UNIVERSITY^{OF} BIRMINGHAM

University of Birmingham Research at Birmingham

Xenogeneic silencing strategies in bacteria are dictated by RNA polymerase promiscuity

Forrest, David; Warman, Emily A; Erkelens, Amanda M; Dame, Remus T; Grainger, David C

DOI:

10.1038/s41467-022-28747-1

License

Creative Commons: Attribution (CC BY)

Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Forrest, D, Warman, EA, Erkelens, AM, Dame, RT & Grainger, DC 2022, 'Xenogeneic silencing strategies in bacteria are dictated by RNA polymerase promiscuity', *Nature Communications*, vol. 13, no. 1, 1149. https://doi.org/10.1038/s41467-022-28747-1

Link to publication on Research at Birmingham portal

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

•Users may freely distribute the URL that is used to identify this publication.

•Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.

•User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)

•Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

Download date: 19. Apr. 2024



ARTICLE



1

https://doi.org/10.1038/s41467-022-28747-1

OPFN

Xenogeneic silencing strategies in bacteria are dictated by RNA polymerase promiscuity

Horizontal gene transfer facilitates dissemination of favourable traits among bacteria. However, foreign DNA can also reduce host fitness: incoming sequences with a higher AT content than the host genome can misdirect transcription. Xenogeneic silencing proteins counteract this by modulating RNA polymerase binding. In this work, we compare xenogeneic silencing strategies of two distantly related model organisms: *Escherichia coli* and *Bacillus subtilis*. In *E. coli*, silencing is mediated by the H-NS protein that binds extensively across horizontally acquired genes. This prevents spurious non-coding transcription, mostly intragenic in origin. By contrast, binding of the *B. subtilis* Rok protein is more targeted and mostly silences expression of functional mRNAs. The difference reflects contrasting transcriptional promiscuity in *E. coli* and *B. subtilis*, largely attributable to housekeeping RNA polymerase σ factors. Thus, whilst RNA polymerase specificity is key to the xenogeneic silencing strategy of *B. subtilis*, transcriptional promiscuity must be overcome to silence horizontally acquired DNA in *E. coli*.

¹ School of Biosciences, University of Birmingham, Edgbaston B15 2TT, UK. ² Leiden Institute of Chemistry, Leiden University, Einsteinweg 55, 2333CC Leiden, The Netherlands. ³ Centre for Microbial Cell Biology, Leiden University, Einsteinweg 55, 2333CC Leiden, The Netherlands. [™]email: d.grainger@bham.ac.uk

ccurate transcription initiation by RNA polymerase depends on DNA sequences called promoters¹. In bacteria, RNA polymerase exists in two states. The core enzyme (β , β' and two α subunits) is catalytically active but cannot bind DNA specifically². Conversely, the holoenzyme contains a dissociable σ factor that delivers RNA polymerase to promoters³. Small accessory subunits, which vary between species, can be components of either core or holoenzyme but are not essential⁴. Housekeeping σ factors, found in all bacterial species, are essential and direct most transcription. Named σ^{70} in Escherichia coli, and σ^{A} in many other bacteria, housekeeping σ factors share four domains: σ_1 , σ_2 , σ_3 and σ_4^5 . The σ_2 , σ_3 and σ_4 domains mediate promoter binding and DNA unwinding during transcription initiation^{3,5}. Particularly important is the interaction between σ_2 and the promoter -10 element (consensus 5'-TATAAT-3'). This drives DNA opening that is the key prerequisite for transcription to begin⁶. Hence, amino acids in σ_2 that mediate DNA melting are conserved in diverse housekeeping σ factors⁵. Nevertheless, the kinetics of open complex formation vary between species and the reasons for this are not fully understood^{7,8}. Once RNA polymerase holoenzyme has escaped the promoter, and entered the elongation phase of transcription, the σ factor dissociates⁹. Core enzyme then completes transcript synthesis before releasing the DNA template and nascent RNA. Reassociation with σ again directs RNA polymerase to a promoter and gene transcription begins afresh.

In addition to canonical genes and their promoters, most bacterial genomes contain large sections of horizontally acquired DNA¹⁰. These include remnants of prophages, pathogenicity islands, and conjugative elements 11,12. Many such regions are conspicuous because their AT-content is higher than the surrounding genome (i.e. they are AT-rich)¹⁰. Thus, whilst encoded proteins can be beneficial, such loci can negatively impact fitness¹³. Working with E. coli, we recently showed that such defects can result from uncontrolled transcription^{14,15}. In particular, sequences resembling promoter -10 elements occur frequently in AT-rich DNA, often within genes, and are used by σ^{70} bound RNA polymerase to initiate transcription 14,16. In turn, this sequesters RNA polymerase and canonical gene expression is reduced globally¹⁵. To avoid this, E. coli masks AT-rich DNA using Histone-like Nucleoid Structuring (H-NS) protein¹⁷. Present at 20,000 molecules per cell, H-NS forms filaments with stretches of AT-rich DNA to prevent promoter recognition and transcription elongation 10,15,18,19. This process is known as xenogeneic silencing²⁰. Genes encoding H-NS homologues are widely distributed in γ-proteobacteria²¹. Amongst the best characterised are the MvaT and MvaU proteins of Pseudomonas aeruginosa²²⁻²⁴. These factors also act to block spurious transcription initiation within horizontally acquired DNA²⁵. Whilst the precise structural details differ, these silencers all recognise AT-rich sequences using arginine side chains that penetrate the DNA minor groove^{23,26}.

Beyond the γ-proteobacteria, most prokaryotes do not encode H-NS homologues. However, unrelated functional analogues have been identified. For example, in actinobacteria, Lsr2 uses a mode of DNA recognition resembling H-NS to silence foreign genes and non-coding antisense transcription^{26,27}. In Gram-positive bacteria, the Rok protein of *B. subtilis* is best understood^{28,29}. First identified as a repressor of natural competence³⁰, Rok was subsequently observed to bind parts of the *B. subtilis* genome that are AT-rich and foreign in origin²⁸. Like H-NS, Rok can oligomerise and bind AT-rich DNA using its N- and C-terminal domains respectively²⁹. However, nothing is known about mechanisms of silencing by Rok or the type of transcript initiation events targeted. It is noteworthy that Rok, present at ~1500 molecules per cell, is much less abundant than H-NS in *E. coli*²⁸.

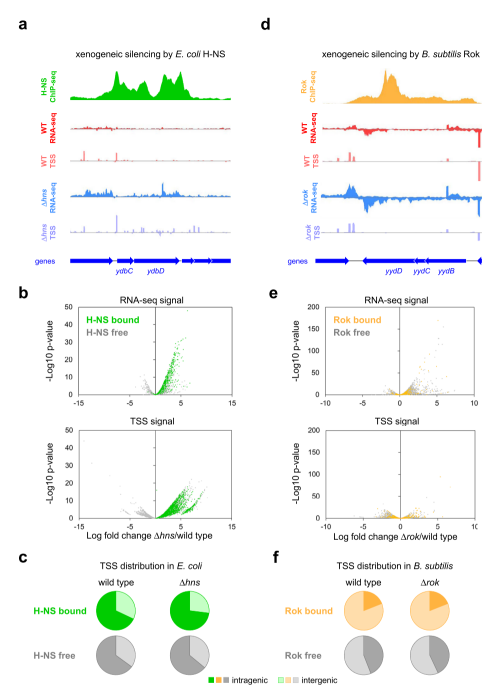
Conversely, the AT-content of the *B. subtilis* genome, and horizontally acquired sequences within, is higher^{11,12}. Hence, the need to prevent spurious transcription initiation, within foreign *B. subtilis* genes, would seem acute.

In this work, we have compared the xenogeneic silencing strategies of H-NS and Rok in E. coli and B. subtilis respectively. We show that, in sharp contrast to H-NS, Rok does not repress spurious intragenic transcription initiation. Instead, Rok specifically targets mRNA synthesis from canonical promoters. The different approaches reflect housekeeping RNA polymerase behaviour in each organism; the E. coli enzyme is more promiscuous, largely by virtue of the σ^{70} factor. Thus, the expression of σ^{70} in B. subtilis results in spurious intragenic transcription that Rok is unable to prevent. Conversely, B. subtilis σ^{A} imposes greater specificity on E. coli RNA polymerase. Genome-wide, the two σ factors direct RNA polymerase to slightly different promoter sequences and E. coli σ^{70} is less fastidious. We conclude that RNA polymerase specificity is key to the xenogeneic silencing strategy in B. subtilis. Conversely, transcriptional promiscuity is a barrier that must be overcome to silence horizontally acquired genes in E. coli.

