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Horizontally acquired AT-rich genes in *Escherichia coli* cause toxicity by sequestering RNA polymerase

Lisa E. Lamberte¹, Gabriele Baniulytc^{1,†}, Shivani S. Singh¹, Anne M. Stringer², Richard P. Bonocora², Mathew Stracy³, Achillefs N. Kapanidis³, Joseph T. Wade^{2,4}, David C. Grainger^{1,*}

¹Institute of Microbiology and Infection, School of Biosciences, University of Birmingham, Edgbaston, Birmingham, B15 2TT, UK

²Wadsworth Center, New York State Department of Health, Albany, NY, 12208, USA

³Biological Physics Research Group, Clarendon Laboratory, Department of Physics, University of Oxford, Oxford OX1 3PU, UK

⁴Department of Biomedical Sciences, School of Public Health, University at Albany, SUNY, Albany, NY, 12201, USA

Abstract

Horizontal gene transfer permits rapid dissemination of genetic elements between individuals in bacterial populations. Transmitted DNA sequences may encode favourable traits. However, if acquired DNA has an atypical base composition, it can reduce host fitness. Consequently, bacteria have evolved strategies to minimise the harmful effects of foreign genes. Most notably, xenogeneic silencing proteins bind incoming DNA that has a higher AT-content than the host genome. An enduring question has been to understand why such sequences are deleterious. Here, we show that the toxicity of AT-rich DNA in *Escherichia coli* frequently results from constitutive transcription initiation within the coding regions of genes. Left unchecked, this causes titration of RNA polymerase and a global downshift in host gene expression. Accordingly, a mutation in RNA polymerase that diminishes the impact of AT-rich DNA on host fitness, reduces transcription from constitutive, but not activator-dependent, promoters.

Introduction

Bacteria obtain DNA from their environment by direct uptake (transformation), the action of viruses (transduction), or the acquisition of transmissible plasmids (conjugation)¹. Thus, “horizontal” DNA transfer allows phenotypes to spread through bacterial populations. Transferred traits can be beneficial¹. However, acquired sequences with a high AT-content

*for correspondence d.grainger@bham.ac.uk Tel: +44 (0)121 4145437.

†Present address: Department of Biomedical Sciences, School of Public Health, University at Albany, SUNY, Albany, NY, 12201, USA

Contributions

D.C.G. and J.T.W. designed the study and wrote the manuscript. L.E.L., G.B., S.S.S., A.M.S., R.P.B., and M.S. generated the data and prepared it for publication. M.S. and A.N.K. provided new analytical tools and critically discussed the manuscript with D.C.G. and J.T.W. All authors contributed to data analysis and interpretation.

may reduce host fitness²⁻⁶. Bacteria have developed mechanisms to diminish the toxicity of AT-rich genes. One approach involves a family of xenogeneic DNA binding proteins^{7,8}. These proteins target DNA that has a higher AT-content than the host genome (“AT-rich” DNA), and silence transcription⁹⁻¹². However, it is unclear why AT-rich DNA is toxic and why transcriptional silencing is beneficial.

The Histone-like Nucleoid Structuring (H-NS) protein of γ -proteobacteria is the best-characterised xenogeneic silencing protein. To reduce transcription, H-NS oligomerises across AT-rich genes. This process is triggered by nucleation at sequences containing a T:A step¹⁴⁻¹⁹. Intriguingly, promoter -10 hexamers (consensus 5'-TATAAT-3', recognised by the RNA polymerase σ^{70} subunit) are excellent inducers of H-NS nucleation^{15,20}. Hence, H-NS can exclude RNA polymerase from, or trap RNA polymerase at, promoter DNA^{8,9,21,22}.

A challenge has been to understand why horizontally acquired AT-rich DNA is toxic. As we noted previously, such regions contain a disproportionately high number of promoter-like sequences²³⁻²⁶. Furthermore, we showed that H-NS suppresses transcription primarily from promoters that are intragenic and/or far from gene starts²⁵. Here, we test the hypothesis that transcription from intragenic promoters is a major cause of toxicity in cells lacking H-NS. Working with a sub-set of AT-rich genes, we demonstrate dramatic effects of H-NS on intragenic transcription. When this transcription is disabled, associated fitness costs decrease. Genome-wide, if derepressed, intragenic transcription sequesters RNA polymerase. This causes a global downshift in canonical gene expression. Accordingly, a mutation in RNA polymerase that reduces transcription from constitutive promoters, but not those that are activator-dependent, compensates for loss of H-NS.

Results

Identification of the canonical *yccE* promoter

We chose *yccE* as a model gene to understand xenogeneic silencing. We selected *yccE* because it is AT-rich (66% A/T), silenced and bound by H-NS²⁷, and contains at least two intragenic promoters^{25,28}. Importantly, *yccE* can be studied in isolation because the adjacent genes are not H-NS bound (Figure 1a). We sought to understand the source of *yccE* transcription in cells lacking H-NS. As a starting point, we identified the canonical *yccE* promoter. We were aided by previous observations that *yccE* is a target for σ^{32} (heat shock σ factor)²⁹. Segments of the *yccE* intergenic region were fused to *lacZ* in reporter plasmid pRW50 (Figure S1a). Promoter activity located to a 55 base pair (bp) DNA fragment immediately upstream of *yccE* (Figure S1b). Figure 1b shows the sequence of this DNA fragment, named *yccE* 200. A σ^{32} -dependent promoter sequence is apparent. We monitored transcripts produced from this promoter (*PyccE*) *in vitro*. To do this, fragments from the *yccE* intergenic region were cloned upstream of the λ oop terminator in plasmid pSR. Transcripts initiating at *PyccE*, and terminating at λ oop, should be ~106 nucleotides (nt) in length. Coincidentally, the RNA-I transcript, derived from the pSR replication origin, is 108/107 nt long. To avoid confusing transcripts, we first monitored RNAs produced using pSR without a *PyccE* insert. The data show that σ^{32} dependent synthesis of the RNA-I transcript is inefficient (Figure 1c, compare lanes 1 and 2). When the entire *yccE* intergenic region was cloned in pSR σ^{32} dependent *yccE* transcripts were identified (Figure 1c, lane 3).

The same transcripts were observed upon cloning the truncated *yccEA200* fragment (Figure 1c, lane 4). A 10 bp truncation at the 5' end of *yccEA200* abolished transcription (Figure 1c, lane 5). To confirm correct identification of *PyccE*, DNA recognition elements for σ^{32} were mutated (detailed in Figure 1b). All mutations greatly reduced promoter activity (Figure 1d).

