Chromosome and plasmid-borne $P_{\text{LacO3O1}}$ promoters differ in sensitivity to critically low temperatures

Samuel M. D. Oliveira1, Nadia S. M. Goncalves1, Vinodh K. Kandavalli1, Leonardo Martins1,2, Ramakanth Neeli-Venkata1, Jan Reyelt3, Jose M. Fonseca2, Jason Lloyd-Price4,5, Harald Kranz3 & Andre S. Ribeiro1,2

Temperature shifts trigger genome-wide changes in Escherichia coli’s gene expression. We studied if chromosome integration impacts on a gene’s sensitivity to these shifts, by comparing the single-RNA production kinetics of a $P_{\text{LacO3O1}}$ promoter, when chromosomally-integrated and when single-copy plasmid-borne. At suboptimal temperatures their induction range, fold change, and response to decreasing temperatures are similar. At critically low temperatures, the chromosome-integrated promoter becomes weaker and noisier. Dissection of its initiation kinetics reveals longer lasting states preceding open complex formation, suggesting enhanced supercoiling buildup. Measurements with Gyrase and Topoisomerase I inhibitors suggest hindrance to escape supercoiling buildup at low temperatures. Consistently, similar phenomena occur in energy-depleted cells by DNP at 30 °C. Transient, critically-low temperatures have no long-term consequences, as raising temperature quickly restores transcription rates. We conclude that the chromosomally-integrated $P_{\text{LacO3O1}}$ has higher sensitivity to low temperatures, due to longer-lasting super-coiled states. A lesser active, chromosome-integrated native $\text{lac}$ is shown to be insensitive to Gyrase overexpression, even at critically low temperatures, indicating that the rate of escaping positive supercoiling buildup is temperature and transcription rate dependent. A genome-wide analysis supports this, since cold-shock genes exhibit atypical supercoiling-sensitivities. This phenomenon might partially explain the temperature-sensitivity of some transcriptional programs of $E$. coli.

Escherichia coli has evolved sophisticated regulatory programs to adapt to fluctuating environments that allow tuning gene expression so as to trigger appropriate responses4–7. In general, gene expression regulation occurs during transcription initiation7 and it can be performed, e.g., by transcription factors4,5, which act locally, affecting specific genes, and by $\sigma$ factors6–9, which have more genome-wide effects.

Similarly, environmental changes can affect chromosomal DNA compaction, which is associated to supercoiling10,11 and is regulated by nucleoid associated proteins (NAPs)12,13. Interestingly, changes in DNA compaction have genome-wide effects14–16, causing the expression of some genes to increase while in others it decreases4,16–18.

DNA compaction and supercoiling have distinct effects on plasmid-borne and chromosome integrated genes (see e.g.19). One reason for this is that the chromosome has topologically constrained segments that allow supercoiling buildup12,16–22, as transcription occurs, since this process generates positive supercoiling ahead of the RNA polymerase (RNAP) and negative supercoiling behind it23,24. Meanwhile, plasmids lack discrete constraints. Thus, when positive and negative supercoiling emerge, they freely diffuse in opposite directions and annihilate each other19. Thus, in general, the transcriptional activity in plasmids is only affected by transient constraints due to, e.g., transient protein binding19,25. Exceptions are, e.g., plasmids encoding membrane-associated proteins that, by anchoring to the membrane26–28, can form longer lasting constraints. Other exceptions are plasmids

1Laboratory of Biosystem Dynamics and Multi-Scaled Biodata Analysis and Modelling Research Community, Faculty of Medicine and Health Technology, Tampere University, Korkeakoulunkatu 7, 33720, Tampere, Finland. 2CA3 CTS/UNINOVA. Faculdade de Ciências e Tecnologia, Universidade Nova de Lisboa, Quinta da Torre, 2829-516, Caparica, Portugal. 3Gene Bridges, Im Neuenheimer Feld 584, 69120, Heidelberg, Germany. 4Biostatistics Department, Harvard T.H. Chan School of Public Health, Boston, MA, 02115, USA. 5Infectious Disease and Microbiome Program, Broad Institute, Cambridge, MA, 02142, USA. Correspondence and requests for materials should be addressed to A.S.R. (email: andre.sanchesribeiro@tuni.fi)
carrying tandem copies of one or two DNA-binding sites and plasmids carrying the T7 promoter, when expressed in topA mutant strains. Nevertheless, it is worth noting that in vivo measurements suggest that, prior to annihilation, transient supercoiling changes can influence transcription rates of both plasmid-borne and chromosomally-integrated promoters.

Temperature shifts affect DNA supercoiling directly as well as indirectly, e.g., by affecting the interactivity between NAPs and chromosomal DNA. This may explain why temperature down-shifts affect the activity of most chromosomal genes in E. coli.

Another temperature-dependent event in transcription is promoter escape, the stage at which the RNAP is freed from the promoter and moves downstream towards the elongation region of the DNA template. The stronger the binding between the RNAP and the promoter, the longer it usually takes for the RNAP to escape the promoter and begin elongation. One reason for this is that, for escape to succeed, the RNAP needs to pull a sufficient amount of downstream DNA into itself (so as to reach its active center), which involves breaking interactions between the RNAP and the promoter, and between the RNAP and initiation factors, which are energy dependent processes.

Given the above, we hypothesized that plasmid-borne and chromosomally-integrated genes can differ in sensitivity to temperature shifts and that these differences may be promoter strength-dependent. To test this, we compared quantitatively the effects of temperature shifts on the in vivo kinetics of transcription of the promoter, when on a plasmid and when chromosomally-integrated (Materials and Methods). Further, we assessed the effects of the native promoter, whose transcription rate is weaker than P_LacO3O1, although we located it in the same position in the chromosome.

For this, we used the MS2-GFP RNA tagging technique in E. coli, along with a recently proposed methodology to resolve the rate-limiting steps governing the in vivo dynamics of initiation of prokaryotic promoters (similar to established steady-state assays to resolve the in vitro dynamics). Further, we studied this process at critically low temperatures (below 23 °C), a regime in which most cellular processes exhibit significant differences due to, e.g., globally-altered transcription rates and increased cytoplasmic viscosity. Using these techniques, we characterized, with single-RNA sensitivity, the RNA production dynamics of these constructs at various temperatures, as well as in the presence of Gyrase and Topoisomerase I inhibitors and of DNP-based energy depletion. Also, we made use of stochastic modelling to show that the observed differences in transcription kinetics between chromosome and plasmid integrated promoters at low temperatures are consistent with current stochastic models of transcription initiation that account for supercoiling buildup, provided that such low temperatures result in the hindrance to escape from DNA super-coiling. Finally, we made use of information of what genes in E. coli have their activity induced following cold-shocks and of what genes are supercoiling sensitive, to assess if these two features are strongly correlated, as our results would suggest.

Results

We studied at the single-RNA level if the kinetics of RNA production under the control of P_LacO3O1 differs in response to temperature changes when the gene is single-copy F-plasmid-borne and when it is chromosomally-integrated. For this, we made use of two identical constructs under the control of the P_LacO3O1 Promoter coding for multiple bindings sites for MS2-GFP followed by the coding region of mCherry.

Both constructs, shown in Figs S1 and S2, are functional and responsive to the inducer (Fig. 1). Also, control tests were performed to verify that spots detected in microscopy images correspond to MS2-GFP tagged RNA molecules (Fig. S3) and that, once appearing, their intensity does not change significantly during the measurement time (Supplementary section “Control tests of the RNA counting method”), as this would affect the counting of MS2-GFP tagged RNA molecules in each cell.

For each strain (carrying the target gene in the plasmid or in the chromosome) and each temperature condition, we performed 3 or more biological repeats and counted RNA-spots (MS2-GFP tagged RNAs) in each cell from the microscopy images (Supplementary section “Image Analysis”). As we did not find statistically significant differences between repeats, the results shown here are from cells from all replicates.