Results

High-resolution mapping of the H-NS regulated transcriptome in Escherichia coli. Previously, we used poly(5'-phosphatase)sequencing to map E. coli transcription start sites (TSSs) genomewide in the presence and absence of H-NS¹⁴. Cappable-seq offers numerous advantages over our prior approach³¹. Most notably, cappable-seq is more sensitive and specifically identifies the triphosphorylated 5' ends of primary transcripts³¹. Hence, we began by remapping H-NS regulated TSSs in E. coli using cappable-seq. Our detection of transcription initiation events increased threefold (Supplementary Data 1). Parallel RNA-seq assays quantified transcript abundance (Supplementary Data 2) and H-NS ChIPseq profiles³² completed a high-resolution view of the regulon. A representative chromosomal region, which typifies the transcriptional response to H-NS, is shown in Fig. 1a. In wild-type cells, we observed low levels of transcription (red track) and few TSSs (pink track) across regions bound by H-NS (green track). By contrast, in the absence of H-NS, transcript abundance increased (blue track) and many TSSs were detected (mauve track). Consistent with our prior work, most of these H-NS repressed TSSs were intragenic^{14,15}. The results are summarised as volcano plots in Fig. 1b. The top panel shows transcript abundance determined by RNA-seq and each data point represents an individual gene. Transcription across H-NS bound genes (green) is derepressed when H-NS is absent. In the lower panel, each data point corresponds to a TSS identified using cappable-seq; a similar trend is evident. Figure 1c illustrates the number of all E. coli TSSs locating to intragenic or intergenic DNA. As expected, for H-NS bound regions, more intragenic TSSs are found when H-NS is absent.

High-resolution mapping of the Rok regulated transcriptome in *Bacillus subtilis*. Like H-NS analogues in other organisms, the *B. subtilis* Rok protein recognises horizontally acquired AT-rich DNA³³. In particular, Rok preferentially binds DNA sequences with TpA steps, rather than continuous A or T tracts^{29,33}. Such sequences are ideal templates for transcription initiation events^{14,15,34}. We next sought to understand how Rok might intervene and applied the approaches described above. The cappable-seq and RNA-seq results are described in Tables S1 and S2 respectively and were compared to existing ChIP-seq data for Rok binding³⁵. A region of the *B. subtilis* genome with a typical transcriptional response to Rok is shown in Fig. 1d. As expected, in the presence of Rok, we observed low levels of transcription (red track) and infrequent TSSs (pink track) across the Rok



bound DNA (orange track). In the absence of Rok, levels of transcription increased (blue track) but additional TSSs were not detected (mauve track). The complete data are visualised as volcano plots in Fig. 1e. Levels of transcription across most Rok bound genes increased in the absence of Rok (top panel). Conversely, Rok had little influence on cappable-seq signals at TSSs (bottom panel). Genome-wide, TSSs were less likely to be intragenic in Rok bound regions and deletion of Rok had no impact on this distribution (Fig. 1f). Hence, despite the high sensitivity of cappable-seq, we did not detect widespread intragenic transcription initiation in the absence of Rok in *B. subtilis*. This differs markedly to the situation in *E. coli* for H-NS (Fig. 1a-c)^{14,15}.

H-NS and Rok have different patterns of DNA binding in vivo. Whilst examining the ChIP-seq data we noticed differences in the distribution of H-NS and Rok (Figs. S1a and S1b). Most notably,

H-NS binding appeared to invade coding sequences more frequently. We quantified this apparent difference in two ways. First, we generated aggregate plots of H-NS and Rok binding, relative to gene start codons, for the complete ChIP-seq datasets. The results are shown in Supplementary Fig. 1c. For Rok, DNA binding is usually centred near gene start codons and tends not to encroach substantially on coding sequences (left panel). Conversely, for H-NS, binding peaks are much broader and encompass large tracts of coding DNA (right panel). Second, we examined the relationship between the binding signal intensity in ChIP-seq experiments and the occurrence of non-coding DNA. The results are shown in Supplementary Fig. 1d. For Rok, increasing the binding signal positively correlates with the occurrence of non-coding DNA. Hence, nearly all of the highest signals for Rok binding located to non-coding sequences (orange line). The pattern for H-NS is more complicated. The likelihood of DNA being non-coding is elevated until 40% of the maximum H-NS

Fig. 1 Xenogeneic silencing impacts transcription differently in E. coli and B. subtilis. a Repression of transcription by H-NS in E. coli. The genomic region encompassing ydbCD is shown. Data for H-NS occupancy are shown by the green graph³². The total RNA abundance determined by RNA-seq in wild-type and Δhns cells is shown by the red and blue graphs respectively. Transcription start sites (TSSs) were identified by cappable-seq for wild-type (pink graph) and Δhns (mauve graph) cells. In the cappable-seq data only RNA 5' ends are sequenced and so the upstream edge of each peak indicates a TSS. Sequence reads mapping to the top and bottom DNA strands are shown above and below the central horizontal line in each plot. The v-axis scales are identical for data obtained using wild-type and Δhns cells for each type of experiment. Genes are shown by blue arrows. **b** Volcano plots illustrating differences in the distribution of signals obtained by RNA-seq (top panel) or cappable-seq (bottom panel) in the presence and absence of H-NS in E. coli. For the RNA-seq analysis, each data point represents the average signal across an individual gene. In the cappable-seq data plot, each data point represents a separate TSS. For both plots, data points are coloured to indicate DNA regions bound by (green) or free from (grey) H-NS. c The pie charts illustrate the distribution of TSSs obtained by cappable-seq from wild-type (left hand side) and Δhns (right hand side) E. coli cells. The TSSs are further separated into those in H-NS bound (green) and H-NS free (grey) regions. For all pie charts, dark shading indicates TSSs in coding DNA whilst pale shading identifies TSSs in non-coding regions. d Repression of transcription by Rok in B. subtilis. The genomic region encompassing yydBCD is shown. Data for Rok occupancy are shown by the orange graph³⁵. Colour coding is otherwise as shown in (a) except that here the comparison is between wild-type and Δrok B. subtilis cells. e Volcano plots illustrating differences signals obtained by RNA-seq (top panel) or cappable-seq (bottom panel) in the presence and absence of Rok in B. subtilis. Data points are as described for (b) and coloured to indicate DNA regions bound by (orange) or free from (grey) Rok. f Pie charts illustrate the distribution of TSSs obtained by cappable-seq from wild-type (left hand side) and Δhns (right hand side) B. subtilis cells. The TSSs are further separated into those in Rok bound (orange) and Rok free (grey) regions. For all pie charts, dark shading indicates TSSs in coding DNA whilst pale shading identifies TSSs in non-coding regions.

binding signal is reached. Hence, regions of maximal H-NS binding are often within coding DNA.

Bacillus subtilis and Escherichia coli RNA polymerase have different propensities for spurious intragenic transcription in vitro. We reasoned that a B. subtilis factor other than Rok might prevent transcription initiation, within AT-rich genes, in vivo. If correct, the B. subtilis RNA polymerase holoenzyme should generate spurious intragenic transcripts from naked DNA templates in vitro. The genomic region that we selected as an in vitro template is shown in Fig. 2a. The DNA segment includes comK, a well-characterised Rok target gene³⁰, and the upstream regulatory DNA. As expected, the comK gene is subject to direct repression by Rok in vivo (compare red and blue tracks in Fig. 2a). However, consistent with the observations described above, we did not detect intragenic transcription initiation in the absence of Rok (compare pink and mauve tracks in Fig. 2a). To measure transcription from this DNA locus in vitro we cloned comK, and the associated regulatory DNA, upstream of the λoop terminator in plasmid pSR (see schematic in Fig. 2b). In this context, RNA polymerase is expected to generate a 696 nucleotide (nt) long mRNA from the comK promoter. Spurious RNAs of intragenic origin, terminated by the λoop signal, should be smaller. The result of the in vitro transcription assay is shown below the Fig. 2b schematic. As expected, a 696 nt transcript was detected (lane 1) and subject to repression by Rok (lanes 2 and 3). However, we did not detect smaller transcripts initiating from sites within the comK gene. Transcripts >1000 nt in size result from transcription units elsewhere in the pSR plasmid. The RNAI transcript is encoded at the plasmid replication origin. The higher RNAI signal upon Rok addition likely results from more RNA polymerase being available when transcription elsewhere, including RNAs encoded by the plasmid backbone, is reduced. Our observation that B. subtilis RNA polymerase holoenzyme did not generate transcripts from sites within the comK coding sequence was surprising. Hence, in a control experiment, we reexamined the propensity of E. coli RNA polymerase holoenzyme to drive spurious transcription. Importantly, we selected an H-NS targeted gene from E. coli that had an AT-content of 56%, identical to that of comK. Consistent with our expectations, H-NS suppressed transcription from sites within the agaB coding sequence in vivo (Fig. 2c). A similar observation was made in vitro (Fig. 2d). Taken together, our data suggest that fundamentally different strategies are needed to ensure specific transcription of horizontally acquired DNA in B. subtilis and E. coli.

In particular, the *E. coli* RNA polymerase holoenzyme is intrinsically promiscuous. Hence, H-NS must block both intragenic transcription and mRNA production. Conversely, *B. subtilis* relies on enhanced RNA polymerase specificity to avoid transcription within genes. We next wanted to understand the basis for these differences.

