in strains either expressing (DLT1472) or not (DLT1215) ParB\textsubscript{F} from an IPTG-inducible promoter.

We first controlled the formation of ParB\textsubscript{F} clusters on these two different DNA molecules using the ParB\textsubscript{F}-mVenus fluorescent fusion protein. ParB\textsubscript{F}-mVenus, fully functional in plasmid partitioning (Supplemental Table S2), was expressed from the endogenous locus on the plasmid F (F1-10B-BmV) or from a low-copy number plasmid under the control of an IPTG-inducible promoter (pJYB294). In both cases, we observed bright and compact foci in
nearly all cells (Fig. 1B and D), indicating that the assembly of highly concentrated ParB<sub>F</sub>
clusters on <i>parS<sub>F</sub></i> from large DNA molecules, plasmid or chromosome, occurs similar to the
smaller plasmid F counterparts (Sanchez et al., 2015). The number of foci from <i>parS<sub>F</sub></i> inserted
on the chromosome is half of what is observed with the plasmid F, as expected from the two-
fold difference in copy-number (Collins and Pritchard, 1973).

We then performed ChIP-sequencing using anti-ParB antibodies and compared the ParB<sub>F</sub>
patterns from the 100-Kbp F1-10B plasmid and the <i>xylE::parS<sub>F</sub></i> chromosome insertion. For
F1-10B, we observed a ParB binding pattern extending over 18-Kbp of <i>parS<sub>F</sub></i>-flanking DNA
nearly identical to the one previously observed on the 60-Kbp plasmid F (Sanchez et al.,
2015), with the asymmetrical distribution arising from RepE nucleoprotein complexes formed
on the left side of <i>parS<sub>F</sub></i> on incC and ori<sub>2</sub> iterons (Fig. 1C). When <i>parS<sub>F</sub></i> is present on the
chromosome, the ParB<sub>F</sub> binding pattern displays a comparable enrichment of <i>xylE::parS<sub>F</sub>-</i>
flanking DNA over 15-Kbp (Fig. 1E). The ParB<sub>F</sub> distribution extends ~9- and 6-Kbp on the
right and left sides of <i>parS<sub>F</sub></i>, respectively. The asymmetry does not depend on <i>parS<sub>F</sub></i>
orientation as an identical ParB<sub>F</sub> binding pattern was observed with <i>parS<sub>F</sub></i> inserted in the
reversed orientation (<i>xylE::parS<sub>F</sub>-rev</i>, Fig. S1B-C). On the left side, ParB<sub>F</sub> binding ends near
the <i>yjbE</i> locus that harbors two promoters (locus A; Fig. 1E, inset and S1A). On the right side,
ParB<sub>F</sub> binding ends at the <i>yjbI</i> gene locus (locus E; Fig. 1E and S1A). A dip in the ParB
binding intensity is also observed ~1-Kbp after <i>parS<sub>F</sub></i> spanning ~300-bp, corresponding to a
promoter region (locus C; Fig. 1E and S1A). Dips and peaks in this ParB<sub>F</sub> binding pattern are
different in terms of position and intensity when compared to the one present on the plasmid
F. Overall, these data clearly indicate that the global ParB<sub>F</sub> binding distribution around <i>parS<sub>F</sub></i>
depends neither on the size nor the DNA molecule, plasmid or chromosome, and that the
ParB<sub>F</sub> binding probability is dependent on the local constraints of each given locus.
The ‘Nucleation & caging’ binding model describes the partition complex assembly from the nucleation point to large genomic distance.

Based on a smaller version of the plasmid F, we previously proposed the ‘Nucleation & caging’ model describing ParB stochastic binding at large distance (>100-bp) from parS due to DNA looping back into the confined ParB cluster. The characteristic asymptotic decay as a power-law with the exponent $b=-3/2$ is also observed with 100-Kbp plasmid F (Fig. 1C) and with parS$_F$-inserted on the E. coli chromosome (Fig. 1E and Fig. S1C). This property is thus an intrinsic parameter of the ParBF binding profile at distance >100-bp from parS$_F$. The abrupt initial drop in ParBF binding at a shorter genomic distance (<100-bp) from parS$_F$ is explained by the difference of ParBF binding affinities between specific parS$_F$ sites (~2 nM) and non-specific DNA (~300 nM) (Ah-Seng et al., 2009). To take into account this initial drop, we now considered explicitly these different binding affinities: the amplitude of the drop, $\exp(\varepsilon_{ns} - \varepsilon_s)$, is given by the ratio of the Boltzmann weights between specific ($\varepsilon_s$) and non-specific ($\varepsilon_{ns}$) binding energies (in units of $kT$). The ParB density was normalized to 1 by the value on the right side of parS, and captured in the following formula:

$$ P_{SB}(s) = \exp(\varepsilon_{ns} - \varepsilon_s) \int dr P(r, s) C(r) $$

where $P(r, s) = s^{-3/2} e^{-3r^2/2R(s)^2}$ is the probability for two DNA loci spaced by a genomic distance $s$ to be at a distance $r$ in space for a Gaussian polymer; $R(s) = a\sqrt{L}$ is the equilibrium size of DNA with linear length $L$ and $a$ is the Kuhn length of the DNA molecule (about twice the persistence length of the corresponding Worm-like chain; (Schiessel, 2013)); $C(r) = C_0 e^{-r^2/2\sigma^2}$ is the density of ParB at a radial distance $r$ from the centromere, with $C_0$ the concentration at the origin of the cluster and $\sigma$ the typical size of the cluster. Note that $C(r)\exp(\varepsilon_{ns} - \varepsilon_s)$ is the linearized form of the Langmuir model (Phillips et al., 2012) offering a more compact and intuitive expression for $P(s)$. From (1) we easily calculate:
\[ P_{SB}(s) = \exp(\epsilon_{ns} - \epsilon_s)(s + \sigma^2/a)^{-3/2} \]

where the decay is asymptotically determined by a power law of exponent -3/2 modulated by an amplitude depending on the concentration and non-specific affinity of ParB. Two of the three parameters of this model are obtained from experiments: \( \sigma = 75 \, \text{nm} \) is determined from superresolution microscopy (Lim et al., 2014; Sanchez et al., 2015) and \( \epsilon_{ns} - \epsilon_s = -0.9 \) is read directly from the ParB density at the nsDNA binding site after \textit{parS} sequence. Note that \( \epsilon_{ns} - \epsilon_s \) estimate depends on the bioinformatics analysis (Fig. S1D). The only remaining free parameter is the Kuhn length \( a \), set at 10- or 23-bp for the plasmid F or \textit{parS}F-chromosomal insertions, respectively, to fully describe the ParBF DNA binding profiles (Fig. 1C, E and Fig. S1C). These fitted values are lower than expected, likely due to the modeling that does not account for supercoiling. Nevertheless, using these defined parameters, the refined ‘Nucleation & caging’ model provides a qualitative prediction of the experimental data over the whole range of genomic positions, from a few bp to more than 10-Kbp.

