Supplemental Material

Transcript Degradation and Noise of Small RNA-Controlled Genes in a Switch Activated Network in *Escherichia coli*

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Supplemental Table of contents

Figure S1  Discrete nature of the intensity of spatially separable fluorescence spots.
Figure S2  Effects of different mutant strains on the stoichiometric degradation of sodB and fumA transcripts with iron deprivation.
Figure S3  Switch-like effects of iron deprivation on mean sodB levels obtained from qRT-PCR experiments.
Figure S4  Effects of iron addition to an iron-free medium on the promoter activity of RyhB and one of its targets.
Figure S5  Switch-like behavior of mean sufC transcript levels as function of iron deprivation.
Figure S6  Effects of iron deprivation on target transcripts with which it does not undergo stoichiometric degradation.
Figure S7  Ratios of promoter activities of RyhB and those of sodB or fumA, as a function of the concentration of DTPA.
Figure S8  Graded monotonic production of YFP induced from an arabinose promoter.
Figure S9  Effects of arabinose-induced RyhB production on the mean sodB transcript number.
Figure S10  Effects of arabinose-induced RyhB production versus iron deprivation on sodB transcript noise.
Figure S11  Effect of an RNase E mutant strain in which co-degradation is blocked on RyhB production.
Figure S12  Average histograms of RyhB fluorescence density for three DTPA concentrations.
Figure S13  Half-life of sodB transcripts for two DTPA concentrations obtained from qRT-PCR experiments.
Figure S14  Effects of iron deprivation on cell growth.

Table S1  Strains used in the study.
Table S2  Sequences of smFISH probes, primers and probes used for qRT-PCR.