Since the two strains are not subject to the same antibiotics (only cells carrying the target gene in the single-copy plasmid are subject to chloramphenicol, see Methods), we tested whether their growth curves differ. Results in Fig. S5 show that these curves are not distinguishable. We also tested whether the two strains produce similar levels of MS2-GFP reporter proteins (as differences could result in different ability to count target RNAs). For this, inducing only the reporter gene, we measured the background fluorescence intensity of cells of both strains (almost exclusively due to MS2-GFP reporters). We then compared the two distributions of single-cell background fluorescence intensity by a Kolmogorov-Smirnov (KS) test of statistical significance and found no significant difference (p-value of 0.46).

**Induction of gene expression is similar, but not identical, in the chromosome and plasmid-integrated constructs.** For each construct (chromosome integrated and single-copy plasmid borne) and IPTG concentration, we quantified the integer-valued RNA numbers in live cells at 30 °C by microscopy imaging. 1 hour after induction of the target promoter by IPTG (Materials and Methods). From Fig. 1, we find that the mean integer-valued RNA numbers per cell, relative to maximum induction (1 mM IPTG) exhibits a similar fold change in both constructs (~4 for the plasmid and ~5 for the chromosome construct), as in previous studies. Also, for both constructs, the transcription rate does not increase beyond 500 µM IPTG, i.e., 1 mM IPTG suffices for full induction, in agreement with previous studies.

However, the RNA production kinetics of the two constructs differs in some aspects. First, from Table 1 and Fig. 1, mean integer-valued RNA numbers per cell are higher in the plasmid construct for all induction conditions. Also, the increase in RNA numbers with IPTG concentration differs. Namely, from Table S3, in the plasmid...
The RNA numbers increase gradually as IPTG is increased, allowing most conditions to differ significantly in a statistical sense, while in the chromosome construct the RNA numbers per cell only differ significantly between 0 µM IPTG and the other conditions (for 50 µM IPTG or higher, there is little increase in RNA numbers with additional increases in IPTG concentrations).

Nevertheless, the same model of transcription (Supplementary Information, section “Model of transcription kinetics”, reactions 1–3) fits both constructs, as tuning $k_{c}$ and/or $k_{unlock}$ suffices to account for differences between them in RNA numbers at 30 ºC.

### Table 1. Number of cells observed, mean, and squared coefficient of variation (CV²) of the absolute integer-valued RNA numbers per cell for the chromosome-integrated and the plasmid-integrated constructs, when induced by 1 mM IPTG. Cells are induced and kept at 30 ºC, 27 ºC, 23 ºC, 20 ºC, 16 ºC and 10 ºC for 60 minutes prior to the acquisition of the results. Results are obtained from 3 biological repeats. Since these exhibited no statistically significant differences, the results presented here are composed of the data from the 3 biological replicates.

| Condition | No. cells | Mean integer-valued RNA no. per cell | CV² |
|-----------|-----------|-------------------------------------|-----|
| **Chromosome construct** | | | |
| 30 ºC | 645 | 2.08 | 2.60 |
| 27 ºC | 632 | 2.00 | 1.93 |
| 23 ºC | 668 | 1.74 | 2.30 |
| 20 ºC | 646 | 0.59 | 7.17 |
| 16 ºC | 668 | 0.22 | 15.81 |
| 10 ºC | 648 | 0.25 | 16.12 |
| **Plasmid construct** | | | |
| 30 ºC | 675 | 2.86 | 1.06 |
| 27 ºC | 654 | 2.46 | 1.42 |
| 23 ºC | 665 | 1.63 | 2.73 |
| 20 ºC | 660 | 1.61 | 3.08 |
| 16 ºC | 663 | 1.50 | 2.99 |
| 10 ºC | 676 | 1.35 | 3.46 |
Transcription by the chromosome-integrated construct is noisier at lower temperatures. We next studied if temperature changes affect differently the chromosome and plasmid constructs. We measured integer-valued RNA numbers in cells under full induction (1 mM IPTG) by microscopy at various temperatures (30, 27, 23, 20, 16 and 10°C). For each condition, from the absolute integer-valued RNA numbers in each cell, we calculated the mean and squared coefficient of variation (CV²) of the RNA numbers in single cells (Table 1). To assess if the RNA production kinetics differs with temperature and between the two constructs, we performed KS tests. The P values comparing the single-cell distributions of RNA numbers between conditions for the chromosome and plasmid constructs and between the constructs at each temperature are shown in Tables S4 and S5, respectively.

From Tables 1 and S4, we find that P_LacO3O1, when in the single-copy plasmid, is highly responsive to temperature decreases until 23°C. Below this temperatures, changes in RNA numbers are only significant for temperature shifts wider than those considered in Table S4 (e.g. p < 0.01 for 23°C and 10°C, not shown in Table S4). This behavior is in line with previous reports for the P_tet and for the P_Lac-Ara-1 promoters, also on single-copy plasmids.

Meanwhile, when chromosome-integrated, P_LacO3O1 activity decreases significantly for a wider range of temperatures. Namely, differences are detectable between all pairs of neighboring conditions, except between 16°C and 10°C. These results are supported by those in Table S5. The P values of the KS tests indicate that, below 23°C, the plasmid and chromosome constructs differ from one another in all temperatures. For 23°C and above, they only differ at 30°C. From this and Table 1, we conclude that the activity of the chromosome-integrated promoter is more heavily reduced as temperature is lowered, and that it remains sensitive to a wider range of temperature shifts.

To validate these results, we used RT-qPCR (Supplementary Information) to obtain the mean RNA numbers relative to the control (30°C) in cells under full induction (1 mM of IPTG) at 23, 16 and 10°C (Fig. S6). From Table 1, we calculated the same quantities from the microscopy measurements. Overall, both the chromosome and plasmid constructs exhibit the same qualitative behavior as temperature decreases when measured by microscopy and RT-qPCR.

We next assessed if the weaker transcriptional activity of the chromosome-integrated promoter at the lowest temperatures could be explained by changes in the spatial distribution of RNAPs or of the nucleoids (Supplementary Information, section “Nucleoid staining with DAPI”). Measurements at 10°C and 30°C (Fig. S7) show no significant differences in these two features, allowing rejecting these hypotheses.

We also performed two additional tests for cells with the chromosome-integrated P_LacO3O1. First, as the mean integer-valued RNA numbers per cell in induced cells at 10°C (Table 1) appears to be smaller than in non-induced cells at 30°C (Table S1), we tested if this difference is statistically significant by performing a KS test between the distributions of single-cell RNA numbers in the two conditions. We obtained a p-value of 0.99 and, thus, we conclude that the RNA numbers in the two conditions do not differ, in a statistical sense (p-value larger than 0.01), implying that induced cells at 10°C produce at least as much RNAs as non-induced cells at 30°C. Second, we tested whether, at 10°C, RNA numbers differ between induced and non-induced cells. A KS test between the distributions of RNA numbers in individual cells in the two conditions (Fig. 2 and Table S1) shows that they can be distinguished in a statistical sense. Thus, we concluded that induction at 10°C tangibly increases the RNA production rate of the chromosome-integrated P_LacO3O1.

Mean relative time prior to commitment to transcription increases in the chromosome-integrated construct at low temperature. To investigate why the two constructs responded differently to lowering temperatures, we assessed whether the changes in the kinetics of transcription with decreasing temperature occur prior to or following the commitment to open complex formation (Supplementary Information, section “Model of transcription kinetics”), by making use of Lineweaver–Burk plots of the inverse of the RNA production rate against the inverse of the RNAP concentration (Supplementary sections “Lineweaver–Burk Plots” and “Tuning intracellular RNAP concentrations”).