**ParBF DNA binding pattern over a wide range of ParB concentrations favors the ‘Nucleation & caging’ model**

The physical modeling for each proposed model (Broedersz et al., 2014; Sanchez et al., 2015) predicts distinct and characteristic responses upon variation of the intracellular ParB concentration (see explanations in Fig. S2A). Briefly, (i) the ‘1-D filament’ model predicts a rapid decrease of ParB binding followed by a constant binding profile dependent on ParB amount, (ii) the ‘Spreading & bridging’ model predicts linear decays with slopes depending on the ParB amount, and (iii) the ‘Nucleation & caging’ model predicts a binding profile which depends only on the size of the foci. The exponent \( b = -3/2 \) of the power-law distribution would not change upon ParB amount variation resulting in an overall similar decay. In order to discriminate between these three model predictions, we performed ChIP-seq experiments
over a large range of intracellular ParB concentrations. To prevent interference with plasmid stability, we used the chromosomally encoded xylE::parSF construct expressing parBF under the control of an IPTG inducible promoter (DLT2075).

Without IPTG induction, ParBF was expressed at ~0.2 of the physiological concentration from plasmid F, as judged by Western blot analyses (Fig. S2B). We also tested an 8- and 14-fold overproduction of ParBF. Assuming the two-fold difference in copy number (Fig. 1B and 1D), these three conditions provided ParBF/parSF ratios of 0.4, 16 and 28, relative to the plasmid F one. At these three ratios, ChIP-seq data revealed that ParBF binding extended similarly over ~15-Kbp around parSF. We analyzed the right side of parSF displaying the longest propagation distance by normalizing each data set (Fig. 2A). It revealed that regardless of ParB concentration (i) the ParB distribution in the vicinity of parSF always displays a good correlation with a power law fitting with an exponent of -3/2, (ii) the ParB binding profile ends at the same genomic location, i.e. 9-kpb from parSF and (iii) the dips and peaks in the pattern are highly conserved. This indicates a highly robust ParB binding pattern that is invariant over a ~70-fold variation of the ParB amount.

To further vary the amount of ParBF available for partition complex assembly, high-copy number (HCN) plasmids containing the parSF sequence were introduced into the xylE::parSF strain to efficiently titrate ParBF by its binding to the excess of specific binding sites (~200- and ~500-fold on pBR322 and pBSKS derivatives, respectively; (Diaz et al., 2015)). Epifluorescence microscopy of these strains reveals that all cells display a diffuse ParB-mVenus fluorescence (Fig. 2B) in contrast to concise foci without titration (Fig. 1A), suggesting a large reduction of ParB availability to non-specific sites in the vicinity of parSF on the chromosome. ChIP-seq analyses in the two titration conditions revealed that ParB binding in the vicinity of parSF was dramatically reduced as expected. However, rescaling the signals by a factor of 10 and 50 for the pBR322 and pBSKS parSF-carrying derivatives,
corresponding to a ParB$_F$/parS$_F$ ratio of 0.04 and 0.016, respectively, revealed a ParB$_F$ binding pattern above the background level (Fig. 2B, inset). In both datasets, ParB$_F$ binding decreases progressively over about the same genomic distance and with a similar power law decay as without titration. Moreover, even with these very low amounts of available ParB$_F$, the dips and peaks in the profiles are present at the same positions.

The invariance of the overall ParB profile over three orders of magnitude of ParB concentration (Fig. 2B, inset) excludes the predictions of both the ‘1-D filament’ and the ‘Spreading & bridging’ models (Fig. S2A). In addition, the conservation in the positions of the dips and peaks indicate that the probability of ParB$_F$ binding at a given location is also not dependent on the amount of ParB$_F$ in the clusters. These results are strongly in favor of the refined ‘Nucleation & caging’ model presented above.

**The size of the dynamic ParB/parS cluster is independent of ParB intracellular concentration**

In all of the ParB induction levels tested, the genomic distance over which ParB$_F$ binds around parS$_F$ is constant and displays a very similar decay (Fig. 2A). This conserved binding behavior could provide information on the cluster size as a function of ParB amount. Indeed, the ‘Nucleation & caging’ model predicts a probability $P(s) \sim (s+C)^{-3/2}$ of ParB binding at a genomic distance $s$, where the constant $C = \sigma^2/a$ is function of the average radius of the foci $\sigma$ and the Kuhn length of the DNA $a$. Thus, the $P(s)$ decay is entirely determined by the geometry of the foci and the intrinsic flexibility of the DNA. Varying the ParB amount could lead to two situations: (i) the density of ParB, but not $\sigma$, is constant (ii) $\sigma$ is fixed and ParB density is variable. We plotted these two situations in the range of ParB/parS ratio considered experimentally (Fig. 2C): with (i), the different $P(s)$ strongly varied, and (ii), $P(s)$ was invariant relative to the ParB amount resulting in overlapping profiles. Experimental data
(Fig. 2A) are in excellent agreement with the latter. From this modeling, we thus concluded that the size of partition complexes is invariant to change in ParB intracellular concentration.

The arginine rich motif (box II) of ParB<sub>F</sub> is critical for partition complex assembly

The ability of ParB to multimerize through dimer-dimer interactions is required for the formation of ParB clusters. A highly-conserved patch of arginine residues present in the N-terminal domain of ParB (box II motif; Yamaichi and Niki, 2000) has been proposed to be involved in ParB multimerization (Breier and Grossman, 2007; Song et al., 2017). To examine to what extent the box II motif is involved in vivo in the assembly of ParB<sub>F</sub> clusters, we changed three arginine residues to alanine (Fig. S3A). The resulting ParB<sub>F</sub>-3R* variant was purified and assayed for DNA binding activity by electro-mobility shift assay (EMSA) in the presence of competitor DNA using a DNA probe containing a single parS<sub>F</sub> site (Fig. 3A). ParB<sub>F</sub>-3R* binds parS<sub>F</sub> with high affinity (B1 complex) indicating no defect in (i) protein folding, (ii) parS<sub>F</sub> binding and (iii) dimerization, a property required for parS binding (Hanai et al., 1996). However, in contrast to WT ParB, the formation of secondary complexes (B’2 and B’3), proposed to result from ParB multimerization (Sanchez et al., 2015), was impaired further suggesting the implication of box II in dimer-dimer interaction. A mini-F carrying the parB<sub>F</sub>-3R* allele (pAS30) was lost at a rate corresponding to random distribution at cell division (Table S2), indicating that this variant is unable to properly segregate the mini-F.

