Review

Hard-Wired Control of Bacterial Processes by Chromosomal Gene Location

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Bacterial processes, such as stress responses and cell differentiation, are controlled at many different levels. While some factors, such as transcriptional regulation, are well appreciated, the importance of chromosomal gene location is often underestimated or even completely neglected. A combination of environmental parameters and the chromosomal location of a gene determine how many copies of its DNA are present at a given time during the cell cycle. Here, we review bacterial processes that rely, completely or partially, on the chromosomal location of involved genes and their fluctuating copy numbers. Special attention will be given to the several different ways in which these copy-number fluctuations can be used for bacterial cell fate determination or coordination of interdependent processes in a bacterial cell.

How Genome Organization and Gene Function Are Connected

For decades, the importance of genome organization has been recognized. Virtually every process that interacts directly or indirectly with the chromosome has left its marks during the course of genome evolution. It has become clear that the order and orientation of features on a chromosome, as well as the three-dimensional structure of the chromosome, is of importance to a cell. Numerous examples of the interplay between genome organization and cellular processes are available. For example, essential genes tend to be located on the strand that is transcribed in the same direction as in which replication proceeds [1].

However, the importance of the genomic location of key elements is still often underestimated. In fact, very little attention is given to the many different ways in which genomic location can impact cell biology. We therefore review the various mechanisms by which the exact genomic location of a feature can play a role in the regulatory landscape and development of bacterial cells. More specifically, we focus on processes in which gene copy number or, more accurately, genome-wide copy number distributions play a role. It is a well-established fact in eukaryotes that having an abnormal number of chromosomes (aneuploidy), leading to atypical gene copy numbers, can have detrimental effects, a well-known example being Down syndrome (trisomy 21 in humans, [2]). Additionally, the need for female mammals to silence one of their two copies of the X-chromosome underlines the importance of DNA copy numbers [3,4]. Furthermore, amplification of specific nutrient transporter genes in Saccharomyces cerevisiae was observed to enhance fitness in nitrogen-limited conditions [5]. The correlation between copy number and gene expression implied by these examples was confirmed recently by Chen and Zhang, who showed that the timing of replication of a gene influences its final expression level in yeast [6]. Nevertheless, copy number effects are still only rarely considered in prokaryotes. During bacterial cell cycle progression, copy numbers around the chromosome fluctuate periodically. Both the periodicity [7] and the amplitude [8,9] of this fluctuation can be employed to regulate certain

Trends

While well appreciated in eukaryotes, the relevance of gene copy numbers in bacteria is largely unexplored.

Copy number imbalance between two genes during DNA replication was found to be responsible for precisely timed activity pulses of the Bacillus subtilis sporulation initiator.

Besides the various ways in which chromosomal gene location has been shown to be involved in B. subtilis sporulation, more examples have recently been found in other bacteria as well.

Changing nutrient availability or malfunction of critical subcellular processes lead to distorted gene copy number distributions, with transcriptional shifts as a result, for example accounting for the activation of pneumococcal competence.

Additionally, the discovery that mRNA diffusibility is unexpectedly low suggests that the chromosomal location of a gene may be in part determined by where in the cell its product is required.

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processes in the cell. Furthermore, global or local (e.g., compartmentalization, see below) distortions of copy number fluctuations can be involved in bacterial ‘decision making’ and even play an important role during virulence [9].

Replication-Associated Copy Number Fluctuations

The majority of bacteria have their DNA organized on a single, circular chromosome, replication of which starts at a well defined origin of replication (oriC). From there, replication proceeds symmetrically in both directions around the chromosome and is terminated at the opposite end (the ter region) of the molecule, where both replication machineries (forks) meet. As a result, the various genes and other features on the chromosome are replicated in a fixed order, leading to periodic fluctuations of their copy numbers that are repeated every cell cycle. After termination of replication, cells still need a specific amount of time to finish cell division (the D-period [9]). The initiation of new rounds of replication is tightly regulated by a variety of factors [10–13]; this ensures that there is exactly one initiation event in each cell cycle, timed in such a way that replication and cell division are properly coordinated. When growth is sufficiently slow, cells have enough time to start and finish DNA replication within one cycle, and local copy numbers will generally only fluctuate between one and two copies of a certain region (Figure 1A). Some bacteria, however, have the capacity to grow so fast that replication of their entire chromosome cannot be completed within one cell cycle [14]. In this case, cells engage in multifork replication; before a replication fork has finished, a new replication initiation event takes place (still exactly once per cell cycle) at all (>2) copies of oriC simultaneously, resulting in copy numbers of oriC-proximal regions of more than 2 (Figure 1B). For example, fast-growing *Escherichia coli* cells have been observed to contain up to 8 origins [15]. Since there is a clear correlation between gene copy number and gene expression [16–18], these fluctuations are relevant to a cell’s transcriptome, as is exemplified by the various cases mentioned in this review.