Increased transcription of *yccE* in *hns* cells does not require *PyccE*

Convention dictates that *PyccE* should cause increased *yccE* transcription in cells lacking H-NS. We considered this unlikely given the low activity, and σ^{32} dependence, of *PyccE*.

Hence, we generated a series of *yccE::lacZ* fusions to investigate the contribution of *PyccE*. The different constructs, labelled *i* through *iv*, are illustrated below the graph in Figure 1e.

We first measured *lacZ* expression in wildtype cells (Figure 1e, grey bars). When *yccE* was in the forward orientation, low-level *lacZ* expression was apparent. This was abolished upon deletion of *PyccE* (compare constructs *ii* and *iii*). In the reverse orientation, *yccE* stimulated higher *lacZ* expression (construct *iv*). This is consistent with our previous identification of a strong antisense promoter at the 5' end of *yccE* (adjacent to *lacZ* in this assay)²⁸.

Remarkably, in the absence of H-NS, *lacZ* expression increased in all scenarios (Figure 1e, white bars). This transcription must be due to intragenic promoters.

The *yccE* coding sequence is enriched for promoters

A search identified 21 possible promoters within *yccE*. To test for function, we isolated promoters on 56 bp DNA fragments and fused them to *lacZ* (Figure 2a). Note that such sequences are too short to be subject to repression by H-NS^{25,30}. Eleven DNA fragments stimulated β -galactosidase expression twofold or more above background (labelled A-K in Figure 2a). Under the conditions of our experiment, 7 of the 11 intragenic promoters were more efficient at driving transcription than *PyccE* (Figure 2a, black bar). Note that fragment "A" contains the strong antisense promoter described previously²⁸.

Transcription initiation within *yccE* is repressed by H-NS *in vitro*

In the context of the full *yccE* gene, intragenic promoters should be repressed by H-NS. We tested this *in vitro* using the pSR system. As noted above, with empty pSR, the 108/107 nt RNA-I transcript is produced. In addition, larger transcripts are generated from genes native to the plasmid. The complete set of transcripts produced from pSR is shown in Lane 1 of Figure 2b. In the figure, large (>1000 nt) pSR derived transcripts are highlighted by a black box and RNA-I is highlighted by a grey box. We introduced *yccE* into pSR, in either the forward or reverse orientation, upstream of the *loop* terminator. Since *yccE* is 1257 bp in length, most intragenic transcripts should be separable from pSR derived transcripts. As expected, numerous transcripts between 100 and 1000 nt in length were detected for both *yccE* containing plasmids (compare Lanes 1, 2 and 6 in Figure 2b). For the plasmid containing *yccE* in the reverse orientation an additional small transcript was detected (highlighted by lower blue box in Figure 2b). This was expected, the transcript generated from the strong antisense promoter at the 5' end of *yccE*, has a size of 90 nt in this assay. Regardless of *yccE* orientation, addition of H-NS inhibited synthesis of most *yccE* derived transcripts (Lanes 3-5 and 6-9). Synthesis of the RNA-I transcript was enhanced by H-NS suggesting that RNA polymerase is titrated by promoters within *yccE*.

Intragenic promoters are the source of increased *yccE* transcription in cells lacking H-NS

To confirm that H-NS repressed intragenic *yccE* promoters *in vivo*, we made derivatives of our *yccE::lacZ* fusions. The derivatives carry mutations in intragenic -10 elements. Schematics are below the graph in Figure 2c. The open arrows show *yccE* lacking internal promoters. Full details are in Figure S2. In the absence of H-NS, the ability of mutated *yccE* alleles to stimulate *lacZ* expression was reduced (Figure 2c). Thus, both biochemical and genetic inspection show silencing of intragenic transcription by H-NS. We refer to this as “pseudo-regulation” that occurs independently of, and may be mistaken for, the control of mRNA synthesis. Thus, supposed gene regulatory effects of H-NS may often be due to intragenic promoters.

The fitness cost of *yccE* is a consequence of intragenic transcription

We predicted a link between intragenic transcription and reduced fitness associated with loss of H-NS. Hence, *E. coli* M182, and the *hns::kan* variant, were transformed with pSR carrying *yccE* with or without internal promoters. Importantly, *yccE* mRNA cannot be expressed in this scenario; no upstream promoter is present. We monitored cultures inoculated with these strains (Figure 2d,e). In all cases, cells lacking *hns* had reduced fitness compared to the parent. However, this fitness defect was smaller for the *yccE* derivative lacking internal promoters (Figure 2e). Complete elimination of the fitness defect was not expected; AT-rich genes present on the *E. coli* chromosome also contribute.

Repression of intragenic transcription by H-NS reduces the fitness cost of many AT-rich genes

We identified other solitary genes targeted by H-NS: *yfdF*, *ykgH*, *yjgN* and *yjgL*. We cloned these, and derivatives lacking intragenic promoters, upstream of the λ oop terminator in pSR. The *fepE* gene, which has an AT-content of 55%, was included as a control. The constructs are illustrated above the gel image in Figure 3a. Full gene sequences are in Figure S3. Figure 3a shows results of *in vitro* transcription experiments. Whilst RNA polymerase did not initiate transcription within *fepE* (Lane 1), intragenic transcription was observed for *yfdF*, *ykgH*, *yjgN* and *yjgL* (Lanes 2, 4, 6 and 8). Mutation of intragenic promoters reduced this transcription (Lanes 3, 5, 7 and 9). The coding regions described above were also cloned upstream of *lacZ* in pRW50 (Figure 3b). Expression of *lacZ* was measured in M182 or the *hns::kan* derivative (Figure 3b). Upon deletion of *hns*, expression of *lacZ* downstream of *yfdF*, *ykgH*, *yjgN* or *yjgL*, but not *fepE*, increased (solid arrows). In contrast, no such increase occurred when intragenic promoters were mutated (open arrows). These analyses are consistent with silencing of intragenic promoters by H-NS at all loci tested. Hence, we measured the fitness cost of multicopy *yfdF*, *ykgH*, *yjgN* and *yjgL* in cells with and without H-NS. In all cases, the fitness deficit between wild-type and *hns::kan* cells decreased upon mutation of intragenic promoters (Figure 3c,d). Furthermore, changes in fitness and *lacZ* expression were significantly correlated (Figure S4a).