For this, we measured RNAP levels in individual cells in each temperature condition, and verified that they differ statistically between conditions (KS-tests in Table S8). From these, we obtained the inverse of the RNAP concentrations relative to the 1X control condition (Table S7). Next, for the same condition, using the region of the target gene coding for mCherry (Fig. S2), we measured the RNA production rates of the two constructs at 10 and 30°C by RT-qPCR, and obtained the inverse of these values (Table S9). Combining both measurements, we obtained Lineweaver–Burk plots for each construct and the two temperature conditions (example Fig. S10).

Next, from these plots, using the same methodology as in, we estimated the mean fraction of time between consecutive transcription events taken by the steps preceding (t_pre) and following (t_post) the commitment to open complex formation at the highest and lowest temperature, with Δt being the mean time length between consecutive transcription events in individual cells (Supplementary Information, section “Relative mean duration prior to and following commitment to transcription”).

Results in Table 2 show that, in all 4 conditions, the most rate-limiting events occur after commitment to open complex formation. However, Table 2 also informs that the lowering temperatures do not cause the same effect in the two constructs.

In particular, in the plasmid construct, in agreement with previous in vitro measurements for the synthetic P_LacUV5 promoter, the reduction in RNA production rate with lowering temperature is mostly due to a reduction in the rate of the events after commitment to open complex formation (with t_post increasing from being 92% to 98% of the Δt as temperature is lowered). Meanwhile, in the chromosome construct, the opposite occurs (with t_post decreasing from 91% to 73% of the Δt as temperature is lowered) suggesting that, in this construct, the events...
whose rates were most reduced occur prior to commitment to open complex formation, provided that the RNA production rate decreases with lowering temperature (as is the case, see Table 1).

Local DNA supercoiling in the chromosomally-integrated gene drives the differences between constructs. To explain the increased time-length of the events preceding the open complex formation in the chromosome-integrated construct at lower temperatures, we considered the model of transcription initiation (Supplementary Information, reactions 1–3). This model allows for this, provided that decreasing temperatures decrease the rate of unlocking (k\textsubscript{unlock}) from locked promoter states (reaction 3, Supplementary Information), or decrease the rate of unbinding of a repressor from a promoter (k\textsubscript{ON}, reaction 2 in Supplementary Information), or both. Either of these possibilities is physically possible since lowering temperatures could affect the efficiency of repressors (see e.g.\textsuperscript{48}), DNA packaging (known to differ between plasmid and chromosomes\textsuperscript{49}, or DNA super-coiling\textsuperscript{35} (known to affect both packaging\textsuperscript{10} and transcription\textsuperscript{19,30,50,51}).

A third possibility would be that decreasing temperature modified the kinetics of closed complex formation, causing increased relative duration of this event, e.g. due to reduced k\textsubscript{1} or k\textsubscript{2}, or instead increased k\textsubscript{−1}. However, this would result in reduced noise in RNA production\textsuperscript{40,52} and thus reduced CV\textsuperscript{2} in RNA numbers in individual

![Graph](image-url)
cells (since, at 30 °C, most time between transcription events is spent in open complex formation, Table 2). The data on CV² in RNA numbers in Table 1 disproves this possibility.

Meanwhile, in the first possibility, where kₜᵢᵦlock or kᵣᵢᵦ are decreased with decreasing temperature, this would result in increased noise in RNA production⁵⁵ and, thus, increased CV² in RNA numbers in individual cells, which was observed (Table 1).

To determine whether it is kₜᵢᵦlock or kᵣᵦ that is decreased, consider that a change in repressors’ efficiency with temperature (i.e. a change in kᵣᵦ) should affect both the chromosome and plasmid constructs similarly since both constructs are affected by this mechanism. However, we observed divergent responses between these two constructs to the lowest temperatures, with the plasmid-borne construct being unable to turn off its RNA production as efficiently as the chromosome-integrated construct (Table 1). Thus, we conclude that the stronger decrease in the chromosome construct in RNA production rate with lowering temperature (at the lowest temperature conditions tested) is likely due to an increased amount of time required to remove the promoter from the locked state, which does not occur in the plasmid construct (i.e. changes in kₜᵢᵦlock with lowering temperature are the most likely explanation for the observed behaviors).

It is further possible to assess if the changes in kₜᵢᵦlock, causing different behaviors of the two constructs in response to lowering temperatures, are associated to DNA packaging and/or super-coiling. For that, we measured the nucleoid size in cells with one nucleoid (Supplementary Information) in the various temperature conditions. If decreasing temperature (in the ranges shown in Table 1) affects DNA packaging significantly, we expect differences in the mean and/or variability of the nucleoid size. However, we found no significant differences between 10 °C and 30 °C (Table S10). Similar results were reported in⁴¹. Thus, we discard DNA packaging as the main cause for the differences between chromosome and plasmid response to temperature shifts.

Given all of the above, we hypothesize that the different response of the chromosome and plasmid-integrated genes with lowering temperature is due to an increased rate of accumulation of local DNA supercoiling in the chromosome-integrated gene, which increases the escape times from locked states (reactions 1 and 3, Supplementary Information).

To validate this hypothesis, we performed several experiments. First, we compared the numbers of RNAs produced over time by cells of each strain. We expect this number to increase near-constantly in the cells carrying the plasmid-borne gene, but not in the cells of the other strain. For this, from the moment of activation of the target gene (t = 0 minutes), we measured integer-valued RNA numbers in individual cells at 10 °C every 15 minutes for 90 minutes (for each time point, new cells were taken from the original culture). If the weaker activity of the chromosome-integrated promoter is due to increased propensity to be in the locked state due to the accumulation of DNA super-coiling, we expect its transcription activity to be blocked after a few events. At a population level, this would result in a sharp decrease in the rate of increase of RNA numbers in the cells, some time after the start of the measurements. Meanwhile, in the plasmid construct, we expect a constant RNA production rate over time, due to the lack of accumulation of local DNA super-coiling⁴⁹. Results in Fig. 2A confirm these predictions. Cells at 10 °C with the chromosome construct only exhibit production in the first 30 minutes, while the plasmid construct shows approximately constant RNA production rate throughout the measurement.

We also performed measurements at 30 °C. Given the similar dynamics of transcription of the two constructs in this condition (Table 2), we expect the RNA production rate to be constant in time in both constructs. Results in Fig. 2B confirm this.

To further test the hypothesis, we next compared the activity of the two constructs at 30 °C when subjecting cells to Novobiocin, an inhibitor of Gyrase activity (Methods)³⁹,⁵⁴. Gyrase releases positive supercoiling⁵⁵ but not negative supercoiling⁶⁰. According to the twin-supercoiled-domain model⁵⁴, which predicts that negative/positive supercoils should accumulate in the absence of supercoil-relaxing enzymes, we expect cells with the chromosome construct to exhibit a similar behavior as when at 10 °C. Meanwhile, cells with the plasmid construct should again exhibit a constant rate of transcription over time⁴⁹. Figure 3A confirms these predictions.

In this regard, in both strains, the gene acrA is present, and thus, Novobiocin is not expected to affect cell division rates⁶⁰. To test this, we measured cell growth rates by OD₆₀₀ for varying Novobiocin concentrations (0, 50, 75, 100 and 150 ng/μl). We found the growth rates to not differ significantly between conditions (data not shown). These results also show that 100 ng/μl Novobiocin concentration suffices to affect (but not halt) the transcription rate of the chromosome-integrated gene (compare the results for this construct in Figs 2B and 3A, at 30 °C).