Function-Associated Gene Order

The amplitude of a gene’s copy number fluctuation will thus depend both on its genomic location, relative to oriC, and on growth rate. The impact of these dependencies is illustrated by the fact that translocations and chromosomal inversions preferentially occur in a copy-number-neutral fashion (i.e., symmetrical with respect to oriC) [19–21]. Another example of the importance of gene order is the strong conservation of the oriC-proximal colocalization of important growth factors involved in replication, transcription and translation [14,22,23]. The colocalization of these factors can be explained by a combination of the importance of their stoichiometry on the one hand and functional compartmentalization (see below) on the other. However, the fact that they are virtually always found close to the origin of replication rather reflects the cells’ need to correlate their expression with their requirement; when growth conditions improve, cells may switch to multifork replication, automatically boosting the expression of these essential growth factors due to the resulting dosage increase. Recent work by Soler-Bistué *et al.* demonstrates the relevance of the genomic position of ribosomal protein genes on the large chromosome of the human pathogen *Vibrio cholerae*, which harbors two circular chromosomes (Figure 1C) [9]. They showed that translocation of a locus bearing half of all ribosomal protein genes from oriC-proximal to various sites further away from the origin of replication results in significant defects in growth and host-invasion capacity. It is worth noting that these defects specifically occur during relatively fast growth, where the difference in copy number between oriC and ter, and therefore the relative effect of translocation of the ribosomal protein genes, is the largest. Both defects are relieved when, instead of one, two copies of the locus are present at an oriC-distal site, effectively restoring absolute ribosomal gene copy numbers and consequently ribosomal production levels. The fact that these genes are then no longer colocalized with other important growth factors is, apparently, of lesser importance in this context.
Figure 1. Replication-Associated Gene Copy Number Fluctuations. Simulated gene copy number distributions throughout the cell cycle (A and B). Each arm of the chromosome has been divided into four quartiles, which are color-coded based on their oriC-proximity. The height of each colored area in the graphs represents the average copy number within the corresponding quartile; as the replisome moves through a quartile, the corresponding graph area steadily increases in height until it is exactly doubled (i.e., the entire quartile is replicated), while the other areas maintain their height. The areas describing the copy number development of the four quartiles are stacked, so their combined height reflects the total DNA content of a cell. Average copy numbers of each quartile at 10%, 50%, and 90% of the cell cycle are shown in the plots. The script to run the simulations is available upon request. Replication initiation is indicated by black arrows. (A) During relatively slow growth (replication time/cell cycle = 0.5, D-period = 10% of cell cycle), only one replication fork is present at a time on each arm of the chromosome (top) and gene copy numbers will fluctuate between 1 and 2 (bottom). (B) During relatively fast growth (replication time/cell cycle = 1.6, D-period = 10% of cell cycle), multifork replication occurs (top) and gene copy numbers can exceed 2 (bottom). (C) The oriC-proximal location of the *Vibrio cholerae* S10 ribosomal protein operon is important for fitness [9]. Top: translocation of these genes to an oriC-distal site leads to lower gene copy numbers and therefore to a growth defect and attenuated infectivity. Merodiploid strains, with two copies of the S10 operon, show restored fitness and infectivity. Bottom: locus-dependent average copy number over the cell cycle for fast-growing cells (same parameters as in (B), closely matching the oriC-ter ratio observed by Soler-Bistué et al. [9]). Inspection of copy numbers at the varying loci of S10 operon placement shows that S10 gene dosage in the merodiploid strain is very similar to that in the wild-type strain.