AT-rich genes titrate RNA polymerase and cause a global downshift in housekeeping transcription

RNA polymerase levels are limited in *E. coli*³¹. Consequently, multiple copies of a strong promoter can hinder growth and titrate transcription of the *lac* operon (Figure S4b). Interestingly, RNA polymerase levels do not increase in cells lacking H-NS (Figure S5). Therefore, competition for the enzyme must increase; more promoters compete for a limited supply of RNA polymerase. Logically, migration of RNA polymerase to spurious promoters should cause a global downshift in canonical transcription. However, despite many studies of the H-NS controlled transcriptome, a universal downshift has never been reported. We reasoned that this might result from data normalisation approaches used previously. Briefly, transcriptome analysis compares RNA levels in two strains. Comparison requires a point of reference believed to be consistent between the strains. For example, it may be assumed that housekeeping genes are similarly transcribed or that averaged transcription across all genes will be equivalent. Problematically, these approaches cannot differentiate between technical variation and genuine shifts in global transcription. To circumvent this problem, spiked-in RNA standards can be used³⁴. We adapted this tactic to quantify global effects of H-NS on transcription. Briefly, we grew *E. coli* MG1655, and the *hns::kan* variant, to mid-log phase. We then counted colony-forming units for each *E. coli* culture. The same was done for a single culture of *Salmonella* Typhimurium. Accordingly, we were able to mix a defined number of clonal *S. Typhimurium* cells with each *E. coli* strain. The cell mixtures were subject to transcriptome analysis. For identically grown *S. Typhimurium* cells, differences in RNA abundance must result from processing variation. Hence, at the end of the procedure, transcripts mapping uniquely to *S. Typhimurium* were used to normalise the data. Figure 4a shows a post-normalisation plot of read depth for each *E. coli* gene in each strain. The diagonal blue line shows the expected position of data points if transcription is unchanged between strains. There is a downshift in transcription of genes not bound by H-NS (black). Conversely, H-NS bound genes (red) are transcribed more frequently. This behaviour is exemplified in Figure 4b.

RNA polymerase titration can be visualised directly

Genes bound by H-NS are overrepresented near the chromosome replication terminus (Ter)^{35,36}. Conversely, most RNA polymerase binds near the origin of replication (Ori)³⁷. These loci occupy distinct intracellular territories; Ter typically frequents mid-cell whilst Ori migrates to the poles³⁷⁻³⁹. Consequently, it should be possible to visualise titration of RNA polymerase at the cellular level. Previously, we used super resolution microscopy to track individual RNA polymerase molecules in live *E. coli*⁴⁰. Here, we repeated this analysis to examine the effect of deleting *hns*. In wild-type cells, RNA polymerase clusters near the quarter cell positions (Figure 4c, top). However, in cells lacking H-NS, RNA polymerase is redistributed and mid-cell is occupied (Figure 4c, middle). Consistent with unaltered RNA polymerase abundance, occupancy of mid-cell corresponds to reduced RNA polymerase abundance elsewhere (Figure 4c).

RNA polymerase mutation can compensate for loss of H-NS by favouring regulated transcription

An aspartic acid substitution in the RNA polymerase σ^{70} subunit (G424D in *E. coli*) can compensate for loss of H-NS⁴¹. This side chain could clash with the promoter -10 element during transcription initiation⁴¹. According to our model, promoters within genes are constitutive; they rely solely on their DNA sequence to bind RNA polymerase. Conversely, canonical promoters are regulated; a complex array of transcriptional activators stabilise RNA polymerase binding and DNA unwinding⁴². We reasoned that transcriptional defects, due to the G424D mutation, might be more pronounced at constitutive promoters. To test this, we purified RNA polymerase containing σ^{70} or the G424D derivative. We then investigated the ability of the RNA polymerase derivatives to stimulate unwinding of a promoter -10 element using KMnO₄ footprinting. The semi-synthetic NM501 promoter was used because it has constitutive activity, by virtue of a consensus -10 element, but can be upregulated by the transcriptional activator CRP⁴³. The G424D mutation completely abolished DNA opening by RNA polymerase in the absence of CRP (Figure 4d, compare lanes 2-5 and 6-9). In contrast, when CRP was present, the σ^{70} G424D mutation had little effect (compare lanes 10-13 and 14-17). Similarly, *in vivo*, the σ^{70} G424D mutation caused transcriptional defects at 7 of 8 constitutive promoters (Figure 4e). However, σ^{70} G424D functioned as well as, or better than, wild type RNA polymerase at activator dependent promoters (Figure 4f).

Discussion

The AT-rich genes examined here impose a fitness cost due to intragenic promoters. This phenomenon is likely to be widespread in bacteria; functional homologues of H-NS are apparent in diverse species⁸. Furthermore, the DNA binding properties of RNA polymerase are highly conserved⁴². We suggest that misappropriation of cellular resources underlies the *hns* phenotype. Redeployment of the finite RNA polymerase pool causes uniform suppression of canonical transcription. Whilst this general effect is likely to be pervasive, we do not exclude organism-specific complications. For example, in *Pseudomonas aeruginosa*, loss of H-NS-like proteins causes prophage induction and cell death^{44,45}. Interestingly, linear H-NS filaments do not pose a barrier to transit of RNA polymerase⁴⁶. Accordingly, transcription of an mRNA, and silencing of intragenic promoters, could occur simultaneously. For example, in this work, *PyccE* was active both in isolation and upstream of H-NS bound *yccE*. We speculate that gene silencing by H-NS may have evolved to discriminate between canonical and spurious RNA synthesis.

Materials and Methods

Strains, plasmids and general methods

E. coli JCB387 *nir lac* and MG1655 have been described previously^{47,48}. M182*hns::kan* and KF26*hns::kan* were constructed by P1 transduction of *hns::kan* from MG1655 into the respective parent strains. The MG1655 *hns::kan* strain was provided by Ding Jin. M182*rpoD::kan* was generated using gene doctoring according to the protocol of Lee *et al*⁴⁹ using the plasmids and oligonucleotides listed in Table S1. Note that, prior to gene

doctoring, M182 strains were transformed with plasmid pVR σ that encodes *rpoD*⁵⁰. Quickchange mutagenesis was used to introduce the G424D mutation into pVR σ encoded *rpoD* (Table S1). Fortuitously, chromosomal and plasmid encoded σ ⁷⁰ were produced at indistinguishable levels (Figure S6). Sample sizes for all experiments were selected to ensure reproducibility in line with our previous work.