The ParB<sub>F</sub>-3R* variant was then expressed in native or fluorescently-tagged (ParB -R3*-mVenus) forms, from pJYB303 or pJYB296, respectively, in the xylE::parS<sub>F</sub> strain. By imaging ParB<sub>F</sub>-3R*-mVenus, we observed only faint foci in a high background of diffuse fluorescence (Fig. 3B). These barely detectable foci may correspond to ParB<sub>F</sub>-3R*-mVenus binding to the ten specific sites present on parS<sub>F</sub> and, if any, to residual ParB<sub>F</sub> cluster formation. We then performed ChIP-seq assays with ParB<sub>F</sub>-3R* present in ~25-fold excess
(relative \textit{ParBF}/par\textit{S}F ratio compared to the plasmid F one; Fig. S3B). The resulting DNA binding profile displayed enrichment only at \textit{parS}F with a total absence of \textit{ParBF} binding on \textit{parS}F-flanking DNA (Fig. 3C). This pattern differs from those observed in conditions of \textit{ParBF} titration (Fig. 2A; inset), indicating that the \textit{ParBF}-3R* box II variant is fully deficient in clustering \textit{in vivo}. The same pattern was also observed with \textit{ParBF}-3R*-mVenus (Fig. S3C) indicating that the mVenus fluorescent-tag fused to \textit{ParBF} does not promote cluster assembly.

Together, these results indicate that the box II variant is specifically deficient in \textit{ParBF} cluster assembly but not in \textit{parS}F binding, and thus reveal that the box II motif is critical for the auto-assembly of the partition complex.

**ParB also propagates stochastically from native chromosomal \textit{parS} sites**

\textit{ParABS} systems are present on most bacterial chromosomes (Gerdes et al., 2000). To determine whether chromosomal \textit{ParB-parS} partition complexes also assembled \textit{in vivo} in a similar manner to the plasmid F, we investigated the bacterium \textit{Vibrio cholerae}, whose genome is composed of two chromosomes. We focused on the largest chromosome to which \textit{ParB}\textsubscript{Vc1} binds to three separated 16-bp \textit{parS} sites comprised within 7-Kbp (Baek et al., 2014; Saint-Dic et al., 2006) (Fig. 4A).

We purified \textit{ParB}\textsubscript{Vc1} antibodies against his-tagged \textit{ParB}\textsubscript{Vc1} and performed ChIP-seq assays on exponentially growing cultures. The \textit{ParB}\textsubscript{Vc1} DNA binding pattern covered \~18-Kbp and displayed three peaks at the exact location of the three \textit{parS}\textsubscript{Vc1} sites (Fig. 4B). Each peak exhibits a distinct but reproducible difference in intensity that might correspond to the slight differences in \textit{parS}\textsubscript{Vc1} sequences (Fig. S4A). An asymmetry in the binding pattern was observed on the left side of \textit{parS}1 with the limit of \textit{ParB}\textsubscript{Vc1} binding corresponding to the end of the rRNA operon located \~4-Kbp upstream from \textit{parS}1 (Fig. 4B). This suggests that highly transcribed genes might significantly interfere with the extent of \textit{ParB} binding.
We modeled ParB$_{Vc1}$ DNA binding profile with the framework of the refined Stochastic Binding model (see above), considering three non-interacting spheres centered on each of the $parS$ sites (Fig. 4C). Here, $\epsilon_{ns} - \epsilon_{s1} = -0.2$, where $\epsilon_{s1}$ is the specific binding energy for $parS1$. The simulated profile was obtained by using the same protocol as performed by the bioinformatic analysis in order to account for the width of the peaks around each $parS$; the same modeling as for E. coli would led to a sharp decay between $parS$ and non-specific sites (Fig. S4B).

The maxima in the ParB binding profile depends on the $parS$ sites (Fig. 4C) and are interpreted as a difference in binding affinity. In the simulations, the ParB density is normalized to 1 by the value on the right of $parS1$. The relative density of the two other $parS$ sites is fixed according to the values read on the ChIP-seq plot (3% and 29% lower affinity for $parS2$ and $parS3$ compared to $parS1$, respectively). We found a good agreement with the ParB$_{Vc1}$ profile by applying a lower difference between the specific and non-specific binding energies than for ParB$_E$, as reported in other ParABS system (Fisher et al., 2017). We also noticed a clear difference at the minima of ParB binding on either side of $parS2$ (64.2 and 68-Kbp; Fig. S4B). In the case of a single cluster constraining the three $parS$, the profile would only depend on the genomic distance from $parS2$ resulting in a symmetrical pattern, while in the case of three independent clusters, an absence of symmetry due to the occupation of the specific sites is expected. This indicates that the system displays three independent clusters nucleated at each $parS$ sites. However, the possibility that these clusters mix together at a frequency dependent on the genomic distance between $parS$ sites is not excluded. At larger distances from $parS$ sites, differences between the experimental data and the simulation probably arise from strong impediments to ParB binding, such as the presence of the rRNA operon.

These data strongly support that the partition complex assembly mechanism is conserved on plasmid and chromosome ParABS systems.
Nucleoprotein complexes, but not active transcription, are the major determinants for the impediment of ParB stochastic binding

The major dips in the ParBF DNA binding signal are often found at promoter loci (Fig. S1A). To investigate the link between gene expression and the impediment to ParB propagation, we reproduced the ChIP-seq assays using the xylE::parS\textsubscript{F} strain grown in the presence of rifampicin, an inhibitor of RNA synthesis that traps RNA polymerases at promoters loci in an abortive complex unable to extend RNAs beyond a few nucleotides (Herring et al., 2005). We did not observe significant changes to the ParB signal on either side of parS\textsubscript{F} (Fig. 5A; compare red and blue curves). Notably, the ParB signal still strongly drops in promoter regions (e.g. loci A, C and E) and the dips and peaks are present at the same locations (Fig. 5B). This indicates that active transcription by RNA polymerase is not a major impediment to ParB binding.

We also measured the ParB binding profile in stationary phase, a growth condition in which gene expression is strongly reduced. On the right side of parS\textsubscript{F}, ParB distribution was similar to all other tested conditions (Fig. 5A), thus confirming the robustness of the binding pattern. On both sides, the strong reduction of ParB binding at loci A, C and E was still observed. However, in contrast to the other conditions, ParB binding recovers after these loci and extends up to ~18-Kbp on both sides, resulting in the location of parS\textsubscript{F} in the middle of a ~36-Kbp propagation zone. Interestingly, the ParB binding profiles after these recoveries could still be fit to a power law exhibiting the same characteristics as at lower genomic distances (Fig. 5 C). In stationary phase, the reduced intracellular dynamics (Parry et al., 2014) and the higher compaction of the DNA (Meyer and Grainger, 2013) may stabilize the partition complex revealing the ParB\textsubscript{F} bound at larger distances from parS\textsubscript{F}. Interestingly, in higher (stationary phase) or lower (rifampicin-treated cells) DNA compaction states (Fig.
S5A), the ParBF DNA binding pattern is not altered, exhibiting a similar profile of dips and peaks (Fig. 5B). This indicates that the assembly of the partition complex is not perturbed by variation in DNA compaction level within the nucleoid.