Subsequently, we subject cells with the chromosome construct to Novobiocin when at 10 °C. Results in Fig. 3B, when compared to Figs 2A and 3A, show that transcription in cells carrying the chromosome is more strongly blocked when combining Novobiocin and low temperatures. I.e. while at 10 °C alone and subject to Novobiocin alone, RNA numbers increase by a factor of 4 (from 0.25 to 1) in a period of 90 min., when subjecting cells to both 10 °C and Novobiocin, the RNA numbers increase only by a factor of 2 (from 0.5 to 1) in the same period of time. Meanwhile, in cells with the plasmid construct, we observe the same RNA production as in Fig. 2A, meaning that, in these cells, Novobiocin has no effect at either temperature. Given this, we suggest that the transcription activity of the chromosome-integrated promoter at 10 °C is hampered by an increased difficulty in unblocking the DNA from supercoiled states, rather than due to a loss of functionality...
Finally, in comparison, subjecting cells with the single-copy F-plasmid to Topotecan causes, qualitatively, the same behavior as adding Novobiocin (at 30 °C and 10 °C) (Fig. 3A–D).

Promoter escape from supercoiling buildup is similarly hampered if cellular energy is depleted.

If the escape from DNA supercoiling buildup in the chromosomally-integrated construct at low temperatures is due to energy deficiency at low temperatures (the energy required for the necessary endothermic reactions to occur should be higher in such conditions), it should be possible to mimic the phenomena by, instead of lowering temperature, depleting cells of energy via DNP treatment45 (Methods). In particular, we expect cells subject to this treatment to, even at 30 °C, be less able to maintain the chromosome integrated promoter active over time when compared to the control, similar to when at 10 °C.

To test this, we subjected cells to DNP for 90 minutes (at 30 °C) prior to imaging (Methods). As expected, we observed similar RNA production dynamics (Fig. 4), as in untreated cells at 10 °C with a chromosome integrated PLacO3O1 (Fig. 2A). I.e., beyond 30 minutes, there is little to no transcription, suggesting that, in this condition, the activity is also being hampered by increased difficulty in escaping from supercoiled states.

Low temperatures have no long-term consequences on transcription blocking by DNA supercoiling.

We performed an additional test to support the hypothesis that the escape from DNA supercoiling buildup in the chromosome construct at low temperatures is due to energy deficiency. Namely, we hypothesized that changing temperature to near-optimal conditions (e.g. 30 °C) should restore the cells’ ability to relax DNA supercoiling (as at higher temperature this process is expected to require less energy). To test this, we subjected cells with a
chromosome integrated P_{lacO3O1} to two temperature shifts, first from high (30 °C) to low (10 °C) and then from low (10 °C) to high (30 °C), and measured the mean integer-valued RNA numbers in the cells over time. Results in Fig. 5 show that both temperature shifts caused smooth transitions in the RNA production rates that are consistent with changes in the kinetics of locking/unlocking of promoters from positive supercoiling buildup. In detail, cells at 30 °C have constant RNA production, as shown previously. Once temperature is shifted to 10 °C, after 15–30 minutes, little to no RNA production is observed (as in Fig. 2A). More importantly, once high temperatures are restored (to 30 °C), RNA production is quickly restored to nearly the original rate. The fast transition between behaviors and the ability to quickly restore the original dynamics reinforce the conclusion that the activity of the chromosome integrated promoter at 10 °C is hampered by an increased difficulty in unblocking the promoter from supercoiled states (due to an increase in the energy required).

Note that in this particular experiment, following the shift from 30 °C to 10 °C, it does not follow a transient of ~15–30 minutes of reduced transcription activity that is visible in Figs 2, 3 and 4. This is because, in this case, when the shift occurs, the cells already contain sufficient IPTG to achieve full transcription rates, while in the previous experiments the inducer was added immediately before the microscopy measurements began, and thus, a transient time to reach quasi-equilibrium RNA production rates is expected, due to the non-negligible time that cells need to intake inducers from the media, particularly at low temperatures. In the case of IPTG, previous measurements suggest that this transient is ~15–30 minutes long, in agreement with the results in Figs 2A, 3A–D and 4.

Stochastic modelling also suggests increased long-lasting super-coiled states at critically low temperatures to be the cause for enhanced sensitivity to shifts to critically low temperatures. We tested whether the increase in the expected time for promoters to escape from a supercoiling state across the cell population is, in accordance with current stochastic models of transcription in E. coli, a plausible explanation for the change with decreasing temperature in the average RNA numbers over time in cells with the chromosome integrated promoter (Fig. 2). For this, we use the stochastic model of transcription initiation (Supplementary Information, reactions 1–3), derived from multiple studies, including genome-wide studies of variability in transcript counts and studies of the transcription dynamics of individual genes.

All parameter values (Table S11) are from single-cell, single-RNA empirical data on the activity of lac derivative promoters. Mean RNAP numbers are set to correspond to the RNAp concentration reported in. Finally, from the results above, we assume that the increase in $k_{\text{unlock}}$ as temperature decreases (Table 2) is mostly due to a decrease in $k_{\text{unlock}}$. Thus, the remaining rate constants are, for simplicity, unchanged.

For each value of $k_{\text{unlock}}$ tested, we performed 500 independent simulations, each 75 minutes long. Data was collected every 15 minutes, as in the experiments (Fig. 2). The values of $k_{\text{unlock}}$ were selected as follows: the highest value, corresponding to high temperatures (30 °C), is reported in. This value was then gradually lowered until the mean number of RNAs per cell at the end of the measurement period was similar to that observed in cells at 10 °C.

We assessed if the model was able to reproduce the observed RNA numbers over time at both high and low temperatures, and if there is a gradual behavioral change between these extreme conditions. For this, the initial
numbers of all molecular species were set to zero, with the exception of P_ON (set to 1, corresponding to one active promoter per cell), RNAp (as noted above), and RNA. Initial RNA numbers were drawn randomly from a Poisson distribution (0.7 RNA/cell) to match the here observed outcome of spurious RNA production events. We observed also (empirically, Supplementary Table S1) that this number did not differ with temperature, as expected, since, prior to moment 0, cells were at the same temperature (30 °C) in both measurements.

In Fig. 6, we compared the results of the model with those in Fig. 2A (10 °C) and Fig. 2B (30 °C) for the chromosome-integrated promoter. For simplicity, as noted, we ignored the first time moment of the empirical data (0 minutes following induction) since, at this stage, the cells did not yet have fully active transcription61. This removed the need to model the intake process for the inducers61.

Results in Fig. 6 support the earlier conclusions. The accuracy with which the model reproduces the measurements suggests that the difference in mean RNA production rates between cells with the chromosome-integrated promoter at critically low (10 °C) and at high (30 °C) temperatures can be explained by a reduced ability to release chromosome-integrated promoters from the effects of DNA supercoiling at critically low temperatures.

Finally, note that setting k_unlock to infinite in reaction 3 (equivalent to having a model that does not allow promoter locking) results in a similar behavior to that of the plasmid-borne construct, and thus to the chromosome-integrated promoter at 30 °C (data not shown).

**Transcription by a lesser active chromosome-integrated promoter construct is less sensitive to gyrase overexpression and temperature shifts.** The influence of positive supercoiling buildup on the dynamics of a chromosome-integrated gene differs with its location on the chromosome due to, among other, differences in the expression rates of operons in different DNA loops64, which will cause the effects of positive supercoiling buildup to differ. Meanwhile, we observed that temperature downshifts have weaker effects on the chromosome-integrated promoter at critically low (10 °C) and at high (30 °C) temperatures can be explained by a reduced ability to release chromosome-integrated promoters from the effects of DNA supercoiling at critically low temperatures.