The Bacillus subtilis RNA polymerase uses a narrower range of promoter configurations and is more sensitive to discriminator **sequence in vivo.** Our attention returned to our cappable-seq data. For each TSS detected we identified the associated promoter -10 element. The histograms in Fig. 3a show the distribution of distances separating TSSs from −10 elements. Consistent with many prior reports, the preferred distance was 7 bp for both B. subtilis and E. coli³⁶⁻³⁸. However, the overall distribution of distances was subtly different. In particular, for $E.\ coli$, more -10elements were separated from TSSs by 8 or 6 bp. Next, we aligned all DNA sequences upstream of TSSs for each organism. The alignments were anchored by TSS position and are illustrated as DNA sequence logos in Fig. 3b. There are three notable differences between the logos derived for B. subtilis and E. coli. First, because -10 elements are less consistently positioned, the 5'-TATAAT-3' consensus sequence is misrepresented for E. coli. Second, there is a clear preference for an AT-rich discriminator sequence in B. subtilis. Finally, the overall information content of the E. coli sequence logo is lower.

The Bacillus subtilis RNA polymerase is sensitive to discriminator sequence and length in vitro. Whilst informative, the DNA sequence logos in Fig. 3b represent aggregate properties of many promoters. We wanted to understand the impact of identified sequence features at a specific promoter. The veg promoter has been widely used as a model to study B. subtilis transcription³⁹. Hence, we made derivatives of the *veg* promoter with different discriminator sequence (AT- or GC-rich) and length (Fig. 3c). A 155 nt transcript is generated from these promoters in our in vitro transcription system. Figure 3d shows a representative gel image from such experiments and a quantification from three independent assays. Transcription using the B. subtilis oA holoenzyme is 2.5-fold lower if the AT-rich discriminator is replaced with a GC-rich sequence. The activity was reduced twofold further if the length of the GC-rich discriminator was increased by 1 base pair. Conversely, E. coli σ⁷⁰ RNA polymerase had indistinguishable activity at the different promoter

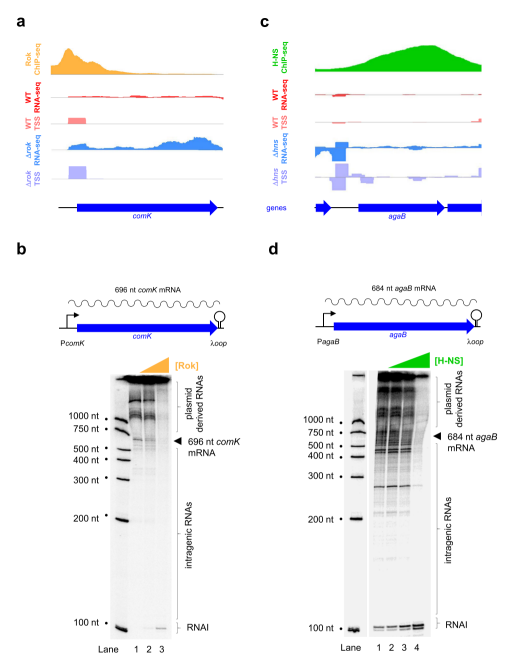


Fig. 2 Housekeeping RNA polymerases of E. coli and B. subtilis differ in their promiscuity. a Rok represses transcription of the B. subtilis comK mRNA in vivo. Data for Rok occupancy (orange)35, total RNA abundance (red and blue) and transcription start site (TSS) usage (pink and mauve) are shown. Sequence reads mapping to the top and bottom DNA strands are shown above and below the central horizontal line in each plot. The y-axis scales are identical for data obtained using wild-type and Δrok cells for each type of experiment. Genes are shown by blue arrows. **b** Rok represses transcription of the B. subtilis comK mRNA in vivo. The schematic illustrates the comK gene and regulatory region, cloned in plasmid pSR, and used as a template for in vitro transcription. The comK TSS is shown as a bent black arrow, the comK gene is shown as a block blue arrow, and the sequence encoding the λοορ transcriptional terminator is indicated by a stem loop schematic. B. subtilis σ^A RNA polymerase (0.5 μ M) and Rok (0, 0.5, or 1.0 μ M) were added as indicated. Note that the 696 nt comK mRNA is easily discernible and there is no evidence for transcription initiation within comK. Species of RNA over ~1000 nt in length are derived from sites elsewhere on the plasmid template. The RNAI transcript is encoded by the plasmid replication origin. The experiment was done twice with similar results. c H-NS represses transcription initiation within the E. coli agaB coding sequence in vivo. Data for H-NS occupancy are in green³² and otherwise as indicated in (a) except that wild-type and Δhns E. coli cells are compared. d H-NS represses transcription initiation within the E. coli agaB coding sequence in vitro. The schematic illustrates a section of DNA cloned in plasmid pSR and used as a template for in vitro transcription. The expected size of the agaB mRNA is 684 nucleotides (nt). The gel image shows transcripts generated by $E.~coli~\sigma^{70}$ RNA polymerase (0.5 μM) using this DNA template. The expected position of agaB mRNA is indicated by an arrow head but is obscured by many similarly sized and smaller transcripts derived from agaB coding sequence. H-NS was added at concentrations of 0, 0.5, 1.0 or 2.0 µM. The experiment was done twice with similar results.

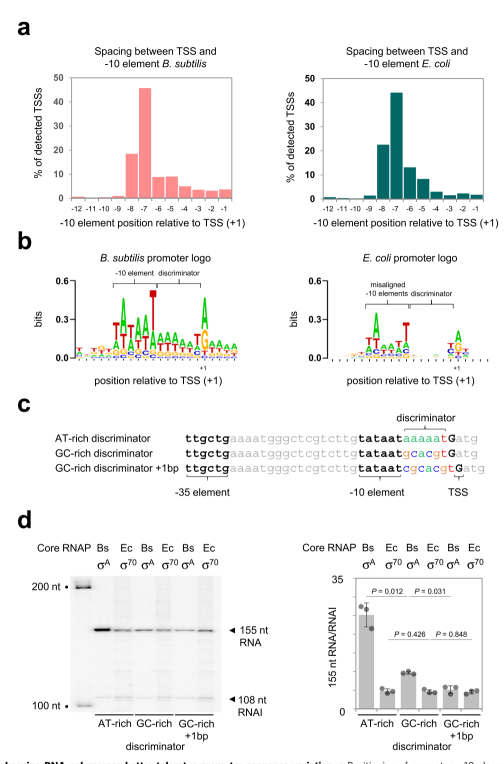


Fig. 3 *E. coli* housekeeping RNA polymerase better tolerates promoter sequence variation. **a** Positioning of promoter -10 elements and transcription start sites in *E. coli* and *B. subtilis*. The bar charts show the percentage of promoter -10 elements located at indicated distances upstream of transcription start sites (TSSs, +1) identified by cappable-seq for *E. coli* and *B. subtilis*. **b** The panel shows DNA sequence logos generated by aligning nucleic acid regions upstream of *B. subtilis* (left) or *E. coli* (right) transcription start sites. The more variable spacing between transcription start sites and promoter -10 elements in *E. coli* generates a motif that misrepresents the consensus -10 element sequence (5′-TATAAT-3′). There is no overall sequence preference for the promoter discriminator region in *E. coli* whilst an AT-rich sequence is common in *B. subtilis*. **c** DNA sequences of the *B. subtilis veg* promoter and derivatives with either an AT-rich or GC-rich discriminator sequence. **d** Results of in vitro transcription assays using DNA templates containing one of the promoter sequences shown in (**c**). Experiments were done with either the *B. subtilis* (Bs) σ ^A, or *E. coli* (Ec) σ ⁷⁰, RNA polymerase holoenzyme (0.5 μM). The 155 nucleotide (nt) transcript is generated from the cloned promoter and the 108 nt RNAl transcript is derived from the plasmid replication origin. The gel image shows a representative result and the bar chart shows a quantification of three independent experiments. Error bars show S.D. and the centre of the error bars is the mean. *P* was calculated using a two-tailed student's *T*-test.

sequences. Supplementary Fig. 2 shows results of KMnO₄ footprints that detect DNA melting. Open complexes formed by the *B. subtilis* enzyme are notoriously unstable and could not be detected using our assay^{7,40,41}. Unwinding dependent on *E. coli* σ^{70} was detected and less stable with the GC-rich discriminator.