To further demonstrate the impediment of ParBF binding in promoter regions, we constructed a strain in which the locus A, carrying two promoters, an IHF and two RcsB binding sites, is replaced by a kanamycin resistance gene (Fig. 5D). The measured ParBF binding pattern remained highly comparable except at the locus A where the dip is absent. This result clearly indicates that site-specific DNA binding proteins are the main factors for restricting locally ParBF binding.

**ParB molecules exchange rapidly between partition complexes**

Single molecule *in vivo* localization experiment have shown that over 90% of ParBF molecules are present at any time in the confined clusters (Sanchez et al., 2015), suggesting that partition complexes are stable structures. However, stochastic binding of most ParBF on non-specific DNA suggests that partition complexes are highly dynamic. To reconcile this apparent discrepancy, we performed fluorescence recovery after photobleaching (FRAP) on two foci cells for measuring ParBF dynamics between partition complexes. By laser-bleaching only one focus, we could determine whether ParBF dimers could exchange between clusters and measure the exchange kinetics. As ParBF foci are mobile, we choose to partially bleach (~50%) the focus enabling immediate measurement of fluorescence recovery (Fig. 6A-B). A few seconds after bleaching, the fluorescence intensity recovers while it decreases in the unbleached focus. This exchange is progressive and the intensity between the two foci equilibrated in ~80 sec on average (between 50 and 120 sec for most individual experiments). We estimate that, when exiting a cluster, each ParBF dimer has the same probability to reach any of the two clusters. Therefore, the time of equilibration between the two foci corresponds
to the exchange of all ParB$_F$. These results thus indicate that the partition complexes are dynamic structures with a rapid exchange of ParB$_F$ molecules between clusters.

**Discussion**

Despite over three decades of biochemical and molecular studies on several ParABS systems, the mechanism of how a few ParB bound to $parS$ sites can attract hundreds of ParB in the vicinity of $parS$ to assemble a high-molecular weight complex remained puzzling. The three main mechanisms proposed for ParB-$parS$ cluster formation have been studied from physico-mathematical perspectives (Broedersz et al., 2014; Sanchez et al., 2015), predicting very different outcomes for the ParB binding profile in the vicinity of $parS$ sites upon change in ParB concentration. Here, the ParB binding patterns were found invariant over a large variation of ParB amount displaying a robust decay function as a power law with the characteristic exponent $b=−3/2$ and a conserved length of the propagation zone (Fig. 2A). Strikingly, even in the titration conditions tested, which resulted in a very low amount of ParB available to bind to non-specific DNA sites, the overall ParB DNA binding pattern remained invariant (Fig. 2A, inset). Neither ‘1-D spreading’ nor ‘Spreading & bridging’ physical models could describe these data in the conditions tested (Broedersz et al., 2014). A variant of the latter model has explored the ParB binding pattern in the low spreading strength limit (Walter et al., 2018). This ‘Looping & clustering’ model also predicts variations in the ParB binding pattern over a simulated 4-fold range of ParB amount, which is in contrast to the invariant pattern observed experimentally over more than three orders of magnitude (Fig. 2). In conclusion, only the ‘Nucleation & caging’ model based on stochastic ParB binding well describes the experimental data and provides accurate predictions for the mechanism of the partition complexes assembly.
We refined the modeling of the dynamic and stochastic ParB binding model by including DNA binding affinities for specific and non-specific sites to describe the initial drop observed immediately after parS. In this framework, we found that ParB clusters have a constant size accommodating important variations in ParB concentration (Fig. 2C). We propose that the cluster size is dependent on the intrinsic ParB-ParB and ParB-nsDNA interactions, and would thus be an inherent characteristic of each ParABS system (Funnell and Gagnier, 1993; Sanchez et al., 2015; Taylor et al., 2015). The refined modeling also well describes the chromosomal partition system of V. cholera, predicting three independent clusters nucleated at each of the three parS sites (Fig. 4C). In all cases reported here, the partition complex assembly is well described by the ‘Nucleation & caging’ model, and we propose that this mechanism of assembly is conserved on chromosome and plasmid partitioning systems.

In addition to its robustness within a large range of ParB concentration (Fig. 2A) and different nucleoid compaction states (Fig. 5A), the in vivo ParB DNA binding pattern also exhibits conserved dips and peaks at particular locations. The major dips are located at promoter regions (Fig. 1E and S1A) but do not depend on active transcription (Fig. 5B). This suggests that these specific signatures mostly depend on the intrinsic local genomic environment. This hypothesis was confirmed by deleting the locus A, carrying several regulator binding sites, which led to the suppression of the dip at this position (Fig. 5D). Therefore, proteins such as transcriptional regulators and NAPs (nucleoid associated proteins) that bind specifically to DNA prevent ParB binding to these sites, thus reducing locally the ParB signal. We propose that this impediment to ParB binding is proportional to the time of occupancy of these regulators at their site-specific DNA binding sites. Larger nucleoprotein complexes, as exemplified on the plasmid F at the iteron sites (ori2 and incC; Fig. 1C) that interact in cis and in trans (Das and Chattoraj, 2004), were previously proposed to be spatially excluded from the vicinity of the ParB cluster with a low probability that DNA beyond these
sites comes back into the cluster preventing ParB binding (Sanchez et al., 2015). Such an exclusion does not occur from smaller protein-DNA complexes, with the recovery of the ParB binding signal that further follows the characteristic power law decay (e.g. locus A; Fig. 5C). These results show that low molecular weight protein-DNA complexes do not impair the overall, only the local, ParB binding pattern.

The formation of highly concentrated clusters of ParB relies on a strong ParB-parS interaction and two other interactions, ParB-ParB and ParB-nsDNA (Fisher et al., 2017; Sanchez et al., 2015). ParB mutants that do not propagate outside parS are impaired in partition activity and in cluster formation in vivo (Breier and Grossman, 2007; Rodionov et al., 1999). The conserved box II motif (Yamaichi and Niki, 2000) was suggested to be part of the dimer-dimer interface (Breier and Grossman, 2007; Graham et al., 2014) but some misfolding caveat has been reported with some mutants, such as ParBsub-G77S (Song et al., 2017). In vivo the box II variant (ParBF-3R*) is totally deficient in partition activity and cluster formation (Fig. 3B) while proficient for parS_F binding (Fig. 3C). The total absence of ParBF-3R* binding outside parS_F (Fig. 3C and S3D) indicates that the box II motif is the major interface for the interaction between ParB dimers and is critical for the partition complexes assembly in vivo and the DNA partition activity.