Similarly, Sobetzko _et al._ demonstrated that nucleoid-associated proteins (NAPs) employed during exponential growth, together with their binding sites, show a tendency to be located closer to oriC than NAPs that act in (near-)stationary phase [23]. Simultaneously, they showed that genes with related functions have a propensity to be distributed at equal distances from oriC, without the necessity of being on the same arm of the chromosome [23]. Taken together, these observations underline that the variation in growth conditions encountered throughout
evolution is directly reflected by the relative positioning on the chromosome of genes with related functions.

**Chromosome Structure and Gene Expression**

Of course, the correlation between gene copy numbers and final expression levels [16–18] represents only part of the puzzle. Bryant et al. showed that changing the chromosomal location of a reporter cassette in the Gram-negative model organism *E. coli* could lead to differences in expression level of several hundredfold [24]. Similar observations were made in the Gram-positive human pathogen *Streptococcus pneumoniae* (the pneumococcus) [25]. These variations cannot be explained by DNA copy number differences alone, especially since the experiments were executed in slow-growth conditions, with no multifork replication and hence with minimal copy number variation along the oriC-ter axis. Possible explanations for these large differences include gyrase activity, transcription-associated supercoiling and the presence of NAP binding sites [24,26,27], many of which play a role in determining chromosome structure [28,29].

Clearly, a myriad of forces have been, and are, at play simultaneously in the evolution of chromosome organization. For that very reason, countless patterns have been, and doubtlessly will still be, discovered in both the order of chromosomal features and chromosome morphology. Although not the focus of this review, the impact of chromosome topology and overall structure on gene expression and chromosome organization cannot go unmentioned. It has been known for decades that a change in DNA supercoiling can affect gene expression [30], and it was recently shown by Sobetzko that preserving the gene-regulatory capacity of supercoiling has been a driving force in the evolution of chromosomal gene order [31]. Furthermore, the folding and compaction of the chromosome can bring genes that are located on very different parts of the DNA molecule into close proximity to each other [32–34]. Interestingly, a periodic distribution of conserved gene pairs around the *E. coli* chromosome was observed with a period of 117 kb [35,36]. This periodicity possibly allows the chromosome to be folded such that these genes, and thereby their corresponding products, end up in close spatial proximity to each other. The surprisingly low diffusibility of some mRNA molecules from their production site suggests that the well defined organization of the chromosome within the cell might serve as a blueprint that determines where in the cell a certain protein is synthesized [37–39]. However, recently, Moffitt et al. probed the localization of 75% of the *E. coli* transcriptome using a combination of fluorescence in situ hybridization and super-resolution microscopy, showing that the role of genome structure in mRNA localization is rather the exception than the rule [40].

As mentioned above, final production levels of a certain protein are determined by many more factors than gene dosage alone. For example, when Gerganova et al. moved the *fis* gene, encoding the *E. coli* global regulator FIS, from an oriC-proximal to a terminus-proximal location, cells were able to maintain the original protein level by upregulating transcription of *fis* [41]. Despite the nearly unchanged *fis* transcript and FIS protein levels, the mutant displayed a significant fitness loss. This was attributed to altered levels of some key NAPs that are directly regulated by FIS and that are, among other things, implicated in promoter binding and DNA supercoiling. While copy-number changes could not be blamed for this observation, the authors speculated that the limited diffusibility of both the *fis* transcript and FIS might be responsible, leading to different activation patterns of its regulon and thereby possibly affecting DNA topology and structure.

**The Various Roles of Chromosome Organization in *B. subtilis* Sporulation**

The Gram-positive model bacterium *Bacillus subtilis* is of great interest when studying culture heterogeneity and cell fate determination. When exposed to stress, it can behave in very different ways [42]: under mild nutritional stress *B. subtilis* can form biofilms, multicellular communities
Figure 2. The Various Roles of Gene Location in Bacillus subtilis Sporulation. (A) Overview of endospore development. Sporulation is initiated by high levels of Spo0A–P, leading to a block of matrix formation. After asymmetric septation, transcriptional differentiation is directed by dedicated sigma-factors, eventually leading to spore maturation and mother cell lysis. (B) Cell-cycle-mediated pulsing controls onset of sporulation [7]. The pulsatile profile of the Spo0A–P concentration during the cell cycle (bottom graph, bottom panel) is made possible by the asymmetrical placement (top left) and therefore copy number development of spo0F and kinA (bottom graph, top panel; red and orange lines, respectively), which are involved in the sporulation phosphorelay, leading to phosphorylation of Spo0A (top right). (C) Temporary diploidy preceding spore development leads to shut-down of matrix production [51]. Simulated copy number distributions throughout the cell cycle (same plotting and simulation parameters as in Figure 1A) demonstrate how gene dosage is affected prior to sporulation (bottom, orange-red areas). Due to the ter-proximal location of regulatory genes sinI and sinR (top left), unusual levels of their respective products are reached (bottom, light orange lines). Due to
Cell-Cycle-Mediated Pulsing Controls Onset of Sporulation