Finally, note that setting k_unlock to infinite in reaction 3 (equivalent to having a model that does not allow promoter locking) results in a similar behavior to that of the plasmid-borne construct, and thus to the chromosome integrated promoter at 30 °C (data not shown).
To test the first hypothesis, we compared the effects of overexpressing Gyrase (Methods) on the mean RNA numbers in cells with the chromosome integrated promoter at 10 °C, when and when not induced. When induced (1 mM IPTG), we find that overexpressing Gyrase increases the mean number of RNAs per cell by 154%. Meanwhile, when not induced (0 mM IPTG), this mean number only increases by 107%. This agrees with the hypothesis that, the higher the expression rate, the bigger is the impact of lowering temperatures due to decreased probability that Gyrases can act in between transcription events.

To test the second hypothesis, we replaced the P LacO3O1 chromosome-integrated promoter by the native lac, which has a similar repression-activation mechanism (Methods) but weaker activity. First, to confirm that its transcription rate is weaker than P LacO3O1, we measured the mean number of RNA molecules per cell, 1 hour after induction, at 30 °C and at 10 °C, under full induction. We found that the native lac produces ~85% less RNAs than PLacO3O1 (both at 30 °C and at 10 °C). Next, to test the hypothesis, we compared the effects of overexpressing Gyrase in cells at 10 °C, when carrying the chromosome-integrated P LacO3O1 and when carrying the chromosome-integrated native lac. Given the above, we expect the overexpression of Gyrase to have a weaker impact on the transcription rate of the native lac. Results in Table S16 confirm this, showing that the overexpression of Gyrase does not significantly affect its number of transcripts, similarly to the plasmid-borne PLacO3O1. Meanwhile, measurements at 30 °C showed, as expected, that no construct is significantly affected by overexpressing Gyrase (Table S16).

Cold shock-inducible genes have atypical sensitivity to supercoiling. Given the above results that show that temperature downshifts can affect the escape times from positive supercoiling buildup, one can hypothesize that genes essential for adaptation to low temperatures (e.g., cold shock inducible genes) should have atypical sensitivity to supercoiling buildup. I.e., these genes should be either significantly more responsive or significantly less responsive to supercoiling buildup than randomly selected genes.

To test this, we combined information on E. coli’s genome-wide sensitivity to supercoiling buildup (enhanced by repressing Gyrase) with information on genome-wide responsiveness to cold shocks. Cold-shock responsive genes were classified as being associated to transient or to prolonged responses. We find that the genes responsible for transient responses (70 genes) are two times more likely (~14% chances) to be supercoiling sensitive than a gene selected randomly from the genome (~7% chances, since there are 306 supercoiling-sensitive genes out of 4452 genes). Meanwhile, genes responsible for long-term responses to cold shocks (35 genes) appear to be impervious to supercoiling (0% chances). The probability that these results would occur by random chance are 1.8% in the case of genes responsible for transient responses, and 8.5% in the case of genes responsible for long-term responses, as estimated from bootstrapped distributions (70 genes each) from 10000 random resamples with replacement using a non-parametric bootstrap method. We thus conclude that both classes of cold shock-related genes have atypical sensitivity (even though opposite) to supercoiling.
Discussion

Temperature-driven changes in genomic DNA supercoiling may be one of E. coli’s mechanisms for sensing and responding to temperature shifts[18]. A previous study[18] using single-molecule mRNA FISH assays to show that, at optimal temperatures, DNA supercoiling buildup eventually halts transcription initiation, which can be resumed upon release of the supercoiling by Gyrase. This buildup only has significant effects in chromosomal genes (in highly expressing operons[41]), since plasmids lack discrete topological constraints, allowing the negative and positive supercoiling emerging in transcription to diffuse freely in opposite directions, until nullifying one another[19].

It is expected that both the time needed for Gyrase to find DNA regions requiring intervention as well as the energy required for this process are temperature-dependent. As such, it may be that the effects of lowering temperature differ both between chromosome-integrated and plasmid-borne genes, as well as between chromosome-integrated genes with different transcription rates.

We studied this phenomenon in the P_{LacO3O1} promoter, under full induction, as a function of temperature (this promoter, due to lacking the O_{2} site and being under full induction, is not expected to form significant discrete topological constraints[49]). We showed that its response to temperature downshifts differs, depending on whether it is chromosomally-integrated or plasmid-borne. Specifically, the chromosome-integrated gene is more sensitive to critically low temperatures (below 23°C), becoming weaker and noisier, and having comparatively longer-length steps preceding the open complex formation than the single-copy plasmid-borne gene. In particular, in these conditions, similar to when inhibiting Gyrase or Topoisomerase I, the transcription activity of the chromosomally-integrated gene is hampered, due to increased time to escape from positive supercoiling buildup. Overall, the results suggest that promoters’ in E. coli differ in sensitivity to shifts to critically low temperatures when chromosome-integrated and when plasmid-borne, due to the temperature-dependence of the kinetics of promoter locking due to positive supercoiling buildup[50]. Simulations of a stochastic model of transcription with a temperature-dependent escape from the locked promoter state are consistent with these observations.

An indirect evidence for the existence of promoter blocking at low temperatures in the chromosome-integrated gene is the high cell-to-cell variability in integer-valued RNA numbers at 10 °C, when compared to the plasmid-borne gene (Table 1). This variability, one hour following induction, suggests that the kinetics of RNA production differed significantly between cells with the chromosome-integrated gene (in agreement with the occurrence of promoter locking in some cells and transcription activity in others). In particular, we estimate that cells with active promoters produced ~1 to 4 RNAs (consistent with mean RNA numbers of native active genes in E. coli[52]), while cells with locked promoters produced ~0 to 2 RNAs at most.

Importantly, we found evidence that the hampering of escape of P_{LacO3O1} from supercoiled buildup states at low temperatures is an energy-associated increase in difficulty to release locked promoters, rather than due to a reduced functionality of Gyrase or Topoisomerases. First, inhibiting the activity of these proteins with Novobiocin and Topotecan at 10°C further reduces transcription reactivation (agreeing with the suggestion that these proteins evolved to act in response to cold-shocks[53,54]). Also, cells with energy depletion due to DNP treatment exhibit a similar transcription dynamics to when at low temperatures. Finally, subjecting cells to consecutive shifts between high and low temperatures results in smooth transitions in the transcription dynamics, which would not be possible if the population of Gyrase or Topoisomerases I had to be renewed.

We do not know whether the transcriptional halting at low temperatures of the chromosome-integrated P_{LacO3O1} is enhanced by the known overexpression at low temperatures of H-NS and similar NAP proteins present in the nucleoid, which, in these conditions, appear to selectivity inhibit early step(s) in transcription initiation by binding to the promoter and acting as transcriptional repressors[49]. However, the fast recovery of the kinetics of RNA production under the control of P_{LacO3O1} observed in Fig. 5, when changing temperature from 10°C and 30°C, could be an indication that these proteins are not involved in the phenomenon observed. Similarly, we also do not know whether there is any influence from stringent response mechanisms. Studies of the roles of, e.g., diskA and ppGpp, may prove to be of value to determine whether, e.g., the biophysical phenomena here reported are affected by these mechanisms.

It is well established that the influence of positive supercoiling buildup on a chromosome-integrated gene differs with its location in the chromosome, because this phenomenon is sensitive to the expression rates of the operons of a DNA loop[54]. The present study indirectly supports this, by showing that the sensitivity of a gene differs with its own transcription rate. In particular, when replacing the promoter (P_{LacO3O1}) controlling the chromosome-integrated gene by the native lac, of weaker activity, we observed much weaker effects when lowering temperature. Also, overexpressing Gyrase no longer had an effect, suggesting absence of positive supercoiling buildup. This can be explained by the gene’s longer time intervals between transcription events, which allow Gyrase to act so as to escape from positive supercoiling buildup[53] prior to these having a significant impact. Thus, we hypothesize that the effects of temperature downshifts on positive supercoiling buildup are promoter activity dependent.