ParA interacts with partition complexes in a ParB-dependent manner both in vitro and in vivo (Bouet and Funnell, 1999; Lemonnier et al., 2000) to ensure the ATP-dependent segregation of centromere sites upon DNA replication (Ah-Seng et al., 2013; Fung et al., 2001; Scholefield et al., 2011). Previous studies from V. cholerae and S. Venezuela have reported contradictory results on the involvement of ParA in the assembly of the partition complex (Baek et al., 2014; Donczew et al., 2016), which may arise from the pleiotropic effects of ParA on cellular processes, such as gene transcription or DNA replication (Murray and Errington, 2008). The ParBF DNA binding profiles on the plasmid F (Fig. 1C) and on the
*E. coli* chromosome (Fig. 1E), in the presence and absence of ParAF, respectively, are highly similar, therefore indicating that they assemble independently of ParA. Partition complexes, composed of hundreds of ParB dimers, were thought to be confined at the interface between the nucleoid and the inner membrane (Vecchiarelli et al., 2012). The observation that they rather are located within the nucleoid in a ParA-dependent manner (Le Gall et al., 2016) raises the question as to how they are not excluded from it. The ‘Nucleation & caging’ model could solve this apparent paradox. Indeed, relying on a strong ParB-parS interaction (nM range) and two other synergistic, but labile interactions, ParB-ParB and ParB-nsDNA (hundreds of nM range; Fisher et al., 2017; Sanchez et al., 2015), it would allow the dynamic confinement of most ParB without forming a rigid static structure. This dynamic organization is further supported by the finding that ParB dimers quickly exchange between clusters (~80 sec; Fig. 6). By comparison, the equilibration times between H-NS or TetR-tetO clusters were 5 or 10 times much longer, respectively (Kumar et al., 2010). Since >90% of ParB are present in clusters (Sanchez et al., 2015), it implies that their time of residency is much longer inside than outside, in agreement with fast diffusion coefficients (~1 µm².s⁻¹) for non-specific DNA binding proteins (Kumar et al., 2010). We propose that, collectively, all the individual but labile interactions for partition complex assembly allow the whole complex attracted by ParA to progress within the mesh of the nucleoid.

**Experimental procedures**

**Bacterial strains and plasmids**

*E. coli* and *V. cholerae* strains and plasmids are listed in Supplemental Table S1. Plasmids and strains constructions, growth cultures and plasmids stability assays are described in Supplemental experimental procedures.
Epifluorescence microscopy

Exponentially growing cultures were deposited on slides coated with a 1% agarose buffered solution and imaged as previously described (Diaz et al., 2015). See conditions in Supplemental experimental procedures.

ChIP-sequencing assay, analysis and fit procedure

ChIP-seq were performed as previously described (Diaz et al., 2017) with minor modifications (Supplemental experimental procedures). Graphing the DNA portion of interest from ChIP-seq data was done using Excel or R softwares. Background levels were determined by normalizing the number of sequence reads between cognate input and IP samples. Data plots superimposed with power law equation were normalized after background subtraction and set to the value of 1 at the last bp of the 10th repeat of parSf.

Western immunoblotting

The determination of ParBF relative intracellular concentrations and antibody purifications were performed as described (Diaz et al., 2015). When indicated, samples were diluted in DLT1215 extract to keep constant the total amount of proteins.

EMSA and proteins purification

EMSA were performed as described (Bouet et al., 2007) in the presence of sonicated salmon sperm DNA as competitor (100 mg.ml⁻¹), using 1 nM radiolabeled 144-bp DNA probe containing a single parSf site generated by PCR. ParBF and ParBF-3R* proteins were purified as previously described (Ah-Seng et al., 2009).

FRAP and FLIM assays
Cells, grown in mid-exponential phase, were subjected to laser-bleaching over 5-9 pixels (Supplemental experimental procedures). Normalization was performed by averaging fluorescence intensity from the three pre-bleached images.

**Accession number**

Raw ChIP-sequencing data for *V. cholera* and *E. coli* are available through the GEO repository with the accession numbers GSE114980 and GSE115274, respectively.

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**Author contributions**

Conceptualization, J.Y.B and A.P.; Methodology, J.Y.B., J.C.W., V.A.L. and A.P.; Investigation, R.E.D., J.C.W, A.S., J.R. and J.Y.B.; Formal Analysis, J.C.W., J.D., F.G., J.P.
and A.P.; Writing – Original Draft, R.E.D, J.C.W. and J.Y.B.; Writing – Review & Editing, J.Y.B., J.C.W., R.E.D and A.P.; Funding Acquisition, J.Y.B, A.P., J.P., and V.A.L.; Resources, D.L. and F.B.; Supervision, J.Y.B., V.A.L. and A.P.
References

Ah-Seng, Y., Lopez, F., Pasta, F., Lane, D., and Bouet, J.Y. (2009). Dual role of DNA in regulating ATP hydrolysis by the SopA partition protein. J Biol Chem 284, 30067-30075.

Ah-Seng, Y., Rech, J., Lane, D., and Bouet, J.Y. (2013). Defining the role of ATP hydrolysis in mitotic segregation of bacterial plasmids. PLoS genetics 9, e1003956.

Baek, J.H., Rajagopala, S.V., and Chattoraj, D.K. (2014). Chromosome segregation proteins of Vibrio cholerae as transcription regulators. MBio 5, e01061-01014.

Baxter, J.C., and Funnell, B.E. (2014). Plasmid Partition Mechanisms. Microbiology spectrum 2.

Bouet, J.Y., Ah-Seng, Y., Benmeradi, N., and Lane, D. (2007). Polymerization of SopA partition ATPase: regulation by DNA binding and SopB. Mol Microbiol 63, 468-481.

Bouet, J.Y., and Funnell, B.E. (1999). P1 ParA interacts with the P1 partition complex at parS and an ATP-ADP switch controls ParA activities. EMBO J 18, 1415-1424.

Bouet, J.Y., Surtees, J.A., and Funnell, B.E. (2000). Stoichiometry of P1 plasmid partition complexes. J Biol Chem 275, 8213-8219.

Breier, A.M., and Grossman, A.D. (2007). Whole-genome analysis of the chromosome partitioning and sporulation protein Spo0J (ParB) reveals spreading and origin-distal sites on the Bacillus subtilis chromosome. Mol Microbiol 64, 703-718.

Broedersz, C.P., Wang, X., Meir, Y., Loparo, J.J., Rudner, D.Z., and Wingreen, N.S. (2014). Condensation and localization of the partitioning protein ParB on the bacterial chromosome. Proc Natl Acad Sci U S A 111, 8809-8814.