The decision to engage in sporulation is taken on a single-cell level and is controlled by a complex phosphorelay system, composed of several kinases and phosphotransferases [46], with master regulator Spo0A at the end of the line. When the amount of active, phosphorylated Spo0A (Spo0A–P) reaches a threshold level, downstream genes are activated and the sporulation program begins [47]. It is, at that point, important for a cell to have completely copied its DNA, so that the spor can receive an intact chromosome and therefore proliferate [43,44,48]. A number of studies showed that B. subtilis realizes the necessary coordination of sporulation activation with the cell cycle by producing a single sharp pulse of active Spo0A–P every cycle, shortly after termination of DNA replication (Figure 2B) [7,49,50]. Thereby, whenever the conditions are such that the threshold level of Spo0A–P is reached, this will occur at a phase of the cell cycle in which two complete copies of the chromosome are present. Narula and coworkers recently showed that the pulse-like dynamics of Spo0A activation largely rely on two requirements, the first one being the presence of negative feedback between Spo0A–P and one of the upstream phosphotransferases, Spo0F [7]. Crucially, this feedback is not instantaneous, but delayed since Spo0A–P transcriptionally increases levels of Spo0F, which in turn is thought to interfere with substrate binding by KinA, the major sporulation kinase, lowering the phosphate flux towards Spo0A. The second requirement is that, during the replication cycle, there is a temporary imbalance between the copy numbers of spo0F and kinA. This imbalance is due to their different locations on the genome: since spo0F is located close to the origin, while kinA is located closer to ter, the kinA:spo0F ratio will temporarily drop after replication of spo0F, leading to a gradual decrease in Spo0A–P. After kinA is also replicated, the Spo0A–P level increases again, then overshoots (i.e., pulses), due to the aforementioned delayed negative feedback loop, and only then falls back to its equilibrium level. In accordance with this model, the researchers showed that translocating kinA and/or spo0F led to sporulation defects, stressing the importance of their chromosomal location for successful endospore formation.

Temporary Diploidy Ensures Mutual Exclusion of Sporulation and Biofilm Formation

Although biofilm formation and sporulation are activated by the same protein, phosphorylated Spo0A (Spo0A–P), both states are mutually exclusive since matrix production is completely absent in cells that have started sporulating [51,52]. The regulation of matrix genes by Spo0A–P is indirect, by activation of the gene encoding Sinl, which counteracts SinR, a repressor of genes responsible for matrix production. Chai et al. showed that while the repression by SinR is cooperative, derepression by Sinl takes place uncopiously, that is, single Sinl molecules bind SinR [51]. They argued that these binding characteristics provided an answer for the somewhat paradoxical mutual exclusion of sporulation and matrix formation. Part of the explanation lies in...
the difference in Spo0A–P binding affinity between biofilm and sporulation gene promoters; while biofilm formation is mainly triggered by lower levels of Spo0A–P, sporulation is activated only when levels are high [47]. More importantly though, the cells make use of the unique situation at the start of sporulation, when there are two complete copies of the chromosome present for a prolonged period of time (Figure 2C) [53]. Due to this temporary diploidy, copy numbers of terminus-proximal genes sinI and sinR are roughly doubled. Due to the cooperative nature of repression by SinR, this may definitively shift the balance towards complete absence of matrix gene expression.

Physical Compartmentalization Is Responsible for Transcriptional Differentiation between Mother Cell and Prespore

In the course of spore development the replicated chromosome of the predivisional cell is segregated such that both the mother cell and the spore receive one copy. Importantly, however, in the early stages of sporulation only the oriC-proximal one-third of the chromosome is translocated into the prespore (Figure 2D–E). The remainder of the molecule is gradually translocated over a time period of around 15 minutes [54,55].