Since the effects of temperature downshifts differ with the kinetics of positive supercoiling buildup, which itself differs with the gene location in the chromosome[54] and own activity level (among other variables), one can expect genome-wide heterogeneity in the response to temperature downshifts. Thus, it is possible that genes associated to the responses to cold-shocks have atypical sensitivity to positive supercoiling buildup. When investigating this possibility, we found that the number of genes that are responsible for transient responses to cold-shock and also have high-sensitivity to supercoiling buildup is above-expected. Further, the number of genes that are responsible for long-term responses that also have high-sensitivity is below-expected.

We interpret these results as follows. Long-term activity following a cold-shock should be facilitated if a gene is only weakly affected by supercoiling buildup (as observed here when comparing the kinetics of the chromosomally integrated P_{LacO3O1} and native lac). Meanwhile, above average sensitivity to supercoiling is expected to contribute to (or be responsible for) the transient nature of the response exhibited by some genes associated to cold-shock. In particular, for a gene to remain active for ~1 hour[53] and then have its activity reduced, there is a need for a mechanism of slow repression, which would be consistent to the ‘repression’ caused by positive...
supercoiling buildup (similar to what is observed in Figs 2, 3 and 4, with activity shutdown occurring after 30–60 minutes).

Overall, our results suggest that it may be possible that the temperature-dependence of the kinetics of promoter locking due to positive supercoiling buildup could be used as a means to introduce temperature sensitivity in some transcriptional programs of E. coli. In this regard, this may be of value to chromosome-integrated, synthetic genetic circuits with temperature-sensitivity. For example, if locating component genes on different DNA loops (with different levels of transcriptional activity), it may be possible for temperature shifts to generate heterogeneity in the responses of the component genes, which can be used to trigger changes in the state of the circuit.

Materials and Methods

E. coli strain BW25993 (lacIq hsdR514 ΔaraBADAH33 ΔrhaBADLD78) cells carry the target and reporter genes. The target gene is controlled by P_LacO301 and codes for an array of 48 binding sites for a modified viral coat protein, MS2-GFP [27-28]. P_LacO301, inducible by IPTG, was engineered from the E. coli native lac promoter, by removing the O₂ repressor binding site downstream of the transcription start site [29]. Due to lacking the site O₂, we expect little to no formation of significant topological constraints [30]. Also, the repression strength of LacI is expected to be 2–3 fold weaker than on the wild-type lac promoter [31]. Finally, in one measurement, we made use of a chromosome-integrated native lac promoter, also followed by an identical array of binding sites for MS2-GFP.

To compare the RNA production rate of P_LacO301 when single-copy plasmid-borne and when chromosome-integrated, two strains were engineered from the original BW25993. One carries a single copy full F-plasmid (~11 kbp) [32], pBELOBAC11 (target plasmid), unknown to form long-lasting bounds to the membrane and originally responsible for the expression of transient DNA-binding proteins [33,34]. In this, we inserted the target gene, P_LacO301, coding for the bindings sites for MS2-GFP (Fig. S1 in Supplementary Information). In the other strain, the target gene, controlled by P_LacO301, was integrated into the lac gene locus of E. coli’s genome using Red/ET recombination (performed by Gene Bridges, Heidelberg, Germany) (Fig. S2 and Tables S13 and S14 in Supplementary Information).

Both strains were also transformed with a medium copy reporter plasmid pZA25-GFP [35] (kind gift from Orna Amster-Choder, Hebrew University of Jerusalem, Israel), coding for the reporter protein MS2-GFP controlled by the BAD promoter. The multiple MS2-GFP binding sites in the target RNAs and the strong binding affinity of each site allow target RNAs to appear as bright spots, soon after produced (Fig. S3) [36]. Their maximum fluorescence is reached in less than 1 min [37] and, once reached, remains constant for hours [38], due to lack of interference from RNA degradation [39,40].

While the strain carrying the target gene in single-copy F-plasmid also contains a native lac promoter in the chromosome (and, thus, has higher number of LacI binding sites overall than the strain carrying the chromosome-integrate target gene as the original lac promoter was replaced by the target one), both strains overexpress LacI, reducing the possibility of significant effects due to shortage of repressors in the strain carrying the F-plasmid. Further, our measurements were conducted under full induction (except the induction curves), further reducing any possibility of effects of differences in number of available repressors.

For overexpressing Gyrase, we engineered a plasmid (pZe11-P_rham-gyrAB, with ampicillin resistance) with the gyrA and gyrB genes under the control of a Rhamnose promoter (Supplementary Section “Gyrase overexpression”).

Cell growth conditions, antibiotics, and means of induction of the target and report genes are described in Supplementary Information (section “Growth Conditions and Induction of the Reporter and Target Gene”).

When analyzing the data, we assumed that, in both strains, only 1 copy of the target gene is present in each cell. This approximation is based on our observation that, 1 hour after starting the measurements, only 15% and 12% of the cells had 2 nucleoids at, respectively, 10 °C and 30 °C (600 cells analyzed per condition) (Supplementary Information, section “Number of promoter copies during the cell lifetime”), suggesting that these cells with the chromosome integrated promoter only carry two copies of the target gene for a relatively short period of time during their lifetime. The same assumption is applied to the cells carrying the gene of interest in the single-copy F-plasmid. This is because F-plasmids replicate at the same time [39] or shortly after [40] the chromosome. Further, we measured the plasmid copy numbers by RT-qPCR relative to the number of chromosomes at any given time (Supplementary Information, section “Plasmid copy number calculation using RT-qPCR”). The measurements showed that the copy number of the single-copy F-plasmid (pBELO) relative to the chromosome copy number is 1.00 and 1.02 at 10 °C and 30 °C, respectively (Fig S9 and Table S12), validating the assumption. Finally, we note that in neither strain did we find any significant difference in the rates of RNA production in the first and second half of the cells lifetime (which would be expected if the replication of the gene occurred early in the cell lifetime, e.g. at midpoint).

We quantified integer-valued RNA numbers in cells with repressed Gyrase activity and with repressed Topoisomerase I activity. For that, we used, respectively, Novobiocin and Topotecan [41,42]. Cells were grown as described in Supplementary Information. Following induction of the reporter gene, cells were incubated at the appropriate temperature (10 °C or 30 °C), at 250 rpm for 15 minutes, prior to induction of the target gene. Afterwards, 1000 μM of IPTG and 100 ng/μl of Novobiocin or 100 μM of Topotecan were added to the cells.

To determine RNA levels in cells treated with 2,4-Dinitrophenol (DNP) (which uncouples the oxidative phosphorylation, causing Adenosine triphosphate depletion) [43] the growth and activation of reporter genes were carried out as described above. Next, 1000 μM of IPTG and 200 μM of DNP were added to the media and cells were incubated at 30 °C.

We used E. coli RL1314 strain to measure RNA polymerases (RNAP) intracellular concentrations, carrying RNAPs fused with GFP (RNAP-GFP) [44]. Changes in fluorescence levels (example image in Fig. S4A) with, e.g.,
media richness, are consistent with RT-PCR (rpoC transcript levels) and plate reading measurements. To visualize the nucleoid we used 4,6-diamidino-2-phenylindole (DAPI) (Fig. S4B).

We performed measurements in cells whose RNAP concentrations differ. To obtain such cells, we employed the method proposed in. In short, we made use of the media differing in Glycerol concentration (Supplementary Information, section “Tuning intracellular RNAP concentrations”). Specifically, cells were grown in media with 0.2, 0.4, 0.6 and 0.8% of Glycerol, denoted as 0.5X, 1X (control), 1.5X and 2X, respectively. These conditions allow cells to differ in RNAP concentration while not differing in mean growth rates (assessed from the OD600 over time by a spectrophotometer), as visible in Fig. S8.