DNA fragments and gene expression assays

Promoter::*lacZ* fusions were made by cloning DNA fragments upstream of *lacZ* in the low copy number plasmid pRW50⁵¹. The nested deletions in the *yccE* intergenic region were generated by PCR and oligonucleotides shown in Table S1. The various *yccE*, *yfdF*, *ykgH*, *yjgN* and *yjgL* alleles were synthesised by Invitrogen and some contain silent mutations to remove *EcoRI* or *HindIII* restriction sites to facilitate cloning (Figures S2a and S3). Oligonucleotides used to amplify the different alleles for cloning into pSR and pRW50 are shown in Table S1. The 56 bp intragenic *yccE* fragments were generated with overlapping oligonucleotides (Table S1). The resulting single stranded overhangs were filled with DNA polymerase before cloning. β -galactosidase assays were done using the protocol of Miller⁵². All assay values are the mean of three biological replicates and the error bars show standard deviation from the mean. Experiments were done at least twice. Cells were grown aerobically at 37°C, to mid-log phase, in LB media.

Bioinformatic analysis of genes and design of new coding regions

Our stringent search criteria selected putative σ ⁷⁰ dependent promoters as described previously²⁵. Thus, sequences were selected that matched the motifs 5'-TAnAAT-3', 5'-TATnAT-3' or 5'-TATAnT-3'. The relaxed search selected the sequence 5'-TAnnnT-3'. To inactivate promoter -10 elements the initial 5'-TA-3' was replaced with 5'-GG-3'

Proteins, KMnO₄ footprinting and in vitro transcription assays

H-NS and RNA polymerase were prepared as described previously⁵³. DNA fragments for KMnO₄ footprinting experiments were derived from Qiagen maxi-preparations of plasmid pSR. Thus, promoter DNA fragments were excised from pSR by sequential digestion with *HindIII* and then *AaII*. After digestion fragments were labelled at the *HindIII* end using [γ -³²P]-ATP and polynucleotide kinase. Footprints were done as described by Grainger *et al.*⁵⁴. The *in vitro* transcription experiments were done as described by Savery *et al.*⁵⁵ using the system of Kolb *et al.*⁵⁶. A Qiagen maxiprep kit was used to purify supercoiled pSR plasmid carrying the different promoter inserts. This template (16 pg ml⁻¹) was pre-incubated with purified H-NS in buffer containing 20 mM Tris pH 7.9, 5 mM MgCl₂, 500 μ M DTT, 50 mM KCl, 100 μ g ml⁻¹ BSA, 200 μ M ATP, 200 μ M GTP, 200 μ M CTP, 10 μ M UTP with 5 μ Ci [α -³²P]-UTP. The reaction was started by adding purified *E. coli* RNA polymerase. Labelled RNA products were analysed on a denaturing polyacrylamide gel. All *in vitro* assays were repeated at least three times in their entirety.

Growth assays

Cells lacking H-NS rapidly acquire compensatory mutations⁴¹. Consequently, reproducible changes in growth were only obtained when precautions were taken to minimise this

phenomenon. The primary precaution was to reduce the number of division cycles that strains passed through during experimental setup. Thus, M182 and the *hns::kan* derivative were taken directly from long-term -80°C storage and used immediately to inoculate LB medium. After incubation for several hours at 37°C cells were harvested and competency was induced using ice cold CaCl₂. The cells were then transformed with desired plasmids and transformants were isolated on selective agar plates. A colony from each plate was suspended in LB medium and aliquots of this were used immediately to inoculate fresh media so that growth could be monitored. Cells were grown either in LB medium at 37°C or in M9 minimal medium at 30°C. Values shown are from three biological replicates and the experiments were done on two separate occasions.

Western blotting

To determine relative protein levels in different strains cells were grown in LB media at 37°C. Aliquots of the culture were harvested at indicated time points and CFUs determined. Following this quantification the same number of cells from each aliquot were resuspended in SDS-PAGE gel loading buffer. After heating to 90°C for two minutes we separated proteins present in each lysate using SDS-PAGE. The proteins were then transferred to a Hybond-ECL nitrocellulose membrane using an Invitrogen XCell II Blot Module. Mouse anti-sera against the σ^{70} and β subunits of RNA polymerase (Neoclone) and H-NS (a gift from Jay Hinton) was used to detect the relevant proteins. Primary antibody binding was probed with horseradish peroxidase-linked rabbit anti-mouse antisera (Sigma-Aldrich A9044). The experiments were done on two separate occasions.

Standard RNA-seq

RNA-seq experiments were done in duplicate. *E. coli* MG1655 and *E. coli* MG1655 *hns* were grown in LB medium at 30°C to an OD₆₀₀ of 0.5-0.7. RNA was harvested, RNA-seq libraries were constructed and libraries were sequenced as described previously⁵⁷.

RNA-seq with a spiked in control

Transcriptome analysis experiments shown were performed in duplicate using *E. coli* MG1655 and *E. coli* MG1655 *hns*. We also ran the same experiment, again in duplicate, using *E. coli* M182 and the *hns::kan* derivative. The results were near identical. For each replicate three mid-log phase bacterial cultures were prepared. The three cultures were *E. coli* MG1655, *E. coli* MG1655 *hns* or *Salmonella* Typhimurium 14028s in LB medium. Before harvesting RNA the number of colony forming units (CFUs) per unit volume was determined for each culture by plating dilutions of each culture on nutrient agar and counting bacterial colonies. Aliquots of the two *E. coli* cultures were mixed *S. Typhimurium* cells. The volume of *Salmonella* cells was normalised to the CFUs for the corresponding *E. coli* culture. This step is crucial because it allows processing artefacts, due to differences in lysis efficiency, RNA recovery or cDNA synthesis, to be removed. Thus, in the final transcriptome analysis, the number of sequencing reads corresponding to the *S. Typhimurium* genome should be identical for each sample; they were derived from the same number of clonal *S. Typhimurium* cells. Hence, differences can only result from downstream processing. RNA was harvested, and RNA-seq libraries were prepared, as described

previously⁵⁷. RNA-seq libraries were sequenced on an Illumina Hi-Seq 2000 instrument (University at Buffalo, SUNY, Buffalo, NY, USA).

Data normalisation and transcriptome analysis

Our normalisation procedure is based on the addition of a proportional number of *S. Typhimurium* cells to each sample of *E. coli* cells immediately before harvesting RNA. All reads were first mapped to the *E. coli* MG1655 genome using CLC Genomics Workbench (Version 8.0; default parameters except that a perfect match was required). Unmapped reads were mapped to the *S. Typhimurium* 14028s genome (same parameters as above). The number of mapped *S. Typhimurium* reads, for each *E. coli* sample, was used to determine a correction factor for each sample. For example, if Sample A has twice as many *S. Typhimurium* reads as Sample B, the correction factor for Sample A will be twice that for Sample B. Having calculated a correction factor for each sample, we remapped all sequence reads to the *S. Typhimurium* 14028s genome using CLC Genomics Workbench (same parameters as above). Unmapped reads were then mapped to the *E. coli* MG1655 genome using CLC Genomics Workbench (same parameters as above). Total read coverage per gene was calculated using a custom Python script. These values were normalised to the length of the gene, and further normalised using the correction factor (described above). Raw data are available from the ArrayExpress database using accession number E-MTAB-4751.