Escherichia coli σ^{70} directs intragenic transcription of Rok bound genes by Bacillus subtilis core RNA polymerase in vivo. Our observations are consistent with the housekeeping B. subtilis RNA polymerase using a narrower range of promoter configurations than the equivalent E. coli enzyme. This may explain the different behaviour of each RNA polymerase at horizontally acquired genes. RNA polymerase-promoter interactions are primarily mediated by the exchangeable σ subunit³. Indeed, prior studies have shown that foreign σ factors, expressed in a given bacterium, can drive the host RNA polymerase to different promoters $^{42-44}$. Hence, we reasoned that E. coli σ^{70} might render B. subtilis core RNA polymerase more promiscuous. To test this prediction, the σ^{70} encoding rpoD gene was cloned in plasmid pDR111. The resulting DNA construct was used to transform B. subtilis strain 168ca. Following transformation, pDR111 integrates with the B. subtilis chromosome at the amyE locus. Hence, transformants encode a single copy of rpoD under the control of an IPTG inducible promoter. We used cappable-seq to identify TSSs in transformed cells. We compared these TSSs to those detected in the absence of σ^{70} . In total, the two cappable-seq analyses identified 8524 TSSs (Fig. 4a, Supplementary Data 1). The vast majority of these are likely to be dependent on either σ^{70} or σ^A rather than an alternative σ factor. Consistent with this, 82% of the TSSs were correctly positioned downstream of a housekeeping promoter -10 element. Of all TSSs, 2510 were identified only in the presence of σ^{70} . These likely represent otherwise silent promoters that can be used only when σ^{70} is expressed. There were 4593 TSSs identified both in the presence and absence of σ^{70} . The associated promoters could be used by both σ^{70} and σ^{A} . Alternatively, some of these promoters could use σ^{A} specifically, even if σ^{70} is present. Finally, we identified 1421 TSSs only in the absence of σ^{70} . Such TSSs are likely to be σ^A specific but not used when σ^{70} is expressed. For instance, σ^{70} could direct the limited pool of core RNA polymerase to alternative locations. Recall that our analysis above-identified subtle differences between E. coli and B. subtilis promoter sequences (Fig. 3a, b). Consistent with this, DNA sequence logos generated from E. coli σ^{70} dependent promoters found in B. subtilis had misaligned -10 elements and less AT-rich discriminators (Supplementary Fig. 3a). Conversely, canonical B. subtilis promoters did not exhibit these sequence features (Supplementary Figs. 3b, c). Most importantly, TSSs dependent on σ^{70} were more abundant in AT-rich islands targeted by Rok (compare left and right charts in Fig. 4b). Some examples are shown in Fig. 4c. In parallel experiments, Rok was similarly unable to silence intragenic transcription in E. coli cells lacking H-NS (Supplementary Fig. 4)

Escherichia coli σ^{70} directs intragenic transcription of Rok bound genes by Bacillus subtilis core RNA polymerase in vitro. We expected that swapping σ^{70} and σ^{A} , in the context of the E. coli and B. subtilis RNA polymerase holoenzymes, would alter transcriptional promiscuity in vitro. We tested this using in vitro transcription assays. The DNA templates were six different B. subtilis genes that we had identified as being targeted by Rok. None of these genes were subject to promiscuous transcription by B. subtilis RNA polymerase in cells lacking Rok. Lanes 1–6 show transcripts generated by B. subtilis core RNA polymerase in conjunction with σ^{A} (Fig. 4d). Note that yydD is located within an operon and so there is no regulatory upstream DNA. For all of

the other DNA templates the expected mRNA species were detected (marked by coloured triangles). Conversely, many intragenic transcripts were generated when the *B. subtilis* core enzyme was provided with σ^{70} (Lanes 7–12). Equivalent experiments using the *E. coli* core RNA polymerase are shown in Lanes 13–24. Again, promiscuity was dictated by the provided σ factor

Side chains R156 and R486 contribute to the promiscuity of E. coli σ^{70} and are absent in B. subtilis σ^{A} . Both σ^{70} and σ^{A} comprise four highly conserved domains (named σ_1 , σ_2 , σ_3 and σ_4) (Fig. 5a). The *E. coli* σ^{70} factor contains a further determinant, known as the non-conserved region (NCR), not present in σ^A . The aligned amino acid sequences of the σ factors are shown in Supplementary Fig. 5. Strikingly, amino acids responsible for interaction with core RNA polymerase and core promoter elements, at each step of transcription initiation, are near identical (Supplementary Fig. 5). Hence, important differences must be subtle. In an initial effort to compare the two proteins we made a series of hybrid σ subunits with different domain combinations. However, in all cases, these hybrid σ factors were poorly active. Hence, in a more nuanced approach, we examined changes between σ^{70} and σ^{A} in the context of the E. coli RNA polymerase structure^{3,45}. In particular, we searched for differences in surfaceexposed amino acid side chains that might alter interactions with core RNA polymerase or the promoter DNA. This identified two residues of interest. Side chain R157 of σ^{70} is located in the NCR and so absent from σ^A . In the context of the closed holoenzymepromoter complex, R157 is 14.5 Å from the DNA backbone. Conformational changes during DNA opening place R157 just 4.1 Å from the DNA. Narayanan et al. previously showed that R157 contacts the DNA backbone in this context to impact DNA opening⁴⁵. Side chain R486 of σ^{70} is located in region 3.2 of domain σ_3 . In the closed complex R486 is 10.6 Å from the DNA and moves to a more distal 22.6 Å in the open complex. In B. subtilis, σ^{A} residue D222, corresponding to σ^{70} R486, is negatively charged. We altered σ^{70} , with mutations R157A and R486D, to more closely resemble σ^A , and refer to this derivative as $\sigma^{70 \text{ Mut}}$. First, we compared the properties of σ^{70} , σ^{A} , and σ^{70} Mut at the B. subtilis veg promoter using either KMnO₄ footprinting (to detect DNA opening) or in vitro transcription assays. Each σ factor was tested with the core RNA polymerases of both B. subtilis and E. coli. As expected, holoenzymes containing the B. subtilis core RNA polymerase formed less stable open complexes than those containing E. coli core enzyme (compare lanes 1-3 with 4-6 in Fig. 5c). We observed subtle differences in patterns of DNA opening for σ^{70} and σ^{A} . For instance, when associated with B. subtilis core RNA polymerase, two sites of KMnO₄ reactivity were observed for each σ factor. These were at promoter positions -1 and -3 (lanes 1-3). Compared to σ^{70} , σ^{A} resulted in greater KMnO₄ reactivity at promoter position -3 (compare lanes 1 and 2). In this context, σ^{70} Mut behaviour resembled σ^{A} rather than σ^{70} (lane 3). Whilst more extensive DNA opening was observed in the context of the *E. coli* core RNA polymerase the different σ factors again resulted in different KMnO₄ reactivity at the -3 position (lanes 4-6). However, in this instance, σ^{70} generated a greater -3 signal (lane 5). Again, $\sigma^{70 \text{ Mut}}$ behaved like σ^{A} rather than σ^{70} . We also observed differences in the amount of transcript generated by RNA polymerase with each σ factor (Fig. 5d). There were significant differences in behaviour σ^A and σ^{70} but not between σ^{A} and $\sigma^{70 \text{ Mut}}$.

Mutation of σ^{70} side chains R156 and R486 improves specificity at high AT-content DNA templates. To determine if $\sigma^{70\,\mathrm{Mut}}$ directed less promiscuous transcription we compared σ factor

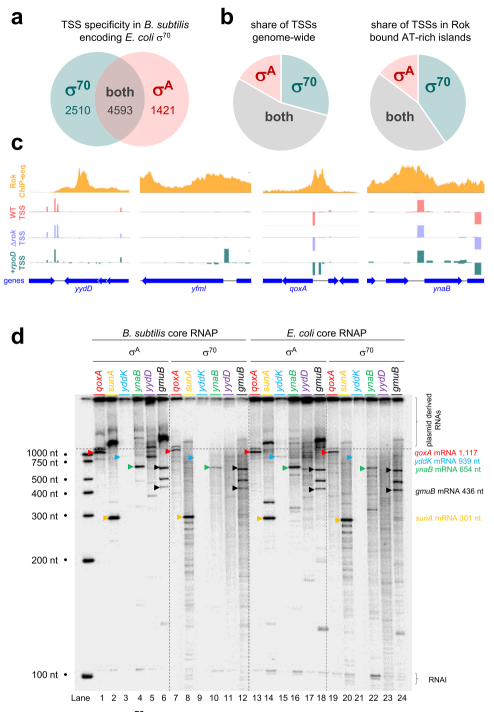
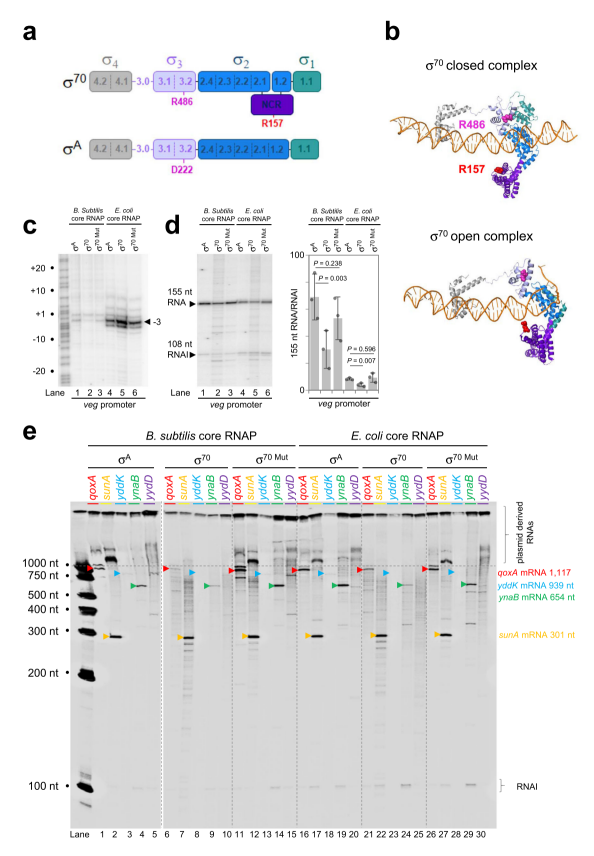