Images acquisition is described in Supplementary Information. It took, on average, ~3 minutes to move cells from the incubator to the microscope, assemble the imaging chamber with slides and cells, and start the observation. Images were then analyzed by the software iCellFusion and CellAging. Simulations of stochastic models of gene expression were performed by SIGNS, a simulator of chemical reaction systems whose dynamics is driven by the Stochastic Simulation Algorithm that allows multi-time-delayed reactions.

References
1. Kannan, G. et al. Rapid acid treatment of Escherichia coli: transcriptomic response and recovery. BMC Microbiol. 8, 37 (2008).
2. Arsène, F., Tomoyasu, T. & Bukau, B. The heat shock response of Escherichia coli. Int. J. Food Microbiol. 55, 3–9 (2000).
3. Brown, D. F. & Busby, S. J. W. Local and global regulation of transcription initiation in bacteria. Nat. Rev. Microbiol. 14, 638–650 (2016).
4. McClure, W. R. Mechanism and control of transcription initiation in prokaryotes. Annu. Rev. Biochem. 54, 171–204 (1985).
5. Brewster, R. C. et al. The transcription factor titration effect dictates level of gene expression. Cell 156, 1312–1323 (2014).
6. Farewell, A., Knütt, K. & Nyström, T. Negative regulation by RpoS: A case of sigma factor competition. Mol. Microbiol. 29, 1039–1051 (1998).
7. Rouvière, P. E. et al. rpoE, the gene encoding the second heat-shock sigma factor, sigma E, in Escherichia coli. EMBO J. 14, 1032–1042 (1995).
8. Dong, T. & Schellhorn, H. E. Global effect of RpoS on gene expression in pathogenic Escherichia coli O157:H7 strain EDL933. BMC Genomics 10, 1–17 (2009).
9. Cho, B.-K., Kim, D., Knight, E. M., Zenger, K. & Palsson, B. O. Genome-scale reconstruction of the sigma factor network in Escherichia coli: topology and functional states. BMC Biol. 12, 4 (2014).
10. Stuger, R. et al. DNA supercoiling by gyrase is linked to nucleoid compaction. Molecular Biology Reports 29, 79–82 (2002).
11. Holmes, V. F. & Cozzarelli, N. R. Closin the ring: links between SMC proteins and chromosome partitioning, condensation, and supercoiling. Proc. Natl. Acad. Sci. USA 97, 1322–4 (2000).
12. Postow, L., Hardy, C. D., Arusuga, J. & Cozzarelli, N. R. Topological domain structure of the Escherichia coli chromosome. Genes Dev. 18, 1766–1779 (2004).
13. Dillon, S. C. & Dorman, C. J. Bacterial nucleoid-associated proteins, nucleoid structure and gene expression. Nat. Rev. Microbiol. 8, 185–195 (2010).
14. Peter, B. J. et al. Genomic transcriptional response to loss of chromosomal supercoiling in Escherichia coli. Genome Biol. 5, R87 (2004).
15. Pruss, G. J. & Drlica, K. DNA supercoiling and prokaryotic transcription. Cell 56, 521–523 (1989).
16. Travers, A. & Mukhbelshivili, G. DNA supercoiling - a global transcriptional regulator for enterobacterial growth? Nat. Rev. Microbiol. 3, 157–169 (2005).
17. Dorman, C. J. DNA supercoiling and bacterial gene expression. Sci. Prog. 89, 151–166 (2006).
18. Dorman, C. J. & Dorman, M. J. DNA supercoiling is a fundamental regulatory principle in the control of bacterial gene expression. Biophys. Rev. 8, 89–100 (2016).
19. Chong, S., Chen, C., Ge, H. & Xie, X. S. Mechanism of Transcriptional Bursting in Bacteria. Cell 158, 314–326 (2014).
20. Hardy, C. D. & Cozzarelli, N. R. A genetic selection for supercoiling mutants of Escherichia coli reveals proteins implicated in chromosome structure. Mol. Microbiol. 57, 1636–1652 (2005).
21. Rovinskiy, N., Agbleke, A. A., Chesnokova, O., Pang, Z. & Higgins, N. P. Rates of Gyrase Supercoiling and Transcription Elongation Control Supercoil Density in a Bacterial Chromosome. PLoS Genet. 8, e1002845 (2012).
22. Higgins, N. P. Species-specific supercoil dynamics of the bacterial nucleoid. Biophys. Rev. 8, 113–121 (2016).
23. Deng, S., Stein, R. A. & Higgins, N. P. Organization of supercoil domains and their reorganization by transcription. Mol. Microbiol. 57, 1511–1521 (2005).
24. Liu, L. F. & Wang, J. C. Supercoiling of the DNA template during transcription. Proc. Natl. Acad. Sci. USA 84, 7024–7027 (1987).
25. Leng, F., Chen, B. & Dunlap, D. D. Dividing a supercoiled DNA molecule into two independent topological domains. Proc. Natl. Acad. Sci. 108, 19973–19978 (2011).
26. Boeke, J. D. & Model, P. A prokaryotic membrane anchor sequence: carboxyl terminus of bacteriophage fi gene III protein retains it in the membrane. Proc. Natl. Acad. Sci. USA 79, 5200–5204 (1982).
27. Pruss, G. J. & Drlica, K. Topoisomerase I mutants: the gene on pBR322 that encodes resistance to tetracycline affects plasmid DNA supercoiling. Proc. Natl. Acad. Sci. USA 83, 8992–6 (1986).
28. Lynch, A. S. & Wang, J. C. Anchoring of DNA to the bacterial cytoplasmic membrane through cotranscriptional synthesis of polypeptides encoding membrane proteins or proteins for export: A mechanism of plasmid hypernegative supercoiling in mutants deficient in DNA topoisomerase I. J. Bacteriol. 175, 1645–1655 (1993).
29. Deng, S., Stein, R. A. & Higgins, N. P. Transcription-induced barriers to supercoil diffusion in the Salmonella typhimurium chromosome. Proc. Natl. Acad. Sci. USA 101, 3398–3403 (2004).
30. Fulcrand, G. et al. DNA supercoiling, a critical signal regulating the basal expression of the lac operon in Escherichia coli. Sci. Rep. 6, 1–12 (2016).
31. Moulin, L., Rahmouni, A. R. & Boccard, F. Topological insulators inhibit diffusion of transcription-induced positive supercoils in the chromosome of Escherichia coli. Molecular Microbiology 55, 601–610 (2005).
32. Samal, R. & Leng, F. Transcription-coupled Hypernegative Supercoiling of Plasmid DNA by T7 RNA Polymerase in Escherichia coli Topoisomerase I-Deficient Strains. J. Mol. Biol. 374, 925–935 (2007).
33. Rahmouni, A. R. & Wells, R. D. Direct Evidence for the Effect of Transcription on Local DNA Supercoiling In vivo. J. Mol. Biol. 223, 131–144 (1992).
34. Goldstein, E. & Drlica, K. Regulation of bacterial DNA supercoiling: plasmid linking numbers vary with growth temperature. Proc. Natl. Acad. Sci. USA 81, 4046–4050 (1984).
35. López-García, P. & Forterre, P. DNA topology and the thermal stress response, a tale from mesophiles and hyperthermophiles. BioEssays 22, 738–746 (2000).
36. Amit, R., Oppenheim, A. B. & Stavans, J. Increased Bending Rigidity of Single DNA Molecules by H-NS, a Temperature and Osmolarity Sensor. Biophys. J. 84, 2467–2473 (2003).
37. Jones, P. G., Van Bogelen, R. A. & Neidhardt, F. C. Induction of proteins in response to low temperature in Escherichia coli. J. Bacteriol. 169, 2092–2095 (1987).
38. Kapanidis, A. N. et al. Initial transcription by RNA polymerase proceeds through a DNA-scrunching mechanism. Science 314, 1144–1147 (2006).
39. Hsu, L. M. Promoter clearance and escape in prokaryotes. Biochim. Biophys. Acta - Gene Struct. Expr. 1577, 191–207 (2002).
40. Lloyd-Price, J. et al. Dissecting the stochastic transcription initiation process in live Escherichia coli. DNA Res. 23, 203–214 (2016).
41. Oliveira, S. M. D. et al. Increased cytoplasm viscosity hampers aggregate polar segregation in Escherichia coli. Mol. Microbiol. 99, 686–699 (2016).
42. Phadare, S. & Inouye, M. Genome-Wide Transcriptional Analysis of the Cold Shock Response in Wild-Type and Cold-Sensitive, Quadruple-csp-Deletion Strains of Escherichia coli. J. Bacteriol. 186(20), 7007–7014 (2004).
43. Kandavalli, V. K., Tran, H. & Ribeiro, A. S. Effects of σ factor competition are promoter initiation kinetics dependent. Biochim. Biophys. Acta (BBA)-Gene Regul. Mech. 1859, 1281–1288 (2016).
44. Oliveira, S. M. D. et al. Temperature-dependent model of multi-step transcription initiation in Escherichia coli based on live single-cell measurements. PLoS Comput. Biol. 12, e1005174 (2016).
45. S. Parry, R. R. et al. The bacterial cytoplasm has glass-like properties and is fluidized by metabolic activity. Cell 156, 183–194 (2014).
46. Lineweaver, H. & Burk, D. The determination of enzyme dissociation constants. J. Am. Chem. Soc. 56, 658–666 (1934).
47. Mäkelä, J., Kandavalli, V. & Ribeiro, A. S. Rate-limiting steps in transcription dictate sensitivity to variability in cellular components. Sci. Rep. 7, 10588 (2017).
48. Oliveira, S. M. D. et al. Single-cell kinetics of a repressilator when implemented in a single-copy plasmid. Mol. Biosyst. 11, 1939–1945 (2015).
49. Higgins, N. P. & Vologodskii, A. V. Topological Behaviour of PlasmidDNA. J. Mol. Biol. 73, 4474–4478 (1976).
50. Drlica, K. Control of bacterial DNA supercoiling. J. Mol. Microbiol. 6, 425–433 (1992).
51. Wang, J. C. DNA Topoisomerases. Annu. Rev. Biochem. 65, 635–692 (1996).
52. Ma, D. et al. Genes acrA and acrB encode a stress–induced efflux system of Escherichia coli. Mol. Microbiol. 16, 45–55 (1995).
53. Patel, K., Craig, S. B., Mcbride, M. G. & Palepu, N. R. Microbial inhibitory properties and stability of topotecan hydrochloride injection. Ann. J. Pharm. Sci. 55, 1584–1587 (1998).
54. Chen, A. Y. & Liu, L. F. DNA topoisomerases: Essential enzymes and lethal targets. Annu. Rev. Pharmacol. Toxicol. 34, 191–218 (1994).
55. Muthukrishnan, A. B., Martikainen, A., Neeli-Venkata, R. & Ribeiro, A. S. In vivo transcription kinetics of a synthetic gene uninvolved in stress-response pathways in stressed Escherichia coli cells. PLoS One 9, e109005 (2014).
56. Tran, H., Oliveira, S. M. D., Goncalves, N. & Ribeiro, A. S. Kinetics of the cellular intake of a gene expression inducer at high concentrations. Mol. Biosyst. 11, 2579–2587 (2015).
57. Taniguchi, Y. et al. Quantifying E. coli proteome and transcriptome with single-molecule sensitivity in single cells. Science 329, 533–538 (2010).
58. Bernstein, I. A., Khodursky, A. B., Lin, P.-H., Lin-Chao, S. & Cohen, S. N. Global analysis of mRNA decay and abundance in Escherichia coli at single-gene resolution using two-color fluorescent DNA microarrays. Proc. Natl. Acad. Sci. USA 99, 9697–9702 (2002).
59. El Hanafi, D. & Bossi, L. Activation and silencing of leu-500 promoter by transcription-induced DNA supercoiling in the Salmonella chromosome. Mol. Microbiol. 37, 583–594 (2000).
60. Oehler, S., Eisemann, E. R., Krämer, H. & Müller-Hill, B. Control of bacterial DNA supercoiling.
61. Lineweaver, H. & Burk, D. The determination of enzyme dissociation constants. J. Am. Chem. Soc. 56, 658–666 (1934).
62. Mäkelä, J., Kandavalli, V. & Ribeiro, A. S. Rate-limiting steps in transcription dictate sensitivity to variability in cellular components. Sci. Rep. 7, 10588 (2017).
63. Oliveira, S. M. D. et al. Single-cell kinetics of a repressilator when implemented in a single-copy plasmid. Mol. Biosyst. 11, 1939–1945 (2015).
64. Higgins, N. P. & Vologodskii, A. V. Topological Behaviour of PlasmidDNA.
82. Bratton, B. P., Mooney, R. A. & Weisshaar, J. C. Spatial distribution and diffusive motion of RNA polymerase in live. Escherichia coli. J. Bacteriol. 193, 5138–5146 (2011).
83. Santinha, J. et al. In Biomedical Image Analysis and Mining Techniques for Improved Health Outcomes. I, 71–99 (IGI Global, 2015).
84. Häkkinen, A., Muthukrishnan, A.-B., Mora, A., Fonseca, J. M. & Ribeiro, A. S. CellAging: a tool to study segregation and partitioning in division in cell lineages of Escherichia coli. Bioinformatics 29, 1708–1709 (2013).
85. Mäkelä, J., Lloyd-Price, J., Yli-Harja, O. & Ribeiro, A. S. Stochastic sequence-level model of coupled transcription and translation in prokaryotes. BMC Bioinformatics 12, 121 (2011).
86. Lloyd-Price, J., Gupta, A. & Ribeiro, A. S. SGNS2: A Compartmentalized Stochastic Chemical Kinetics Simulator for Dynamic Cell Populations. Bioinformatics 28, 3004–3005 (2012).
87. Gillespie, D. T. Exact Stochastic Simulation of Coupled Chemical Reactions. J. Phys. Chem. 81, 2340–2361 (1977).
88. Roussel, M. R. & Zhu, R. Validation of an algorithm for delay stochastic simulation of transcription and translation in prokaryotic gene expression. Phys. Biol. 3, 274–284 (2006).