Collins, J., and Pritchard, R.H. (1973). Relationship between chromosome replication and F'lac episome replication in Escherichia coli. J Mol Biol 78, 143-155.

Das, N., and Chattoraj, D.K. (2004). Origin pairing ('handcuffing') and unpairing in the control of P1 plasmid replication. Molecular microbiology 54, 836-849.

Diaz, R., Rech, J., and Bouet, J.Y. (2015). Imaging centromere-based incompatibilities: Insights into the mechanism of incompatibility mediated by low-copy number plasmids. Plasmid 80, 54-62.

Diaz, R.E., Sanchez, A., Anton Le Berre, V., and Bouet, J.Y. (2017). High-Resolution Chromatin Immunoprecipitation: ChIP-Sequencing. Methods in molecular biology (Clifton, NJ 1624, 61-73.)
Donczew, M., Mackiewicz, P., Wrobel, A., Flardh, K., Zakrzewska-Czerwinska, J., and Jakimowicz, D. (2016). ParA and ParB coordinate chromosome segregation with cell elongation and division during Streptomyces sporulation. Open biology 6.

Fisher, G.L., Pastrana, C.L., Higman, V.A., Koh, A., Taylor, J.A., Butterer, A., Craggs, T., Sobott, F., Murray, H., Crump, M.P., et al. (2017). The structural basis for dynamic DNA binding and bridging interactions which condense the bacterial centromere. eLife 6.

Fung, E., Bouet, J.Y., and Funnell, B.E. (2001). Probing the ATP-binding site of P1 ParA: partition and repression have different requirements for ATP binding and hydrolysis. EMBO J 20, 4901-4911.

Funnell, B.E. (2016). ParB Partition Proteins: Complex Formation and Spreading at Bacterial and Plasmid Centromeres. Frontiers in molecular biosciences 3, 44.

Funnell, B.E., and Gagnier, L. (1993). The P1 plasmid partition complex at parS: II. Analysis of ParB protein binding activity and specificity. J Biol Chem 268, 3616-3624.

Gerdes, K., Moller-Jensen, J., and Bugge Jensen, R. (2000). Plasmid and chromosome partitioning: surprises from phylogeny. Mol Microbiol 37, 455-466.

Graham, T.G., Wang, X., Song, D., Etson, C.M., van Oijen, A.M., Rudner, D.Z., and Loparo, J.J. (2014). ParB spreading requires DNA bridging. Genes Dev 28, 1228-1238.

Hanai, R., Liu, R.P., Benedetti, P., Caron, P.R., Lynch, A.S., and Wang, J.C. (1996). Molecular dissection of a protein SopB essential for Escherichia coli F plasmid partition. J Biol Chem 271, 17469-17475.

Helsberg, M., and Eichenlaub, R. (1986). Twelve 43-base-pair repeats map in a cis-acting region essential for partition of plasmid mini-F. J Bacteriol 165, 1043-1045.

Herring, C.D., Raffaelle, M., Allen, T.E., Kanin, E.I., Landick, R., Ansari, A.Z., and Palsson, B.O. (2005). Immobilization of Escherichia coli RNA polymerase and location of binding sites by use of chromatin immunoprecipitation and microarrays. J Bacteriol 187, 6166-6174.

Hu, L., Vecchiarelli, A.G., Mizuuchi, K., Neuman, K.C., and Liu, J. (2017). Brownian Ratchet Mechanism for Faithful Segregation of Low-Copy-Number Plasmids. Biophys J 112, 1489-1502.

Hwang, L.C., Vecchiarelli, A.G., Han, Y.W., Mizuuchi, M., Harada, Y., Funnell, B.E., and Mizuuchi, K. (2013). ParA-mediated plasmid partition driven by protein pattern self-organization. Embo J 32, 1238-1249.

Kumar, M., Mommer, M.S., and Sourjik, V. (2010). Mobility of cytoplasmic, membrane, and DNA-binding proteins in Escherichia coli. Biophys J 98, 552-559.
Lagage, V., Boccard, F., and Vallet-Gely, I. (2016). Regional Control of Chromosome Segregation in Pseudomonas aeruginosa. PLoS genetics 12, e1006428.

Le Gall, A., Cattoni, D.I., Guilhas, B., Mathieu-Demaziere, C., Oudjedi, L., Fiche, J.B., Rech, J., Abrahamsson, S., Murray, H., Bouet, J.Y., et al. (2016). Bacterial partition complexes segregate within the volume of the nucleoid. Nature communications 7, 12107.

Lemonnier, M., Bouet, J.Y., Libante, V., and Lane, D. (2000). Disruption of the F plasmid partition complex in vivo by partition protein SopA. Mol Microbiol 38, 493-505.

Lim, H.C., Surovtsev, I.V., Beltran, B.G., Huang, F., Bewersdorf, J., and Jacobs-Wagner, C. (2014). Evidence for a DNA-relay mechanism in ParABS-mediated chromosome segregation. eLife, e02758.

Lin, D.C.H., and Grossman, A.D. (1998). Identification and characterization of a bacterial chromosome partitioning site. Cell 92, 675-685.

Lobocka, M., and Yarmolinsky, M. (1996). P1 plasmid partition: a mutational analysis of ParB. J Mol Biol 259, 366-382.

Lynch, A.S., and Wang, J.C. (1995). SopB protein-mediated silencing of genes linked to the sopC locus of Escherichia coli F plasmid. Proc Natl Acad Sci USA 92, 1896-1900.

Meyer, A.S., and Grainger, D.C. (2013). The Escherichia coli Nucleoid in Stationary Phase. Adv Appl Microbiol 83, 69-86.

Mori, H., Kondo, A., Ohshima, A., Ogura, T., and Hiraga, S. (1986). Structure and function of the F plasmid genes essential for partitioning. J Mol Biol 192, 1-15.

Murray, H., and Errington, J. (2008). Dynamic control of the DNA replication initiation protein DnaA by Soj/ParA. Cell 135, 74-84.

Murray, H., Ferreira, H., and Errington, J. (2006). The bacterial chromosome segregation protein Spo0J spreads along DNA from parS nucleation sites. Mol Microbiol 61, 1352-1361.

Parry, B.R., Surovtsev, I.V., Cabeen, M.T., O'Hern, C.S., Dufresne, E.R., and Jacobs-Wagner, C. (2014). The bacterial cytoplasm has glass-like properties and is fluidized by metabolic activity. Cell 156, 183-194.

Phillips, R.B., Kondev, J., Theriot, J., and Garcia, H. (2012). Physical biology of the cell. Garland Science New York.

Pillet, F., Sanchez, A., Lane, D., Anton Leberre, V., and Bouet, J.Y. (2011). Centromere binding specificity in assembly of the F plasmid partition complex. Nucleic Acids Res 39, 7477-7486.