Fig. 4 Expression of the *E. coli* **RNA polymerase** σ^{70} **subunit is sufficient for promiscuous transcription in** *B. subtilis.* **a** The Venn diagram shows the distribution of TSSs identified in *B. subtilis* by cappable-seq in different genetic backgrounds. The teal area represents TSSs only detected upon expression of *E. coli rpoD* whilst the pink section represents TSSs only identified in the absence of *rpoD* expression. The overlap identifies those TSSs detected both with and without *rpoD* expression. **b** The pie charts show the distribution of *B. subtilis* TSSs identified in different genetic backgrounds and in different parts of the genome. The number of TSSs dependent on *E. coli* σ^{70} expression is higher in horizontally acquired AT-rich sections of DNA targeted by Rok. Conversely, the number of σ^A dependent TSSs is lower in these regions. **c** Examples of *E. coli* σ^{70} dependent transcription initiation within horizontally acquired *B. subtilis* genes. Data from ChIP-seq experiments for Rok occupancy³⁵ are shown by the orange graph. Transcription start sites (TSSs) were identified by cappable-seq for wild-type (pink graph), Δrok (mauve graph) and *B. subtilis* cells carrying the σ^{70} encoding *rpoD* gene (teal graph). Sequence reads mapping to the top and bottom DNA strands are shown above and below the central horizontal line in each plot. Genes are indicated with block blue arrows. **d** Results of in vitro transcription assays using AT-rich horizontally acquired genes targeted by Rok cloned in plasmid pSR as the DNA template. Transcription reactions were done using core RNA polymerase, from either *B. subtilis* or *E. coli*, in conjunction with either σ^{A} or σ^{70} (0.5 μM final holoenzyme concentration). For each DNA template, bands corresponding to full-length mRNAs are indicated by coloured arrowheads. The experiment was done twice with similar results.



behaviour with DNA templates derived from Rok targeted genes (Fig. 5e). As described above, σ^A specifically stimulated the initiation of transcription at intergenic *B. subtilis* promoters to produce mRNAs (marked with coloured triangles, lanes 1–5 and 16–20). Conversely, σ^{70} resulted in promiscuous initiation (lanes 6–10 and 21–25). The promiscuity of σ^{70} Mut was reduced; we

detected greater production of expected mRNA species and reduced transcription of spurious RNAs (lanes 11–15 and 26–30). We conclude that σ^{70} residues R156 and R486, which influence patterns of DNA opening, are important for the more promiscuous behaviour of core RNA polymerase associated with $\it E.~coli~\sigma^{70}$.

Fig. 5 Mutation of E. coli σ^{70} to resemble B. subtilis σ^{A} reduces transcriptional promiscuity. a Schematic representations of E. coli σ^{70} and B. subtilis σ^{A} . Individual domains are labelled σ^1 through σ^4 and the non-conserved region (NCR) specific to E. coli σ^{70} is also shown. Sub-regions of each σ factor are labelled 1.1 through 4.2 and are separated by dashed lines where required. Side chains R157 and R486 important for the promiscuous behaviour of σ^{70} are shown. Side chain D222 of σ^A is in the position equivalent to that of R486 in σ^{70} . **b** Location of R157 and R486 in *E. coli* σ^{70} RNA polymerase bound to promoter DNA. The top and bottom images are derived from PDB accession numbers 6PSO3 and 6CAO45 respectively. Whilst present in the structures. RNA polymerase core enzyme has been hidden from view for clarity. Colour coding of σ^{70} matches (a) and DNA is shown in orange. c The E. coli σ^{70} Mut derivative has DNA opening properties similar to B. subtilis σ^A . The gel image shows KMnO₄ reactivity patterns at the B. subtilis veg promoter due to DNA opening by B. subtilis (lanes 1-3) or E. coli (lanes 4-6) core RNA polymerase in complex with either σ^A , σ^{70} or σ^{70} Mut as indicated (0.5 μ M final holoenzyme concentration). The gel is calibrated with a Maxam-Gilbert G + A sequencing reaction. The experiment was done twice with similar results. d Results of in vitro transcription assays using B. subtilis (lanes 1-3) or E. coli (lanes 4-6) core RNA polymerase in complex with either σ^A , σ^{70} or σ^{70} Mut as indicated (0.5 µM final holoenzyme concentration). The 155 nucleotide (nt) transcript is generated from the B. subtilis veg promoter and the 108 nt RNAI transcript is derived from the plasmid replication origin. The gel image shows a representative result and the bar chart shows a quantification of 3 independent experiments. Error bars show S.D. and the centre of the error bars is the mean. P was calculated using a two-tailed student's T-test. e Results of in vitro transcription assays using AT-rich horizontally acquired genes targeted by Rok cloned in plasmid pSR as DNA templates. Transcription reactions were done using core RNA polymerase, from either B. subtilis or E. coli, in conjunction with either σ^A , σ^{70} or σ^{70} Mut (0.5 μ M final holoenzyme concentration). For each DNA template, bands corresponding to full-length mRNAs are indicated by coloured arrowheads. The experiment was done twice with similar results.

The B. subtilis δ subunit also contributes to specific transcription of horizontally acquired genes. The B. subtilis RNA polymerase contains two small ancillary subunits, named δ and ϵ , that are not found in E. coli⁴. Comparatively, the role of these subunits is poorly understood. However, previous work has implicated δ in control of transcriptional specificity and DNA opening $^{40,46-49}$. Thus, in a final experiment, we remapped B. subtilis TSSs in the absence of δ. This identified 6786 TSSs and 1916 of these were unique to cells lacking the δ encoding rpoE gene. Interestingly, GTP was the most frequently used initiating nucleotide triphosphate (iNTP) at TSSs specific to the $\Delta rpoE$ strain (47% of TSSs). Conversely, ATP was the most common iNTP globally (52% of TSSs). This is intriguing since Rabatinová and co-workers implicated δ in the control of transcription by iNTP concentration⁵⁰. There were no other sequence differences (Supplementary Fig. 6a). A greater proportion of TSSs in Rok targeted AT-rich islands were specific to $\Delta rpoE$ cells compared to the rest of the genome (Supplementary Fig. 6b). An example is shown in Supplementary Fig. 6c.

Discussion

We propose fundamentally different xenogeneic silencing strategies in E. coli and B. subtilis (Fig. 6). In the former, RNA polymerase promiscuity creates a need for widespread transcriptional silencing by H-NS^{14,15,18,34}. Hence, xenogeneic silencing inhibits both mRNA expression and spurious intragenic transcription initiation¹⁴. Conversely, Rok binds mostly to noncoding DNA and better RNA polymerase specificity reduces intragenic transcription (Figs. S1 and 5). We attribute differences in RNA polymerase promiscuity primarily to the housekeeping σ factor (Figs. 4 and 5). Consistent with this, Rok is unable to block intragenic transcription when E. coli σ^{70} is expressed in B. subtilis (Supplementary Fig. 4). Similarly, compared to σ^{70} , σ^{A} imposes greater specificity on core E. coli RNA polymerase (Figs. 4 and 5). Previous work, with a small number of model promoters, also noted the greater specificity of B. subtilis housekeeping RNA polymerase 38,51,52. One such study concluded that core enzyme, not the associated σ factor, enforced greater specificity⁴². The latter conclusion was based on in vitro analysis of two phage promoters, \(\lambda \) PR' and T7A1. We do not exclude a role for core RNA polymerase in promoter selection related to transcriptional promiscuity. Indeed, core RNA polymerase has a clear influence on DNA opening in our assays (Fig. 5c). However, the ability of E. coli σ^{70} to divert B. subtilis core enzyme to many different promoter sequences, both in vivo and in vitro, suggests σ must play a key role (Figs. 4 and 5). Consistent with prior

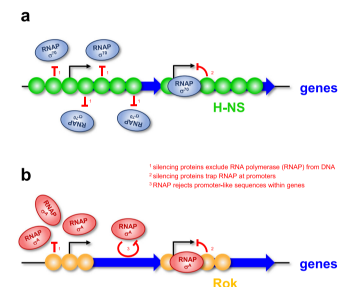


Fig. 6 Different xenogeneic silencing strategies in *E. coli* **and** *B. subtilis.* **a** In *E. coli*, H-NS (green) binds to extensive tracts of DNA. Consequently, the comparatively promiscuous *E. coli* housekeeping RNA polymerase (blue ovals) is prevented from synthesising mRNAs, and many spurious intragenic RNAs, from sections of AT-rich horizontally acquired DNA (blue block arrows). H-NS may repress transcription by blocking access of RNA polymerase to the DNA and by trapping RNA polymerase at promoters (bent arrows). **b** In *B. subtilis*, Rok (orange) binds to shorter tracts of DNA around promoters (bent arrows) at gene 5' ends. This stops the synthesis of mRNAs from sections of AT-rich horizontally acquired DNA (blue block arrows). The less promiscuous housekeeping RNA polymerase of *B. subtilis* containing σ^A (red ovals) is not prone to spurious intragenic transcription initiation. Trapping of RNA polymerase early during elongation appears commonplace.

reports, the *B. subtilis* δ subunit also contributes^{7,53}. However, σ is dominant and impacts spurious intragenic promoters even when δ is present (Figs. 3 and 4). Our genome-scale TSS analyses identified sequence properties consistent with prior work^{38,50,52}. For example, Henkin and Sonenshein noted that an AT-rich discriminator allowed the *E. coli lacUV5* promoter to be used by *B. subtilis* holoenzyme⁵¹. We speculate that greater specificity of *B. subtilis* RNA polymerase holoenzyme may have evolved to accommodate a higher AT-content genome in this organism.