Super resolution microscopy of RNA polymerase

Cell preparation, microscopy, and analysis were performed as described previously^{40,58}. In brief, we used an endogenous fusion of photoactivatable fluorescent protein PAMCherry, with the β' subunit of RNA polymerase, encoded by *E. coli* strain KF26⁵⁸. Glycerol stocks of KF26, and the *hns::kan* derivative, were used to inoculate fresh Rich Defined Media (RDM, Teknova). When the culture attained an OD₆₅₀ value of ~0.2 cells were collected by centrifugation and resuspended. Cells in 1 μ l of the suspension were placed on an RDM agarose pad and imaged for 300000 frames at 15 ms exposure on a custom built single-molecule TIRF microscope. For each strain, 9 fields of view were imaged. The experiment was done on 3 separate occasions. Molecules were imaged by photoactivating and localising fluorophores, and joining localisations over multiple frames to obtain trajectories of individual molecules. To measure RNA polymerase mobility, we calculated an apparent diffusion coefficient from the mean squared displacement of trajectories of individual molecules, and used a threshold to distinguish transcriptionally active and promoter bound molecules from the rest of the population. Cell outlines were determined from the brightfield image, and the average intracellular location of RNA polymerase was established from a 2D histogram of the mean trajectory positions relative to the cell outline for all DNA-bound molecules in at least 100 cells. Ter positioning was determined using similar analysis of strain PZ111 carrying a *tetO* array inserted at Ter, and expressing a TetR-mYpet fusion. The strain was constructed by P1 transduction of *tetR*-mYPet kan, into an AB1157 strain with an array of 240 *tetO* sequences 50 kb clockwise of *dif*(ter3)⁵⁹.

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Data availability

The data that support these findings are available from the corresponding author upon request. The raw data for RNA-seq experiments are available from the ArrayExpress database using accession number E-MTAB-4751. Original gel images are shown in Figure S7.

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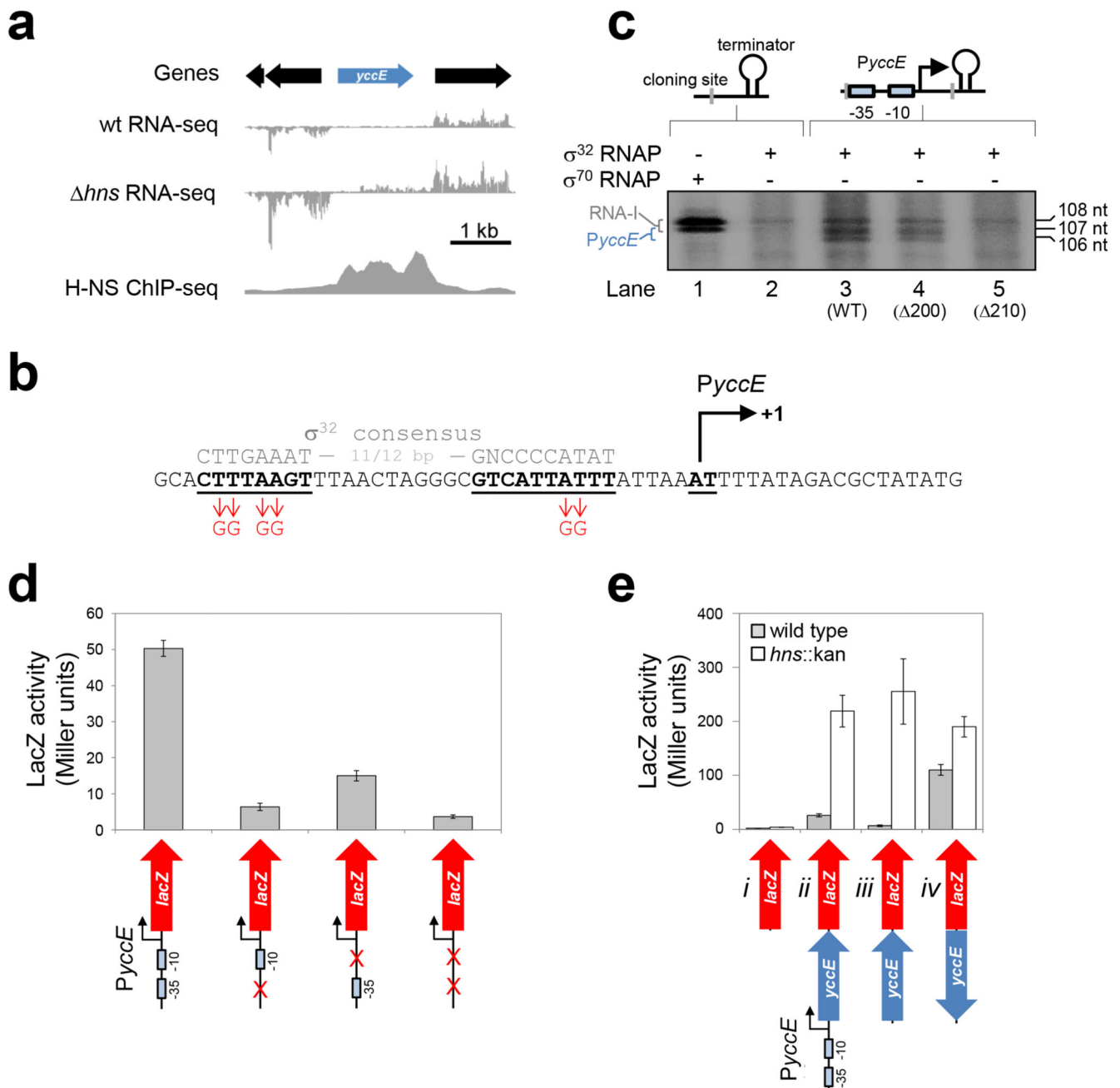


Figure 1. Characterisation of the *yccE* locus.

a) Genomic context of *yccE* and its promoter. The panel shows *yccE*, and surrounding genes, alongside data describing H-NS binding (ChIP-seq²⁷) and RNA abundance (standard RNA-seq; this work, done in duplicate). Data are representative.