Rodionov, O., Lobocka, M., and Yarmolinsky, M. (1999). Silencing of genes flanking the P1 plasmid centromere. Science 283, 546-549.
Saint-Dic, D., Frushour, B.P., Kehrl, J.H., and Kahng, L.S. (2006). A parA homolog selectively influences positioning of the large chromosome origin in Vibrio cholerae. J Bacteriol 188, 5626-5631.
Sanchez, A., Cattoni, D.I., Walter, J.C., Rech, J., Parmeggiani, A., Nollmann, M., and Bouet, J.Y. (2015). Stochastic Self-Assembly of ParB Proteins Builds the Bacterial DNA Segregation Apparatus. Cell Syst 1, 163-173.
Schiessel, H. (2013). Biophysics for Beginners: A Journey through the Cell Nucleus. Pan Stanford Publishing.
Scholefield, G., Whiting, R., Errington, J., and Murray, H. (2011). Spo0J regulates the oligomeric state of Soj to trigger its switch from an activator to an inhibitor of DNA replication initiation. Mol Microbiol 79, 1089-1100.
Schumacher, M.A., Tonthat, N.K., Lee, J., Rodriguez-Castaneda, F.A., Chinnam, N.B., Kalliomaan-Sanford, A.K., Ng, I.W., Barge, M.T., Shaw, P.L., and Barilla, D. (2015). Structures of archaeal DNA segregation machinery reveal bacterial and eukaryotic linkages. Science 349, 1120-1124.
Song, D., Rodrigues, K., Graham, T.G., and Loparo, J.J. (2017). A network of cis and trans interactions is required for ParB spreading. Nucleic Acids Res.
Taylor, J.A., Pastrana, C.L., Butterer, A., Pernstich, C., Gwynn, E.J., Sobott, F., Moreno-Herrero, F., and Dillingham, M.S. (2015). Specific and non-specific interactions of ParB with DNA: implications for chromosome segregation. Nucleic Acids Res 43, 719-731.
Vecchiarelli, A.G., Mizuuchi, K., and Funnell, B.E. (2012). Surfing biological surfaces: exploiting the nucleoid for partition and transport in bacteria. Mol Microbiol 86, 513-523.
Walter, J.C., Dorignac, J., Lorman, V., Rech, J., Bouet, J.Y., Nollmann, M., Palmeri, J., Parmeggiani, A., and Geniet, F. (2017). Surfing on Protein Waves: Proteophoresis as a Mechanism for Bacterial Genome Partitioning. Phys Rev Lett 119, 028101.
Walter, J.C., Walliser, N.O., David, G., Dorignac, J., Geniet, F., Palmeri, J., Parmeggiani, A., Wingreen, N.S., and Broedersz, C.P. (2018). Looping and clustering model for the organization of protein-DNA complexes on the bacterial genome. New J Phys 20, 035002.
Yamaichi, Y., and Niki, H. (2000). Active segregation by the Bacillus subtilis partitioning system in Escherichia coli. Proc Natl Acad Sci U S A 97, 14656-14661.
Figure legends

Figure 1. ParB_F binding outside of parS centromere on plasmid and chromosome

(A) Schematic representation of the ‘Nucleation & caging’ model. Most ParB dimers (green dots) are highly confined in a cluster (dotted circle) centered on the parS sites (black rectangles) onto which some ParB are stably bound (red dots). The DNA entering the cluster is bound stochastically by ParB. Red and blue lines represent DNA present at small and large (or on a different molecule) genomic distance from parS, respectively.

(B) ParB clusters on plasmid F in vivo. Typical E. coli cells (DLT3594) displays foci of ParB_F-mVenus protein (top) expressed from the endogenous genetic locus of the plasmid F (F1-10B-mVenus). The nucleoid is labelled with Hu-mCherry (central). The overlay (bottom) combines the two fluorescent channels. Over 99 % of cells harbor ParB_F foci. White bars: 1 µm.

(C) ParB_F binding outside parS_F on the plasmid F follows a power-law decay. High resolution ChIP-seq performed on DLT3586 carrying the plasmid F (F1-10B). The ParB density, normalized to 1 at the first bp after the last parS_F binding repeat, is displayed over 14-Kbp on the right side of parS_F. Monte Carlo simulations and analytic formula are represented in red and dotted black lines, respectively. MC simulations are performed with a Freely-Jointed Chain of length N=4000 monomers of size a=10-bp, preventing finite size effect on the range of genomic coordinate considered. The cluster radius is σ=75nm and the binding energy difference ε_m-ε_s=-0.9 is obtained by reading the density at the 1st site after parS (see text and supplemental information). As a benchmark for simulations, the analytics are obtained from Eq.(1) with the same parameters. Insert: The ParB_F binding profile is represented as the number of nucleotide reads over 80-Kbp centered at parS.

(D) and (E) Same as B and C with parS_F inserted at the xylE locus on E. coli chromosome from DLT3584 and DLT2075, respectively. The Kuhn length was adjusted to a=23-bp in the
simulations and analytics. The characteristics of the A-F genetic loci are presented in Fig. S1A.

**Figure 2. ParBF DNA binding pattern is robust over a large range of intracellular ParBF concentrations.**

(A) Normalized ParBF binding profiles at different ParBF/parS ratio. ChIP-seq density on the right side of parS inserted at xylE were measured in DLT2075 induced (16, 28) or not (0.4) with IPTG, or carrying HCN plasmids pZC302 (0.04) or pJYB57 (0.016). The ParB/parS ratio is calculated relative to the one of plasmid F as determined from Western blot analyses (Fig. S2B). Monte Carlo simulations and analytical formula are plotted with the same parameters as in Fig. 1E. Inset; the amplitudes of the curves are rescaled by the indicated factors to overlap with the curves of highest amplitude.

(B) ParBF are dispersed in the cell upon titration by HCN plasmids. ParB-mVenus expressed from pJYB294 were imaged as in Fig. 1D in DLT3577 (left) and DLT3576 (right) carrying pZC302 and pJYB57, respectively. The number of extra parS per cell, indicated on top of each raw, are estimated from the copy number per cell of HCN plasmids carrying 10 specific binding sites. White bars: 1 µm.

(C) The size of ParBF clusters is independent of the intracellular ParBF concentration. We considered two possible evolutions of the cluster size upon variations of ParB amount in the framework of ‘Nucleation & caging’ with corresponding schematics drawn on the right. Top: constant ParB concentration; supposing that clusters are compact, the cluster radius $\sigma$ would depend on the number $m$ of ParB like $\sigma = m^{1/3}$. Predictions profiles, plotted at different ratio of ParB/parS, vary within the range of the experimental levels tested. Bottom: constant cluster size; ParB concentrations vary but the range of exploration remains the same resulting in overlapping prediction profiles.
Figure 3. The box II motif of ParBF is crucial for ParBF binding in the vicinity of parSF and cluster formation.