Together, our cappable-seq and RNA-seq analyses suggest that Rok primarily silences mRNA transcription early during elongation; TSS signals for Rok silenced mRNAs are frequently similar both in the presence and absence of Rok. Conversely, there are substantial differences in signals for full-length mRNAs in RNA-seq experiments. Figures 1d and 2a both illustrate this behaviour that is also consistent with global patterns of transcription (Fig. 1e). In *E. coli*, H-NS can silence transcription both by excluding RNA polymerase from the DNA template and by trapping RNA polymerase at the promoter⁵⁴. This property of H-NS reflects the protein's ability to switch between linear and bridging modes of DNA binding. Hence, we speculate that Rok may be less versatile in this regard. Consistent with this, biophysical measurements indicate that Rok is a DNA bridging protein⁵⁵.

The differences in promiscuity of *E. coli* and *B. subtilis* RNA polymerase are intriguing given the different role of the transcription termination factor Rho in these organisms. In *E. coli*, Rho is essential and reducing Rho activity results in greater spurious transcription across horizontally acquired genes. Numerous genetic interactions between *hns* and *rho* have been reported and are consistent with the factors working together to suppress unwanted transcription^{56,57}. By sharp contrast, Rho is not essential in *B. subtilis*⁵⁸. Indeed, loss of *rho* has only a minor impact on cell fitness⁵⁸. In conclusion, natural selection has identified diverse strategies for managing horizontally acquired DNA in different bacteria. Most likely, the correct balance of silencing factors and transcriptional specificity is essential.

Methods

Strains, plasmids, and oligonucleotides. Bacterial strains, plasmids, synthesised gene strands and oligonucleotides used in this study are listened in Supplementary Table 1. Bacterial cultures were grown at 37 °C in Lennox Broth (LB) medium. Cloning was done using HiFi Gibson assembly (New England Biolabs). Insert DNA was either synthesised as gene strands (Eurofins Genomics) or amplified from genomic DNA using oligonucleotides (Eurofins Genomics). The *B. subtilis rpoD* expression strain was constructed by cloning *E. coli rpoD*, with an optimal ribosome binding site, in plasmid pDR111. The resulting constructs were used to transform *B. subtilis* 168c and double crossover integration at the chromosomal *amyE* locus was confirmed by iodine testing patched cells grown on LB agar with 1 % (*w/v*) starch.

Cappable-seq and RNA-seq. All experiments were done twice using total RNA isolated from biological replicates. Strains were grown in LB with shaking at 37 °C until mid-log phase. Aliquots of 2 ml were then pelleted and flash frozen in liquid nitrogen. For expression of *rpoD* in *B. subtilis*, 3 mM IPTG (final) was added at exponential phase for 1 h before harvesting cells. For both *B. subtilis* and *E. coli* RNA-seq and cappable-seq experiments were done as described in Warman et al.³⁷. Vertis Biotechnologie AG (Germany) did library preparation steps followed by sequencing with an Illumina NextSeq 500 system (75 bp read length). Raw data in FASTQ format are available from ArrayExpress (accession number E-MTAB-10777, https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-10777/).

Bioinformatics. Individual sequence reads in FASTQ files were aligned to the reference genomes NC000964.3 (B. subtilis) or U00096.3 (E. coli) using Bowtie2 (Galaxy version 2.4.2)⁵⁹. Coverage for each genome position was extracted from resulting Binary Alignment Map (BAM) files using the genomcov function of BedTools (Galaxy version 2.30.0)60. For each strand, coverage was used to call TSSs at positions where read depth increased more than threefold, compared to the previous base, in both replicates. Positions where the read depth was zero are excluded to avoid erroneous TSS selection. To generate DNA sequence logos, sequences upstream of the desired TSSs were aligned by TSS position and submitted to WebLogo (version 2.8.2)⁶¹. Standard settings were used. The R package GenomicRanges (version 1.44)62 was used to identify which TSSs fell within different genomic contexts (e.g. H-NS or Rok bound regions) using the coordinates in Supplementary Data 3. FeatureCounts (version 2.6)⁶³ of the Rsubread (version 2.6)⁶⁴ package was used to determine gene read counts, which were inputted into the exact function of edgeR (Galaxy version 3.34.0)65 to determine differential gene expression. For differential TSS activity, TSSs for the two samples to be compared were pooled, with duplicate TSSs removed. The coverage at each TSS position on the genome was calculated for biological duplicates submitted to the exact function of edgeR⁶⁵. Note that edgeR automatically adjusts for sequencing depth discrepancies when calculating fold-changes and P values.

To quantify differences in Rok and H-NS distribution we used existing ChIP-seq data^{32,35}. Binding signals for each factor were averaged every 10 bp across the entire genome. The average ChIP-seq signal for all 10 bp bins was then calculated and this value was subtracted from each bin to remove background binding signals. The binned data were processed in two different ways. First, the distance between each 10 bp bin and the nearest gene start codon was determined. The sum of ChIP-seq binding signals was then calculated for all bins falling in a given range of distances upstream or downstream of a start codon (Supplementary Fig. 1c). Second, we determined the ChIP-seq signal for each bin as a percentage of the maximum ChIP-seq signal for H-NS or Rok. These values were compared to the percentage of bins locating to non-coding DNA in Supplementary Fig. 1d. Distances between promoter —10 elements and TSSs were determined as described previously³⁷.

Proteins. B. subtilis RNA polymerase was purified from a strain expressing a chromosomal β' C-terminal domain His6 tag fusion66. B. subtilis cells were grown at 37 °C with shaking to late exponential phase in LB medium supplemented with 1% (w/v) glucose. Cell pellets were resuspended in lysis buffer (50 mM Tris pH 7.9, 300 mM NaCl, 3 mM β Mercaptoethanol, 5% and cOmplete EDTA-free protease inhibit tablet) and lysed by sonication. Lysate was clarified by centrifugation at $45,000 \times g$ for 20 min at 4 °C followed by filtration through a 0.45 μ M PES filter. Lysates were applied to a His-Trap HP column (GE Healthcare). Unbound protein was removed by washing the column with lysis buffer followed by lysis buffer with 25 mM imidazole. Bound protein was eluted in a gradient to 200 mM imidazole in lysis buffer. RNA polymerase containing fraction eluted between 120 and 150 mM imidazole. These fractions were pooled, diluted 3-fold in HiTrap buffer (40 mM Tris pH 7.9, 1 mM EDTA, 5% glycerol) and loaded onto a HiTrap Q column. RNA polymerase was eluted in a gradient from 100 mM to 1 M NaCl. RNA polymerase eluting at ~500 mM NaCl was pooled, concentrated, and made to 50% glycerol for -20 °C storage. E. coli RNA polymerase was purchased from New England Biolabs and σ^{70} was purified as described previously 16. The B. subtilis sigA gene, cloned in pET-28a, was used to transform T7 Express E. coli (New England Biolabs). Cells were grown in LB to exponential phase at 37 °C with shaking before protein expression was induced for 3 h with 1 mM IPTG. Overexpressed of forms inclusion bodies, which were isolated as outlined by Borukhov and Goldfarb⁶⁷. Inclusion bodies were solubilised in denaturing His-Trap buffer (20 mM Tris pH 7.9, 5% glycerol, 600 mM NaCl, 8 M urea) and cell debris removed by centrifugation at 45,000 × g for 20 min. Solubilised inclusion bodies were loaded on a His-Trap column (GE Healthcare), which was washed with denaturing His-Trap buffer then stepwise increasing concentrations of imidazole. Fractions containing σ^A were pooled and diluted with an equal volume of denaturing buffer (50 mM Tris pH 7.9, 8 M urea, 10% glycerol, 10 mM MgCl₂, 10 μM ZnCl₂, 1 mM EDTA, 10 mM DTT) before dialysis overnight in 21 of reconstitution buffer (50 mM Tris pH 7.9, 200 mM NaCl, 20% glycerol, 10 mM MgCl₂, 10 μM ZnCl₂, 1 mM EDTA, 1 mM DTT). Refolded σ^A was diluted 4 times in HiTrap buffer and loaded onto a HiTrap Q column (GE Healthcare). σ^A was eluted in HiTrap buffer with a gradient of 50 mM NaCl to 1 M NaCl. Fractions containing σ^A were pooled, concentrated, and made to 50% glycerol for −20 °C storage.