Acknowledgements
We thank Sofia Startceva and Huy Tran for valuable advices on data analysis and models. Work supported by Portuguese Foundation for Science and Technology FCT/MCTES (SFRH/BD/88987/2012 to L.M.), Vilho, Yrjo and Kalle Vaisala Foundation (to S.M.D.O.), Tampere University of Technology President’s Graduate Programme (R.N.-V.), Pirkanmaa Regional Fund (to V.K.K.), FCT Strategic Program (UID/EEA/00066/203 to J.M.F.), Academy of Finland (295027 to A.S.R. and 305342 to A.S.R.), and Jane and Aatos Erkko Foundation (610536 to A.S.R.). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. The authors declare that they have no conflict of interest.

Author Contributions
A.S.R. conceived the study. S.M.D.O., N.S.M.G. and V.K.K. conceived and performed experiments. L.M. and S.M.D.O. performed image and data analysis. S.M.D.O. and R.N.-V. designed the genetic constructs. V.K.K. performed plasmid constructions. J.R. and H.K. performed genome integrations. J.M.F. assisted in the data and image analysis. A.S.R., L.M. and S.M.D.O. created the models. A.S.R. and S.M.D.O. wrote the manuscript, assisted by N.S.M.G., L.M., V.K.K. and J.L.-P. All authors performed research and contributed to the experimental design, discussion, and paper writing.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-39618-z.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019