b) Sequence of the *yccE* 200 DNA fragment containing *PyccE*. The *PyccE*-10 and -35 elements are in bold and underlined. A consensus σ^{32} promoter sequence is in grey for comparison. Mutations made to disrupt sequence elements are in red. Transcription can initiate at adjacent nucleotides. These are labelled (+1) and are highlighted by a bent arrow.

c) Analysis of transcripts generated from *Pycce* *in vitro*. The gel image shows transcripts generated by RNA polymerase, associated with either σ^{70} or σ^{32} , from pSR plasmid DNA templates. The schematic diagrams above the gel image represent the native cloning site of pSR (left hand side) or derivatives containing a *Pycce* insert (right hand side). The different *Pycce* containing DNA fragments inserted are indicated below the gel image in parenthesis. *Pycce* derived transcripts manifest as a 107/106 nucleotide (nt) doublet. The 108/107 nt RNA-I transcript is derived from the plasmid replication origin. The experiment was done three times. Data are representative.

d) Effect of *Pycce* mutation *in vivo*. The graph shows LacZ activity data obtained from *E. coli* JCB387 cells carrying different *yccE* 200 derivatives cloned in pRW50. Assays were done in triplicate and error bars show standard deviation from the mean.

e) Induction of *yccE* transcription in the absence of H-NS does not require *Pycce* *in vivo*. The panel illustrates a series of *yccE::lacZ* fusions labelled *i-iv*. Genes are shown as block arrows and *Pycce* is shown as a bent line arrow. The β -galactosidase activity was measured in lysates of M182 (grey bars) and M182*hns::kan* cells (open bars). Assays were done in triplicate and error bars show standard deviation from the mean.

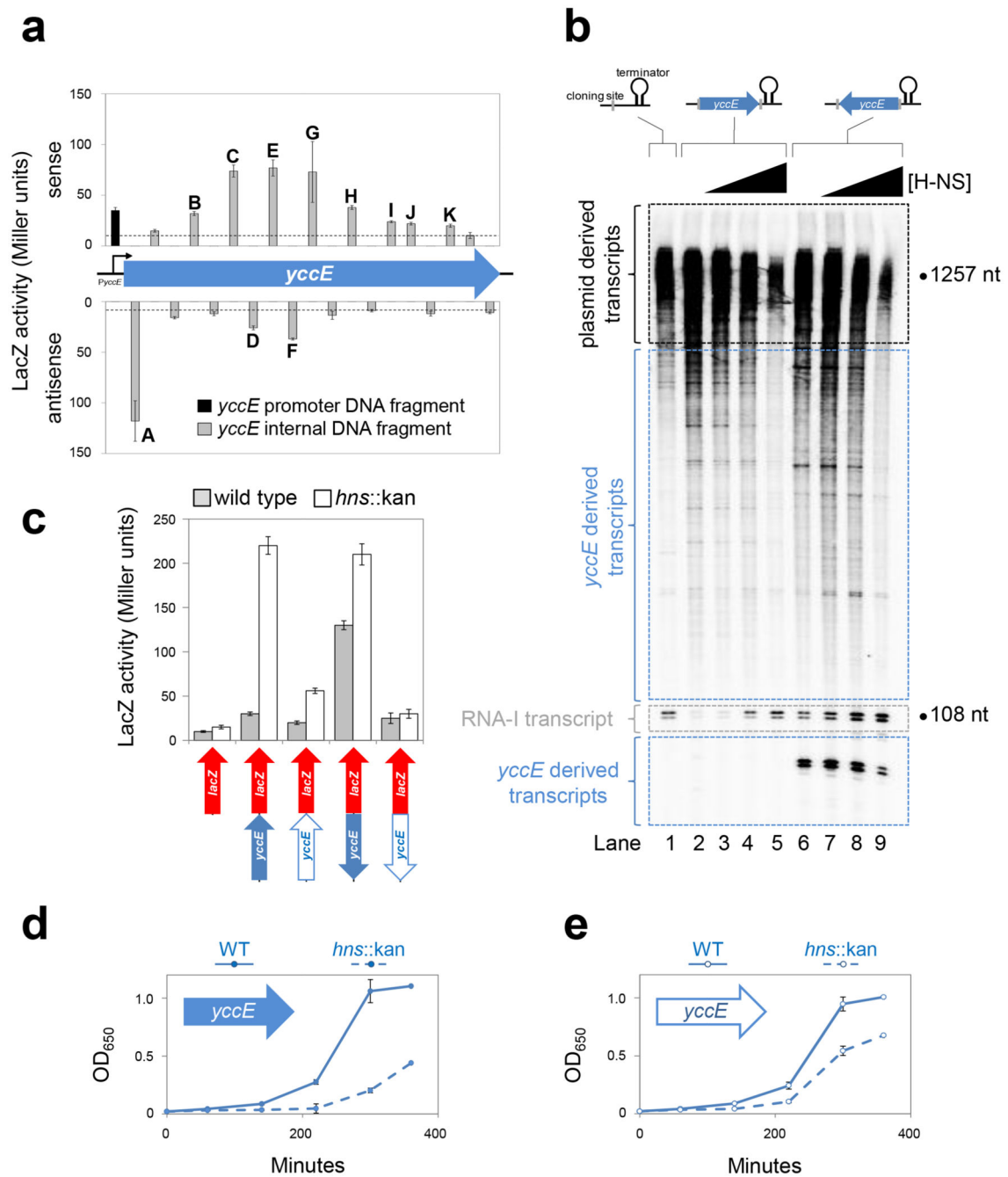


Figure 2. H-NS represses intragenic *yccE* transcription and associated fitness costs

a) Identification of intragenic *yccE* promoters. The data are β -galactosidase activities driven by short intragenic *yccE* DNA fragments in strain JCB387. The bars align with the location of the DNA fragment relative to *yccE* and show sense (upper) and antisense (lower) transcription. Bars labelled “a”-“k” have at least 2-fold over background activity (empty pRW50; shown by dashed line). The black bar represents the canonical promoter *PyccE*. Note that, two DNA fragments resisted cloning. Hence, 19 of the 21 potential promoters

were tested. Assays were done in triplicate. Error bars show standard deviation from the mean.

b) Transcription can initiate at multiple sites within *yccE* *in vitro*. Gel image showing RNA generated *in vitro* separated by denaturing gel electrophoresis. DNA templates, with the *yccE* gene cloned upstream of the λ oop terminator in plasmid pSR, are illustrated above the gel. Transcripts generated by RNA polymerase (400 nM) with empty pSR plasmid (lane 1) are highlighted by a black dashed line. Transcripts initiating within *yccE* in the forward (lanes 2-5) or reverse (lanes 6-9) orientation are highlighted by a blue dashed line. The control RNA-I transcripts are highlighted by a grey dashed line. H-NS was added at concentrations of 0.8, 1.5, or 3.0 μ M. The experiment, done three times, is representative.