The rok gene was cloned in pET-21a and used to transform $E.\ coli$ T7 Express (New England Biolabs). Cells were grown in LB to exponential phase at 37 °C with shaking before protein expression was induced for 3 h with 1 mM IPTG. Cell pellets were resuspended in His-Trap buffer (20 mM Tris pH 7.9, 5% glycerol, 600 mM NaCl) and lysed by sonication. Lysate was clarified by centrifugation at $45,000 \times g$ for 20 min at 4° C followed by filtration through a $0.45 \ \mu M$ PES filter. Lysate was applied to a His-Trap column which was then washed with His-Trap buffer followed by stepwise increases in imidazole concentration. Protein eluted by 200 mM imidazole was diluted three times in 20 mM Tris HCl pH 7.9, 5% (v/v) glycerol and applied to a Heparin column (GE Healthcare). Protein was eluted with step wise increasing concentrations of NaCl ranging from 150 mM NaCl to 1 M NaCl. The fractions containing Rok were pooled, concentrated, and made to 50% glycerol for -20 °C storage. $E.\ coli$ H-NS was purified as described previously 16.

in vitro transcription assays. Each in vitro transcription reaction contained 0.01 µM DNA template, 0.05 µM B. subtilis RNAP/0.5 units E. coli core RNAP (New England Biolabs), 0.25 µM $\sigma^{\rm A}/\sigma^{\rm 70}$ in Transcription buffer (20 mM Tris pH 7.9, 40 mM KCl, and 10 mM MgCl₂). Reactions were started by addition of NTP mix to give final NTP concentrations of 200 µM ATP/GTP/CTP, 10 µM UTP and 2 µCi [$\alpha^{\rm -32}$ P] UTP. After 10 min incubation at 37 °C, reactions were stopped by the addition of an equal volume of formamide containing stop buffer. DNA templates consisted of the promoter and/or gene of interest cloned in plasmid pSR. Plasmid templates were isolated from E. coli using Qiagen Maxiprep kits. Reactions were resolved on an 8% (w/v) denaturing polyacrylamide gel, exposed on a Bio-Rad phosphor screen then visualised on a Bio-Rad Personal Molecular Imager using Quantity One software (version 4.6.9).

Potassium permanganate footprinting. KMnO₄ footprints were done using standard procedures with the following alterations ¹⁶. DNA fragments were made by polymerase chain reaction amplification of pVEG cloned in plasmid pSR and labelled with 50 μ Ci [γ -³²P] ATP using T4 Polynucleotide Kinase (New England

Biolabs). Reactions contained 10 nM labelled DNA, 100 nM RNAP, 500 nM $\sigma^{70}/$ $\sigma^A,$ 10 mM HEPES pH 8, 20 mM CH₃COOK, 0.1 mM DTT, 10 mM Mg(C₂H₃O₂)₂ and 100 μg ml $^{-1}$ BSA.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The raw RNA-seq and cappable-seq data generated in this study have been deposited in ArrayExpress under accession code E-MTAB-10777. Reference genomes NC000964.3 or U00096.3 (https://www.ncbi.nlm.nih.gov/nuccore/545778205) were used as appropriate. Results generated from processing of the sequencing data (e.g. to determine changes in gene expression or TSS signals) are available as Source Data or Supplementary Data files. The ChIP-seq data used were obtained from the ArrayExpress or NCBI databases for E. coli H-NS (E-MTAB-332) or B. subtilis Rok (PRJNA272948) respectively. All Original gel images are in Supplementary Fig. 7 and results obtained from quantification of band intensities are provided as Source Data. Source data are provided with this paper.

Received: 9 September 2021; Accepted: 7 February 2022; Published online: 03 March 2022

References

- Browning, D. F. & Busby, S. J. W. The regulation of bacterial transcription initiation. *Nat. Rev. Microbiol.* 2, 57–65 (2004).
- Zhang, G. et al. Crystal structure of Thermus aquaticus core RNA polymerase at 3.3 Å resolution. Cell 98, 811–824 (1999).
- Chen, J. et al. Stepwise promoter melting by bacterial RNA polymerase. Mol. Cell 78, 275–288.e6 (2020).
- Weiss, A. & Shaw, L. N. Small things considered: the small accessory subunits of RNA polymerase in Gram-positive bacteria. FEMS Microbiol. Rev. 39, 541–554 (2015).
- Campbell, E. A. et al. Structure of the bacterial RNA polymerase promoter specificity σ subunit. Mol. Cell 9, 527–539 (2002).
- Feklistov, A. et al. RNA polymerase motions during promoter melting. Science 356, 863–866 (2017).
- Dobinson, K. F. & Spiegelman, G. B. Effect of the δ subunit of *Bacillus subtilis* RNA polymerase on initiation of RNA synthesis at two bacteriophage φ29 promoters. *Biochemistry* 26, 8206–8213 (1987).
- Artsimovitch, I., Svetlov, V., Murakami, K. S. & Landick, R. Co-overexpression of *Escherichia coli* RNA polymerase subunits allows isolation and analysis of mutant enzymes lacking lineage-specific sequence insertions. *J. Biol. Chem.* 278, 12344–12355 (2003).
- Reppas, N. B., Wade, J. T., Church, G. M. M. & Struhl, K. The transition between transcriptional initiation and elongation in *E. coli* is highly variable and often rate limiting. *Mol. Cell* 24, 747–757 (2006).
- Dorman, C. J. H-NS, the genome sentinel. Nat. Rev. Microbiol. 5, 157–161 (2007)
- 11. Blattner, F. R. et al. The complete genome sequence of *Escherichia coli* K-12. *Science* 277, 1453–1462 (1997).
- 12. Kunst, F. et al. The complete genome sequence of the gram-positive bacterium *Bacillus subtilis. Nature* **390**, 249–256 (1997).
- Doyle, M. et al. An H-NS-like stealth protein aids horizontal DNA transmission in bacteria. Science 315, 251–252 (2007).
- Singh, S. S. et al. Widespread suppression of intragenic transcription initiation by H-NS. Genes Dev. 28, 214–219 (2014).
- Lamberte, L. E. et al. Horizontally acquired AT-rich genes in *Escherichia coli* cause toxicity by sequestering RNA polymerase. *Nat. Microbiol.* 2, 16249 (2017).
- Singh, S. S. & Grainger, D. C. H-NS can facilitate specific DNA-binding by RNA polymerase in AT-rich gene regulatory regions. *PLoS Genet.* 9, e1003589 (2013).
- Oshima, T., Ishikawa, S., Kurokawa, K., Aiba, H. & Ogasawara, N. Escherichia coli histone-like protein H-NS preferentially binds to horizontally acquired DNA in association with RNA polymerase. DNA Res. 13, 141–153 (2006).
- Landick, R., Wade, J. T. & Grainger, D. C. H-NS and RNA polymerase: a lovehate relationship? Curr. Opin. Microbiol. 24, 53–59 (2015).
- Kotlajich, M. V. et al. Bridged filaments of histone-like nucleoid structuring protein pause RNA polymerase and aid termination in bacteria. *Elife* 4, e04970 (2015).
- Singh, K., Milstein, J. N. & Navarre, W. W. Xenogeneic silencing and its impact on bacterial genomes. Annu. Rev. Microbiol. 70, 199–213 (2016).
- Tendeng, C. & Bertin, P. N. H-NS in Gram-negative bacteria: a family of multifaceted proteins. *Trends Microbiol.* 11, 511–518 (2003).