The transcriptional differentiation between the two compartments (prespore and mother cell) is largely initiated by two different sigma factors, $\sigma^F$ and $\sigma^E$, that activate two separate gene expression programs in the mother cell and prespore, respectively [44].

As discussed by Hilbert and Piggot, $\sigma^F$ activity is obstructed by anti-$\sigma$ protein SpoIIAB (Figure 2D). The release of $\sigma^F$ is promoted by the dephosphorylated form of SpoIAB (an anti-anti-$\sigma$ protein), which can, however, become phosphorylated through the kinase activity of SpoIIAB. Dephosphorylation of SpoIAB is performed by protein phosphatase SpoIE [44].

While $\sigma^F$, SpoIAB, and SpoIIAB are encoded at a terminus-proximal site on a single transcriptional unit, spoIIIE is located close to oriC. It was suggested that the transient overproduction of SpoIIIE may be responsible for the eventual release of active $\sigma^F$ in the prespore (Figure 2D) [56,57]. Frandsen et al. showed that moving the coding sequence for $\sigma^F$ to an origin-proximal site allowed spore production even in the absence of the normally required factors SpoIAB and SpoIIIE [56]. McBride et al. showed that displacement of spoIIIE to origin-distal sites partially repressed prespore-specific genes, but did not affect overall sporulation efficiency [57]. These findings suggest that more factors play a role in initiating the prespore’s transcriptional program.

In the mother cell, the precursor of $\sigma^F$ (pro-$\sigma^F$), encoded by sigE, is inactive due to an N-terminal 27-amino-acid extension that can be cleaved off by SpoIIGA, resulting in active $\sigma^E$. Even though the N-terminal extension may be relevant for proper development of sporulation, it is the terminus-proximal location of the sigE-SpoIIGA operon that seems to be responsible for the initial accumulation in the mother cell [57]. To ensure that the mother cell only enters its transcriptional program when that of the prespore has fully initiated, $\sigma^F$-dependent, prespore-localized expression of another gene, spoIIIR, is required for activation of $\sigma^F$ in the mother cell (Figure 2E) [58,59]. SpoIIIR is thought to function as a signal for the septum-localized SpoIIGA, which then processes pro-$\sigma^F$ into $\sigma^E$ [44]. The oriC-proximal location of spoIIIR is required for sufficient expression levels in the prespore during asymmetrical segregation. This became clear when Khvorova et al. saw a drastic delay in timing and decrease in level of sporulation gene induction upon the translocation of spoIIIR to a ter-proximal site [60].

In the mother cell, $\sigma^F$ activity is prevented, perhaps in part due to lower copy number of SpoIIIE, but probably mainly due to mother-cell-specific degradation of SpoIIIE (Figure 2E) [57]. At the same time, $\sigma^F$ activity in the prespore is repressed by binding to anti-$\sigma$ protein CsfB, which is again encoded on the prespore-localized one-third of the chromosome (Figure 2D) [61].
B. subtilis sporulation thus provides us with many examples of the relevance of chromosome organization for key processes in a bacterial cell, through a variety of mechanisms.

Distortion of Natural Gene Dosage Fluctuation Induces Bacterial Competence

Whether or not a bacterium will perform multifork replication largely depends on the combination of its growth rate and its genome size. As discussed earlier, the oriC-proximal location of genes encoding important growth factors automatically correlates their production and requirement levels. A different way in which oriC-proximity is utilized is found in the pneumococcus (Figure 3). With its relatively small genome (~2 Mb, [62]), multifork replication in rapidly dividing cells has not been observed [63]. This situation alters, however, when replication fork progression is directly or indirectly perturbed and slowed down. Since, as far as we know, there is no instantaneous feedback to the pneumococcal replication initiation system, new replication complexes may be loaded onto the genome before the stalled or slowed replication forks have finished, leading to increased dosage of oriC-proximal genes.