c) Mutation of promoters within *yccE* prevents induction in *hns* cells. The lower illustrations show *yccE* (blue arrows) cloned upstream of *lacZ* (red arrow). A solid blue arrow represents wild type *yccE* whereas open arrows indicate *yccE* with mutated intragenic promoter -10 elements. For each *lacZ* fusion, β -galactosidase activity was measured in lysates of M182 (grey bars) and M182*hns*:kan cells (open bars). Assays were done in triplicate and error bars show standard deviation from the mean.

d,e) The fitness cost of *yccE* is reduced when intragenic promoters are mutated. The figure illustrates changes in culture OD650 following inoculation of LB medium. The inoculum was M182 (solid line) or M182*hns*:kan (dashed line) transformed with the pSR plasmid carrying d) wild type *yccE* or e) *yccE* with internal promoter -10 elements mutated. Cells were grown at 37°C. The experiment was done in triplicate. Error bars show standard deviation from the mean.

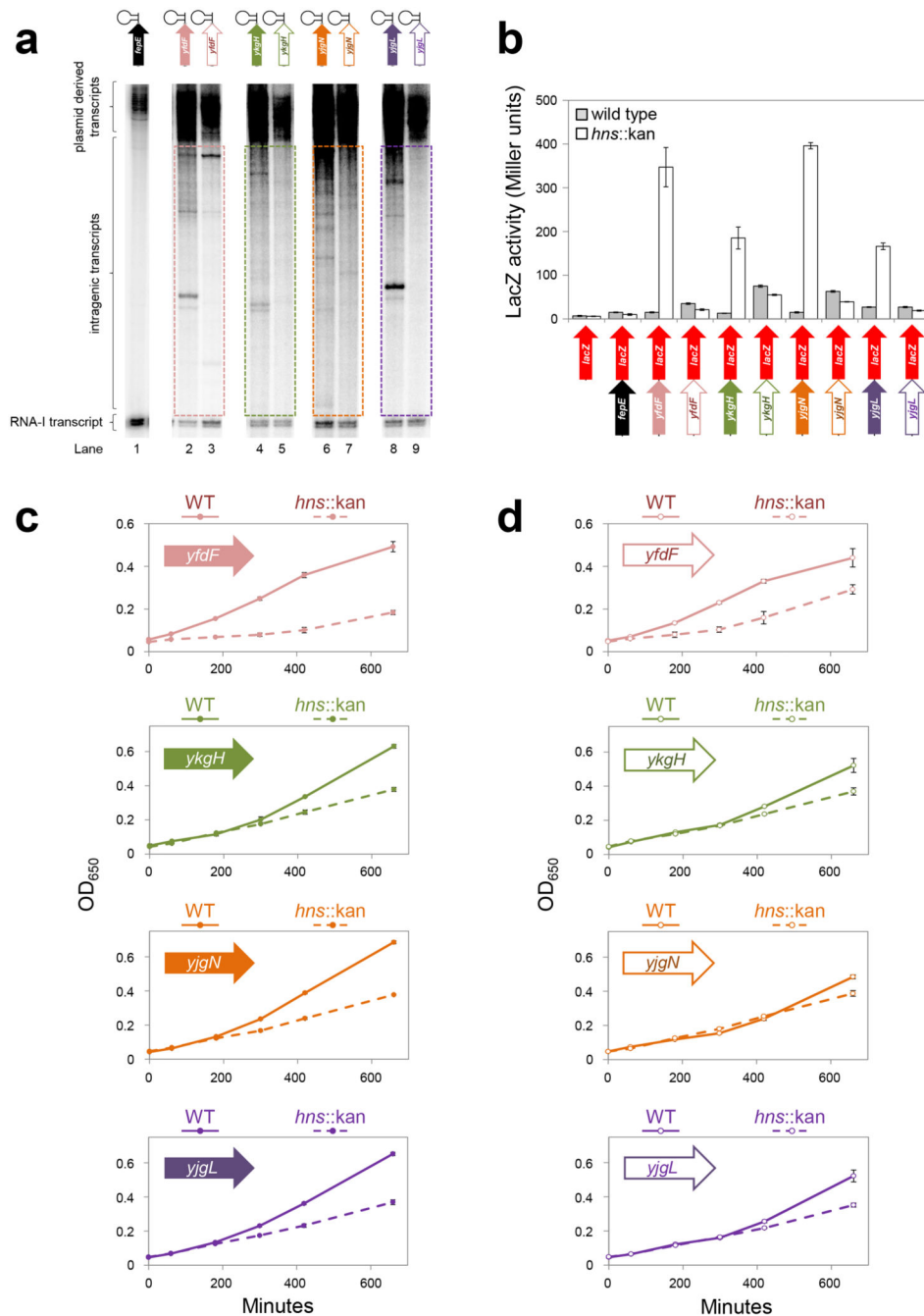


Figure 3. H-NS represses intragenic transcription and associated fitness costs at many loci.
a) Transcription initiation within the coding regions of H-NS target genes *in vitro*. An *in vitro* transcription assay using different AT-rich genes, cloned upstream of the *loop* terminator in plasmid pSR, as a template. The different DNA constructs are illustrated above the gel. The cloned genes have an AT-content of 65% (*yfdF*), 63% (*ykgH*), 63% (*yjgN*) and 68% (*yjgL*). For each cloned gene a solid arrow represents the wild type DNA sequence whereas an open arrow is a derivative where intragenic promoter -10 elements have point mutations. Note that the *fepA* gene was used as a control and has an AT-content of 55%. The

positions of transcripts generated by RNA polymerase (400 nM) from the *fepE* control (lane 1) or the other cloned genes (lanes 2-9) are labelled. The *in vitro* transcription assays were run on three separate occasions. Data are representative.

b) Increased transcription in cells lacking H-NS frequently requires intragenic promoters *in vivo*. The panel illustrates a series of DNA constructs where different gene coding regions have been cloned upstream of *lacZ* (red arrow). For each cloned gene a solid arrow represents the wild type DNA sequence whereas an open arrow is a derivative where intragenic promoter -10 elements have point mutations. The cloned genes have an AT-content of 65% (*yfdF*), 63% (*ykgH*), 63% (*yjgN*) and 68% (*yjgL*). Note that the *fepA* gene is used as a control and has an AT-content of 55%. For each *lacZ* fusion β -galactosidase activity was measured in lysates of M182 (grey bars) and M182 *hns* (open bars) cells. Assays were done in triplicate and error bars show standard deviation from the mean.

c,d) The toxicity of many AT-rich genes is a consequence of spurious intragenic transcription. The figure illustrates growth of M182 (solid line) or M182*hns::kan* (dashed line) cells transformed with the pSR plasmid carrying different AT-rich genes. Panel c) shows wild type gene derivatives and

d) shows derivatives with internal promoter -10 elements mutated (open arrows). Experiments were done using M9 minimal media at 30°C. The experiment was done in triplicate and error bars show standard deviation from the mean.