- Castang, S., McManus, H. R., Turner, K. H. & Dove, S. L. H-NS family members function coordinately in an opportunistic pathogen. *Proc. Natl Acad.* Sci. USA 105, 18947–18952 (2008).
- Ding, P. et al. A novel AT-rich DNA Recognition mechanism for bacterial xenogeneic silencer MvaT. PLoS Pathog. 11, 1004967 (2015).
- Ali, S. S., Xia, B., Liu, J. & Navarre, W. W. Silencing of foreign DNA in bacteria. Curr. Opin. Microbiol. 15, 175–181 (2012).
- Lippa, A. M., Gebhardt, M. J. & Dove, S. L. H-NS-like proteins in Pseudomonas aeruginosa coordinately silence intragenic transcription. Mol. Microbiol. 115, 1138–1151 (2021).
- Gordon, B. R. G. et al. Structural basis for recognition of AT-rich DNA by unrelated xenogeneic silencing proteins. *Proc. Natl Acad. Sci. USA* 108, 10690–10695 (2011).
- Gehrke, E. J. et al. Silencing cryptic specialized metabolism in Streptomyces by the nucleoid-associated protein Lsr2. Elife 8, e47691 (2019).
- Smits, W. K. & Grossman, A. D. The transcriptional regulator Rok binds A +T-rich DNA and is involved in repression of a mobile genetic element in Bacillus subtilis. PLoS Genet. 6, e1001207 (2010).
- Duan, B. et al. How bacterial xenogeneic silencer rok distinguishes foreign from self DNA in its resident genome. *Nucleic Acids Res.* 46, 10514–10529 (2018).
- Hoa, T. T., Tortosa, P., Albano, M. & Dubnau, D. Rok (YkuW) regulates genetic competence in Bacillus subtilis by directly repressing comK. *Mol. Microbiol.* 43, 15–26 (2002).
- Ettwiller, L., Buswell, J., Yigit, E. & Schildkraut, I. A novel enrichment strategy reveals unprecedented number of novel transcription start sites at single base resolution in a model prokaryote and the gut microbiome. *BMC Genomics* 17, 199 (2016).
- Kahramanoglou, C. et al. Direct and indirect effects of H-NS and Fis on global gene expression control in *Escherichia coli. Nucleic Acids Res.* 39, 2073–2091 (2011).
- Duan, B., Ding, P., Navarre, W. W., Liu, J. & Xia, B. Xenogeneic silencing and bacterial genome evolution: mechanisms for DNA recognition imply multifaceted roles of xenogeneic silencers. *Mol. Biol. Evol.* 38, 4135–4148 (2021).
- Warman, E. A., Singh, S. S., Gubieda, A. G. & Grainger, D. C. A non-canonical promoter element drives spurious transcription of horizontally acquired bacterial genes. *Nucleic Acids Res.* 48, 4891–4901 (2020).
- Seid, C. A., Smith, J. L. & Grossman, A. D. Genetic and biochemical interactions between the bacterial replication initiator DnaA and the nucleoidassociated protein Rok in *Bacillus subtilis*. *Mol. Microbiol.* 103, 798–817 (2017)
- Hawley, D. K. & Mcclure, W. R. Compilation and analysis of Escherichia coli promoter DNA sequences. Nucleic Acids Res. 11, 2237–2255 (1983).
- Warman, E. A. et al. Widespread divergent transcription from bacterial and archaeal promoters is a consequence of DNA-sequence symmetry. *Nat. Microbiol.* 6, 746–756 (2021).
- Helmann, J. D. Compilation and analysus of Bacillus Subtilis σA-dependent promoter sequences: evidence for extended contact between RNA polymerse and upstream promoter DNA. Nucleic Acids Res. 23, 2351–2360 (1995).
- Moran Jr, C. P. et al. Nucleotide sequences that signal the initiation of transcription and translation in *Bacillus subtilis. Mol. Gen. Genet.* 186, 339–346 (1982).
- Juang, Y. L. & Helmann, J. D. The δ subunit of *Bacillus subtilis* RNA polymerase: an allosteric effector of the initiation and core-recycling phases of transcription. *J. Mol. Biol.* 239, 1–14 (1994).
- 41. Whipple, F. W. & Sonenshein, A. L. Mechanism of initiation of transcription by *Bacillus subtilis* RNA polymerase at several promoters. *J. Mol. Biol.* **223**, 399–414 (1992).
- Artsimovitch, I., Svetlov, V., Anthony, L., Burgess, R. R. & Landick, R. RNA polymerases from *Bacillus subtilis* and *Escherichia coli* differ in recognition of regulatory signals in vitro. *J. Bacteriol.* https://journals.asm.org/journal/jb (2000)
- 43. Bervoets, I. et al. A sigma factor toolbox for orthogonal gene expression in *Escherichia coli. Nucleic Acids Res.* 46, 2133–2144 (2018).
- Gaida, S. M. et al. Expression of heterologous sigma factors enables functional screening of metagenomic and heterologous genomic libraries. *Nat. Commun.* 6, 1–10 (2015).
- 45. Narayanan, A. et al. Cryo-EM structure of *Escherichia coli* σ 70 RNA polymerase and promoter DNA complex revealed a role of σ non-conserved region during the open complex formation. *J. Biol. Chem.* **293**, 7367–7375 (2018).
- Pero, J., Nelson, J. & Fox, T. D. Highly asymmetric transcription by RNA polymerase containing phage-SP01-induced polypeptides and a new host protein. PNAS 72, 1589 (1975).
- Achberger, E. C., Hilton, M. D. & Whiteley, H. R. The effect of the delta snbunit on the interaction of Bacillus subtilis RNA polymerase with bases in a SP82 early gene promoter. *Nucleic Acids Res.* 10, 2893–2910 (1982).

- Dickel, C. D., Burtis, K. C. & Doi, R. H. Delta factor increases promoter selectivity by *Bacillus subtilis* vegetative cell RNA polymerase. *Biochem. Biophys. Res. Commun.* 95, 1789–1795 (1980).
- Tjian, R., Losick, R., Pero, J. & Hinnebush, A. Purification and comparative properties of the delta and sigma subunits of RNA polymerase from *Bacillus* subtilis. Eur. J. Biochem. 74, 149–154 (1977).
- 50. Rabatinová, A. et al. The δ subunit of RNA polymerase is required for rapid changes in gene expression and competitive fitness of the cell. *J. Bacteriol.* **195**, 2603–2611 (2013).
- Henkin, T. M. & Sonenshein, A. L. Mutations of the Escherichia coli lacUV5 promoter resulting in increased expression in Bacillus subtilis. MGG Mol. Gen. Genet. 209, 467–474 (1987).
- Graves, M. C. & Rabinowitz, J. C. In vivo and in vitro transcription of the Clostridium pasteurianum ferredoxin gene. Evidence for 'extended' promoter elements in Gram-positive organisms. J. Biol. Chem. 261, 11409–11415 (1986).
- Achberger, E. C. & Whiteley, H. R. The role of the delta peptide of the Bacillus subtilis RNA polymerase in promoter selection. *J. Biol. Chem.* 256, 7424–7432 (1981).
- Grainger, D. C. Structure and function of bacterial H-NS protein. Biochem. Soc. Trans. 44, 1561–1569 (2016).
- Qin, L., Erkelens, A. M., Markus, D. & Dame, R. T. The B. subtilis Rok protein compacts and organizes DNA by bridging. bioRxiv https://doi.org/10.1101/ 769117 (2019).
- Tran, L., van Baarsel, J. A., Washburn, R. S., Gottesman, M. E. & Miller, J. H. Single-gene deletion mutants of *Escherichia coli* with altered sensitivity to bicyclomycin, an inhibitor of transcription termination factor Rho. *J. Bacteriol.* 193, 2229–2235 (2011).
- Saxena, S. & Gowrishankar, J. Modulation of Rho-dependent transcription termination in *Escherichia coli* by the H-NS family of proteins. *J. Bacteriol.* 193, 3832–3841 (2011).
- Quirk, P. G., Dunkley, E. A., Lee, P. & Krulwich, T. A. Identification of a putative *Bacillus subtilis* rho gene. *J. Bacteriol.* 175, 647–654 (1993).
- Langmead, B. & Salzberg, S. L. Fast gapped-read alignment with Bowtie 2. Nat. Methods 9, 357–359 (2012).
- Quinlan, A. R. & Hall, I. M. BEDTools: A flexible suite of utilities for comparing genomic features. *Bioinformatics* 26, 841–842 (2010).
- Crooks, G. E., Hon, G., Chandonia, J. M. & Brenner, S. E. WebLogo: A sequence logo generator. Genome Res. 14, 1188–1190 (2004).
- Lawrence, M. et al. Software for computing and annotating genomic ranges. PLOS Comput. Biol. 9, e1003118 (2013).
- Liao, Y., Smyth, G. K. & Shi, W. featureCounts: an efficient general purpose program for assigning sequence reads to genomic features. *Bioinformatics* 30, 923–930 (2014).
- Liao, Y., Smyth, G. K. & Shi, W. The R package Rsubread is easier, faster, cheaper and better for alignment and quantification of RNA sequencing reads. Nucleic Acids Res. 47, e47–e47 (2019).
- Robinson, M. D., McCarthy, D. J. & Smyth, G. K. edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. *Bioinformatics* 26, 139–140 (2010).
- 66. Qi, Y. & Hulett, F. M. PhoP~P and RNA polymerase σ(A) holoenzyme are sufficient for transcription of Pho regulon promoters in Bacillus subtilis: PhoP~P activator sites within the coding region stimulate transcription in vitro. *Mol. Microbiol.* 28, 1187–1197 (1998).

 Borukhov, S. & Goldfarb, A. Recombinant *Escherichia coli* rna polymerase: purification of individually overexpressed subunits and in vitro assembly. *Protein Expr. Purif.* 4, 503–511 (1993).

Acknowledgements

This work was funded by a Wellcome Trust Investigator award (no. 212193/Z/18/Z) and a Leverhulme Trust project grant (no. RPG-2018-198) to D.C.G. We thank Joseph Wade, Steve Busby, and John Helmann for critical reading of the manuscript prior to submission. We also thank James Haycocks for proofreading the final version of the text

Author contributions

D.F., E.A.W. and D.C.G. designed the project. All experimental work was done by D.F. with support and advice from E.A.W. Bioinformatics was done by D.F. and D.C.G. A.E.M. and R.T.D. provided materials. D.F. and D.C.G. wrote the paper with all authors making edits as appropriate.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-022-28747-1.

Correspondence and requests for materials should be addressed to David C. Grainger.

Peer review information Nature Communications thanks William Navarre and the other, anonymous, reviewers for their contribution to the peer review of this work. Peer reviewer reports are available.

Reprints and permission information is available at http://www.nature.com/reprints

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 https://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022