Figure 3. Competence Activation in Streptococcus pneumoniae Due to Dosage Upshift of oriC-Proximal Regulator Genes. The oriC-proximal location of early competence genes allows the pneumococcus to activate this state in response to replication stress [63]. Simulated development of copy-number distribution during replication stress is shown in the bottom graph (bottom panel; same plotting parameters and (initially) same simulation parameters as in Figure 1A). Halfway in the second cell cycle, replication stress is applied (red star; new replication rate is one-third of original replication rate), while timing of replication initiation events is unaltered (black arrows). Note that time units indicated with an asterisk are multiples of the cell cycle time in the absence of replication stress. Due to the oriC-proximal location of comAB and comCDE, their expression levels increase (bottom graph, top panel) and once a certain threshold activity is reached, competence is activated via the positive feedback loop in its regulatory system (top right).
Various factors can lead to this form of overinitiation: DNA damage (e.g., double-strand breaks induced by mitomycin C); insufficient functioning of type II topoisomerases, which are responsible for the relaxation of DNA required for replication forks to progress (e.g., induced by fluoroquinolone antibiotics); or limited nucleotide availability (e.g., induced by trimethoprim and hydroxyurea). *S. pneumoniae* makes use of this exceptional situation to activate competence or the so-called X-state, one of the most important stress response mechanisms it has at its disposal, allowing cells to take up exogenous DNA [64]. The activation of this system encompasses the expression of over a hundred genes [65–67], blocks cell division [68], and thus represents a significant burden for the cell. It is therefore important for the cell to somehow regulate the activation of this system. Despite the large number of genes eventually being activated, the on/off switch of the X-state is constituted by a positive feedback loop containing a set of only five genes organized into two operons [69], comAB and comCDE. Very-low-level basal expression occurs for both operons. ComC is a 41-residue peptide containing a double-glycine leader of 24 amino acids in length. Membrane-associated transporter complex ComAB exports ComC, cleaving off the leader peptide, and extracellularly releasing the 17-residue competence-stimulating peptide (CSP), which acts as a quorum-sensing autoinducer [70]. ComCDE constitutes a typical two-component regulatory system: the membrane-bound histidine kinase ComD binds the extracellular CSP and consecutively transfers a phosphate group to the response regulator ComE, resulting in ComE-P. ComE-P then completes the positive feedback loop by enhancing expression of both comAB and comCDE [71]. Additionally, it induces the expression of comX, coding for the X-state-specific sigma factor σ^X_, required for the activation of the entire competence regulon [72]. However, processes such as mRNA and protein degradation and dilution by growth will counteract this positive feedback loop and may prevent the X-state from switching on. Additionally, the autocatalytic efficiency of the system is dependent on medium parameters such as pH. Only when the local extracellular CSP concentration exceeds a certain threshold, the positive feedback may outcompete the counteracting forces and X-state gene expression may dramatically increase (possibly with several orders of magnitude). Hence, whether or not the X-state is activated depends on a complex set of parameters, including the copy numbers of comAB and comCDE; because of their oriC-proximal location on the chromosome (8° and −1°, respectively), relative overinitiation (e.g., due to replication fork stalling) can push up the dosage of early X-state genes. It was shown that even a slight increase in dosage, of below twofold, can suffice to reach threshold CSP concentrations and lead to X-state activation [63]. Interestingly, it was recently shown that the production of pneumococcal bacteriocins (pneumocins) is also potentiated by X-state activation [73,74]. Since pneumocins play an important role in intra- and interspecies competition in their natural niche (the human nasopharynx), the gene-dosage-induced activation of the X-state may cause the composition of the nasopharyngeal flora to change, for better or for worse.

**Implications for Future Research**

**mRNA Noise Prediction**

Over the past 15 years it has become more and more clear that stochastic ‘noise’, which is intrinsically present in any process that takes place in a cell, can have tremendous effects on cell fate determination [75,76]. Phenotypic heterogeneity can arise when noisy expression of certain regulators only exceeds a threshold in a subpopulation of cells. To fully understand the impact of molecular noise on macroscopic properties, such as a bacterial phenotype, it is important to know all the relevant parameters that contribute to the absolute level of variation in a specific process. Recently, Peterson et al. showed that, for accurate modeling of mRNA distributions, it is imperative to account for DNA replication progressing and at some point doubling the copy number of the gene under study. Since replication is very tightly regulated, replication noise is negligible in this context, and incorporation into noise models should be
relatively straightforward. Ignoring the effect of replication leads to an—in some cases very severe—overestimation of mRNA noise [77].