(A) The formation of secondary ParBF-DNA complexes requires the box II motif. EMSA were performed with a 144-bp 32P-labelled DNA fragments (C144) carrying a single 16-bp parS binding motif. Reaction mixtures containing 100 µg.ml⁻¹ sonicated salmon sperm DNA were incubated in the absence (-) or the presence of increasing concentrations (grey triangle; 10, 30, 100, 300, and 1000 nM) of ParBF or ParBF-3R*. Positions of free and bound probes are indicated on the left. B1 represents complexes involving the specific interaction on the 16-bp binding site, while B’2 and B’3 complexes represent secondary complexes involving the parSF site with one or two additional nsDNA-binding interactions, respectively.

(B) ParBF cluster formation requires the box II motif. Epifluorescence microscopy of ParBF-3R*-mVenus from DLT3566 is displayed as in Fig. 1D. White bars: 1 µm.

(C) ParBF in vivo DNA binding in the vicinity of parSF sites requires the box II motif. ChIP-seq was performed on DLT3726 carrying parSF in the xylE chromosomal locus and expressing ParBF-3R* variant. ParBF-3R* DNA binding profile displayed the number of nucleotide reads as a function of the E. coli genomic coordinates. The peak at parSF covered approximately 950-bp, which corresponds to the 402-bp between the 1st and 10th specific binding sites and ~280-bp on each sides (representing the average size of the DNA library; see Fig. S3D). No ParBF-3R* enrichment was found on parSF-flanking DNA and elsewhere on the chromosome. Inset, zoom in on the right side of parSF over 5-Kbp with the ParB density, normalized to 1 at the first bp after the last parS binding repeat, plotted as a function of the distance from parSF.

Figure 4. ParB of V. cholerae assembled in cluster similarly to ParBF.
(A) Schematic representation of the genomic locus of the chromosome 1 of *V. cholerae* with the three *parS* sites, named *parS*1-3. The rRNA operon (blue rectangle) spans the genomic coordinates 53823 to 59123.

(B) ChIP-seq performed on strain N16961 is displayed as the number of nucleotide reads in function of the genomic coordinates. Correspondence to the *parS*1-3 location in (A) is indicated by grey dotted lines.

(C) We model the ChIP-Seq data as in Fig.1C-E by means of MC simulations with a Freely jointed chain of size *N*=2000 monomers of size *a*=16-bp. The best fit was achieved with *σ*=25nm and the difference *ε*ns-*ε*s1=-0.2 between non-specific binding energy and chemical potential (read directly from ChIP-Seq data). In the MC simulation, we accounted for the finite width of the distribution around *parS* sites by including the average size of the DNA library (304-bp; Fig. S1D and S4B).

**Figure 5. Robust dips and peaks signatures in ParB DNA binding profiles.**

ChIP-sequencing assays were performed on DLT2075 (*xylE::parSF*) expressing ParB*F* grown in exponential (expo) or stationary (stat) phases with addition of rifampicin when indicated (+Rif).

(A) ParB*F* DNA binding around *parSF* is independent of active transcription. The color-coded ParB*F* profiles are represented over 50-Kbp as the relative ParB density normalized to 1 at the first bp after the last *parSF* binding site. Loci A, C, E and F are defined in Fig. S1A.

(B) The dips and peaks are highly similar in the three indicated conditions. Same as in (A) with zoom in on the right side of *parSF* up to 9-Kpb and normalization to 1 at genomic coordinate 230. The dotted line corresponds to the analytics description of Stochastic Binding (see details in Fig. 1C-E).
(C) ParBF binding profile upstream of the locus A. Same as in (A) with zoom in from -6.5 to -16.5-Kbp and normalization to 1 at genomic coordinate -6.5-Kbp (upstream of the dip at the locus A). ParBF binding profile follows the power-law characteristic of stochastic binding, represented by the analytics description (dotted line), upstream of the locus A in stationary phase (black) and in exponential phase (blue).

(D) The promoter region at locus A prevents ParBF DNA binding. Chip-seq assays were performed in isogenic xylE::parSF strains (DLT2075; black curve) in which the locus A is replaced by a kanamycin gene (DLT3651; red curve). The relative ParB density as a function of the distance from parSF is drawn and normalized as in (A). The promoter region is depicted as in Fig. S1A.

Figure 6. ParB dynamics between partition complexes.

ParBF exchange between foci was measured by FRAP and FLIM (fluorescence lifetime imaging microscopy) from two-foci cell of DLT1215 carrying pJYB234.

A- Representative images of a photobleached cell during a FRAP experiment. The 488 nm laser was pulsed (Bleach) on one of the two foci at ~2.4 sec (black arrow). Red and blue arrows correspond to the bleached and unbleached focus, respectively. Time is indicated in second (upper right). The cell outline is drawn in red. Scale bar: 1 μm.

B- Quantification of ParBF-mVenus fluorescence intensity over time. The dynamics of fluorescence intensity is shown from averaging 18 independent measurements of the bleached (FRAP, red line) and unbleached (FLIM, blue line) foci. Foci fluorescence intensity in each experiment was normalized to the average intensity of each focus before photobleaching. Natural bleaching during the course of the experiments (green curve) was estimated for each measurement by averaging the fluorescence intensity of 15 foci present in each field of view.
Figure 1
Figure 2

A

![Graph showing ParB density vs. genomic distance from parS](image)

B

**Extra parS binding sites**

- 200
- 500

ParB

Nucleoid

Overlay

C

![Graph showing ParB density vs. genomic distance from parS](image)
Figure 3

(A) Electrophoretic image showing PCR products for ParB\textsubscript{F} and ParB\textsubscript{F}-3R\textsuperscript{*} with bands designated as B', B, and C144.

(B) Images of ParB\textsubscript{F}-3R\textsuperscript{*}, Nucleoid, and Overlay.

(C) Graph showing number of reads (x10\textsuperscript{3}) plotted against genomic distance from par\textsuperscript{S}.

Legend:
- ParB\textsubscript{F}: ParB\textsubscript{F} without the 3R region.
- ParB\textsubscript{F}-3R\textsuperscript{*}: ParB\textsubscript{F} with the 3R region.
- xylE::par\textsubscript{S}\textsubscript{F}: Insertion of xylE::par\textsubscript{S}\textsubscript{F} into the genome.
Figure 4

(A) Genomic coordinates (Kbp)

(B) ParB density

(C) ChIP-Seq data vs. MC simulations
Figure 5
Figure 6

A.

Pre-bleached | Bleach | Recovery

0.2
0.9
3.4
8.2
13.2
23.2
28.2
33.2
68
98
139
169

B.

FRAP | FLIM | Bleaching over time

Relative intensity

N = 18

Time (sec)