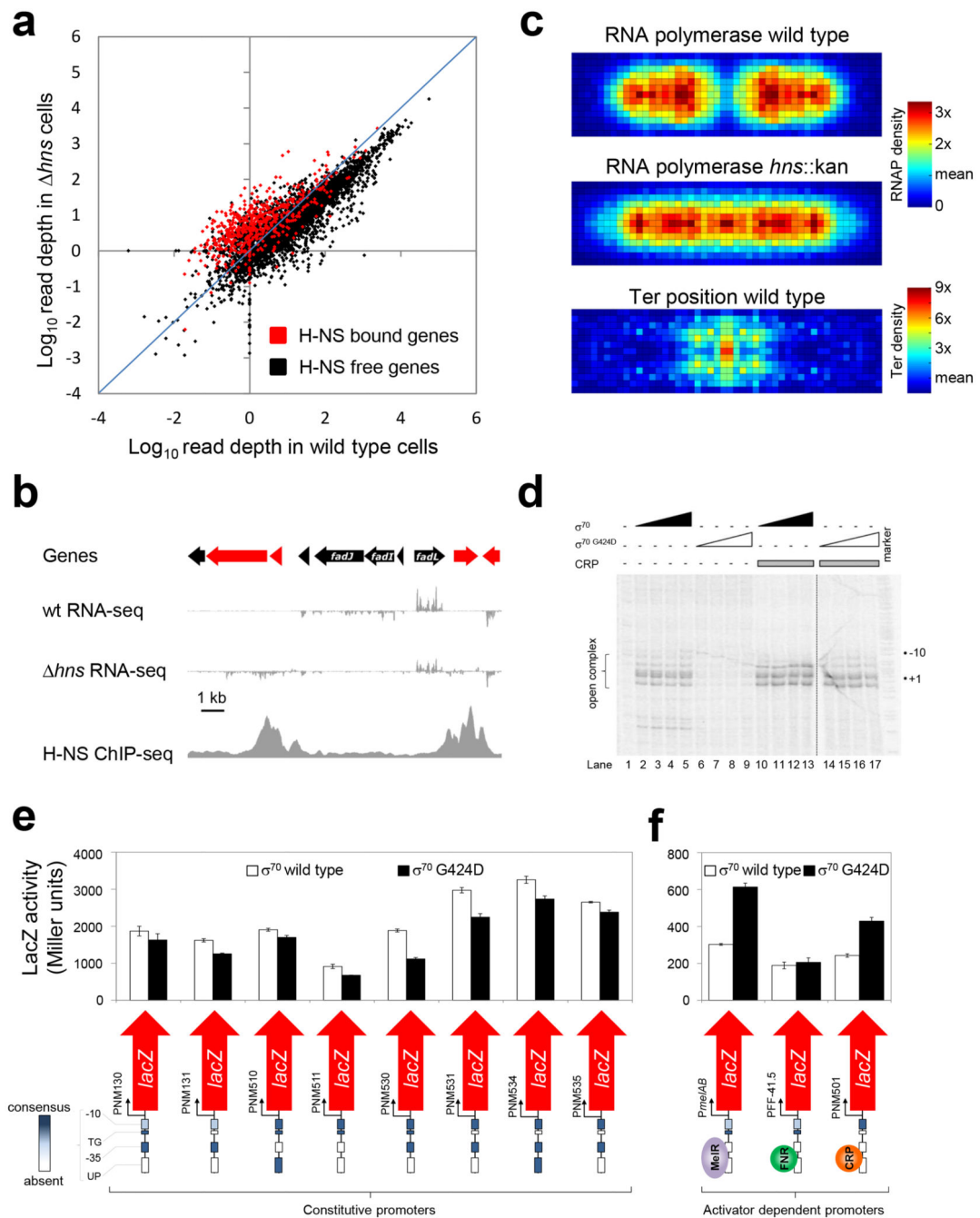


Figure 4. Most transcription is uniformly downregulated in cells lacking H-NS.

a,b) Most transcription is uniformly downregulated in cells lacking H-NS. a) the plot illustrates changes in global transcription caused by loss of H-NS. Data points represent H-NS bound (red) and unbound (black) genes. Genes with unaltered transcription should fall on the diagonal blue line. Data are from duplicate RNA-seq experiments. b) The basally expressed *fad* genes whose transcription is reduced in the absence of H-NS. Genes bound by H-NS are in red and other genes are black. Graphs show H-NS binding²⁷ and RNA abundance (RNA-seq with spiked in control; this work). Data are representative.

c) RNA polymerase is redistributed in cells lacking H-NS. RNA polymerase distribution in wild type (top) and *hns::kan* cells (middle). Each heat map shows the average position of DNA-bound (i.e. transcribing or interacting with a promoter) RNA polymerase molecules within the cell. The bottom panel shows the average position of Ter as determined by visualising a TetR-mYpet fusion bound at an array of Ter proximal *tetO* sequences. Each distribution was generated from 100 cells between 3.5 and 4.5 μm in length. Each square is 1/624 of total cell area.

d) The σ^{70} G424D mutation hinders constitutive but not activator dependent NM501 promoter activity *in vitro*. KMnO₄ footprinting reactions analysed on a denaturing polyacrylamide gel. Bands are indicative of open complex near the transcription start site (+1). RNA polymerase added at concentrations of 200, 250, 300 or 350 nM and CRP was 1.0 μM . The experiment, done three times, is representative.

e,f) The σ^{70} G424D mutation hinders constitutive but not activator dependent promoter activity *in vivo*. Different promoter DNA fragments were cloned upstream of *lacZ* (red arrow). For each promoter the location of key DNA sequence elements is represented by a box. In each case, the box is coloured according to the relationship between the DNA sequence and the consensus sequence for that element: perfect (dark blue), imperfect (pale blue) or completely absent (white). For each *lacZ* fusion β -galactosidase activity was measured in lysates of JCB387 *rpoD::kan* carrying pVR σ (white bars) or pVR σ^{G424D} (black bars). The experiment was done in triplicate. Error bars show standard deviation from the mean.