Normalization for Differential Gene Expression Analysis
With the upsurge of Next-Generation Sequencing (NGS) techniques, such as RNA-Seq, over the past decade, transcriptome studies have become much more attainable for many microbiologists around the globe. Analysis of the obtained datasets is often performed in automated pipelines that do not require much input from the user. The relative ease of use of these techniques has led to large amounts of highly valuable data. To our knowledge, however, none of the existing analysis packages take into account the possibility of a genome-wide copy number shift, which is expected in several conditions, including treatment with certain antibiotics [63] and changes in growth rate, temperature, or replication rate. As a result of such an altered gene-dosage pattern around the chromosome, it may become difficult to set a proper baseline of expression or differential expression; most normalization methods depend, at least in part, either on the assumption that most genes will have an unaltered expression level (e.g., upper-quartile or median normalization), or on the assumption that the total number of transcripts per cell remains roughly the same (e.g., transcripts per million (TPM, [78]) as a measure of expression). In case of a global shift in copy-number distribution, neither of these assumptions is necessarily valid, and failing to acknowledge this may lead to overestimation of the number of truly differentially expressed genes, and simultaneously camouflage the changes of interest. The most accurate interpretation of data in these situations depends on the question one is trying to answer, but it is important to be aware of the role that copy-number changes have in these experiments.

Additionally, normalization methods for quantitative reverse-transcriptase-PCR (qRT-PCR) are usually based on the assumption that certain reference genes will keep a constant expression level in the various conditions that are being compared. This may be, even in the absence of copy-number effects, a very dangerous assumption, but it is especially so when gene-dosage distribution shifts are in play. It is therefore advisable to, in addition to using more than one reference gene, confirm that copy-number shifts are not responsible for the observed results. This can be accomplished by a qPCR experiment using chromosomal DNA as a template.

Copy-Number Effects in Synthetic and Natural Systems
In this review we highlighted the role of copy-number fluctuations in bacterial processes. In synthetic constructs the copy number of an integrated DNA sequence can be critical for its proper functioning and in these cases the location of chromosomal integration should be carefully deliberated [18].

Surprisingly, beyond the well known replication-associated gene dosage, not too many examples of bacterial decision-making are available that have been ascribed to the genomic location of key factors involved. Over the past couple of years, however, several of these examples have emerged [7,9,51,63]. Combined with the fact that DNA replication is universally present in all living organisms, this is highly suggestive of the possibility that these effects are much more abundant in bacterial biology than currently acknowledged. Going further, the reviewed phenomena are not necessarily limited to bacteria; archaeal chromosomes typically contain no more than a few replication origins, and since bacterial and archaeal chromosomes share several organizational traits [79], some of the mechanisms discussed here may very well be active in archaea as well.

Concluding Remarks
Various aspects of chromosome organization, including chromosome structure and topology, have been described and are increasingly being studied. The growing pool of knowledge on
properties related to the spatial organization of genes, including accessibility to transcription-related proteins, spatial colocalization of genes, and mRNA and protein diffusibility, will be very important for the understanding of bacterial gene regulation. However, while not unrelated, chromosomal gene order is yet another aspect that affects the regulatory landscape of a bacterial cell, and has not received as much attention as necessary. As described in this review, the exact position of a gene on the chromosome determines when, where, and how often its DNA is copied. Although gene order is not critical for cell survival in laboratory conditions [80], the significance of this facet of chromosome organization is emphasized by the several examples that have emerged of regulatory processes that depend on the dynamic copy number fluctuations during DNA replication [7,9,51,63]. Future research will have to determine how widespread these mechanisms are (see Outstanding Questions).

Additionally, regardless of the role of the chromosomal location of a gene under natural circumstances, it is important to keep in mind the potential impact certain experiments will have on copy number distributions in a cell; translocation of genes, antibiotic treatment, nutrient limitation, and other types of stress can each in their own way induce transcriptional changes by affecting gene dosage, either locally or genome-wide. A better understanding and increased awareness of the role of chromosomal gene order in the regulation of key processes is therefore paramount in understanding bacteria, and possibly also archaea, both in nature and in the laboratory.

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