Supplemental Materials and Methods

Supplemental References
Figure S1  **Discrete nature of the intensity of spatially separable fluorescence spots.** Histogram of spot fluorescence intensity levels in smFISH images of sodB transcripts, at 0 µM DTPA. The histogram has been decomposed into a sum (red curve) of Gaussian contributions (black lines), each corresponding to a different discrete number of transcripts. The position of the peak corresponding to 1 mRNA was determined from a histogram obtained at 75 µM DTPA.
Figure S2  Effects of different mutant strains on the stoichiometric degradation of sodB and fumA transcripts with iron deprivation. Mean transcript numbers of sodB (blue) and fumA (red) in individual cells as function of the concentration of DTPA measured by smFISH, following procedures as described in Materials and Methods. (A) Measurements in a ∆ryhB mutant strain (IRE-106). The transcript numbers are nearly independent of iron deprivation. For the sake of comparison, the corresponding data measured in the wild type strain (shown in Figure 2) is also plotted (empty circles). (B) Measurements in a ∆hfq mutant strain (HL1188). The growth is severely affected in this strain. (C) Measurements in an RNase E mutant strain (TM528, rne701-Flag-cat, full circles) and in the background strain (TM338, rne-Flag-cat, empty circles). Activity in the RNase E mutant strain is not fully abrogated under iron deprivation. (D) Measurements in an RNase III mutant strain (NC499), show behavior similar to the wild type strain. The gradual reduction observed in the mean transcript number of sodB in all panels is due in part to a known reduction in the promoter activity of sodB with iron deprivation (1). Error bars denote standard errors over three independent experimental repeats.
Figure S3  Switch-like effects of iron deprivation on mean sodB levels obtained from qRT-PCR experiments. Transcript sodB levels as a function of DTPA, normalized by the value without DTPA (red circles). Error bars denote standard errors from two independent experimental runs. Switch-like behavior is observed at ~30 µM DPTA, in agreement with the data obtained from the smFISH experiments of Figure 2A shown for comparison (black circles). Cells were grown in the presence of different concentrations of DTPA and the sodB transcript level was measured using qRT-PCR techniques as outlined in Materials and Methods.
Figure S4  Effects of iron addition to an iron-free medium on the promoter activity of RyhB and one of its targets. *E. coli* cell cultures (MG1655 strain) were grown overnight at 37°C in LB medium, then diluted 1:50 into fresh M9 medium containing L-glucose (0.4%) and casamino acids (0.2%) and allowed to reach an OD_{600} of 0.3-0.4. Cells were then further diluted 1:50 in the same medium, containing various ferric citrate concentrations and allowed to reach an OD_{600} of 0.4-0.6. Left: Promoter activity of RyhB measured with reporter plasmids encoding a *pryhB-yfp* fusion. Right: Mean μ_{sodB} transcript number in individual cells as function of the concentration of iron citrate. Data were obtained from two independent experimental runs. Error bars represent standard errors. The black line through the data, represents a fit with a Hill function \( y = a + (b - a)(x^n/(k^n + x^n)) \) with \( a = 1.8 \pm 0.2, b = 5.2 \pm 0.4, k = 1.8 \pm 0.1, n = 6 \pm 2 \). The value of the Hill coefficient 6±2 in the present plot, albeit smaller than when adding DTPA to LB, is consistent with a high degree of cooperativity.
Figure S5 Switch-like behavior of mean sufC transcript levels as function of iron deprivation. Mean sufC transcript levels as a function of DTPA concentration, measured using smFISH. Note that sufC is under direct transcriptional repression by Fur but its transcripts are not RyhB targets. The data represent an average over three experimental runs carried out under the same conditions, while error bars denote standard errors. The black line through the data is a fit with a Hill function ($y = a + (b - a)(x^n / (k^n + x^n))$) with $a = 0.5 \pm 0.3$, $b = 5 \pm 1$, $k = 38 \pm 9$, and a Hill coefficient $n = 3 \pm 1$.)
Figure S6  Effects of iron deprivation on target transcripts with which it does not undergo stoichiometric degradation. Left: mean fepB transcript number in individual cells as function of the concentration of DTPA. Binding of RyhB to fepB transcripts blocks the latter’s Shine Dalgarno sequence, thereby inhibiting translation. Data were obtained from four independent experiments. Right: mean shiA transcript number in individual cells as function of the concentration of DTPA. Binding of RyhB to shiA transcripts activates translation by altering transcript secondary structure, exposing their Shine Dalgarno sequence. Data were obtained from four independent experimental runs. Error bars in both panels represent standard errors.
Figure S7  Ratios of promoter activities of RyhB and those of sodB or fumA, as a function of the concentration of DTPA. E. coli cell cultures (MG1655 strain) bearing plasmids with the promoter fusions of either ryhB, sodB or fumA to YFP were grown overnight at 37°C in LB medium, then diluted 1:50 into fresh LB medium and allowed to reach an OD600 of 0.3-0.4. Cells were then further diluted 1:50 in the same medium for 3 hours. The values of the promoter activities of the three genes \( \alpha_{RyhB} \), and \( \alpha_{m} = \alpha_{sodB} \) or \( \alpha_{fumA} \) were measured simultaneously in a plate reader as a function of DTPA concentration (see Materials and Methods). Shown are the ratios \( \alpha_{RyhB}/\alpha_{m} \) of either sodB (empty circles) or fumA (full circles). Data were obtained from two independent experimental runs. Error bars represent standard errors. Inset: blowup of the small DTPA portion of the data, illustrating that the value at which the rate of production of RyhB is equal to the rates of production of sodB and fumA is accessible in our experiments (dashed line).
Figure S8  Graded monotonic production of YFP induced from an arabinose promoter. 
*E. coli* cell cultures (strain IRE-111) were grown overnight at 37°C in LB medium, then 
diluted 1:100 into fresh LB and allowed to reach an OD_{600} of 0.3-0.4. Cells were then 
further diluted 1:40 in LB containing various arabinose concentrations for 3 hours and 
then diluted 1:5 in M9 medium with the same arabinose concentration in a 96-well plate. 
Plates were maintained at 37°C with shaking throughout the YFP fluorescence and optical 
density (OD_{600}) measurements. The mean fluorescence was obtained over six 
independent replicates and the error bars represent standard deviations.
**Figure S9** Effects of arabinose-induced RyhB production on the mean sodB transcript number. Mean sodB transcript number in individual cells as function of the concentration of arabinose measured using smFISH in a strain in which RyhB is produced from an arabinose-induced promoter (IRE-109). The fluorescence represents the mean over three independent experiments and the error bars represent the standard error.
Figure S10  Effects of arabinose-induced RyhB production versus iron deprivation on sodB transcript noise. Comparison between the noise levels of sodB mRNA as function of the mean sodB transcript number $\mu_{\text{sodB}}$ measured in a wild-type strain in which RyhB is produced by iron depletion (empty circles), and in a strain (IRE-109) in which RyhB is produced from an arabinose-inducible promoter (full circles). The latter data was obtained from three independent experiments, while the data from the wild type strain is the same as in Figure 3A. The noise levels in both strains coincide within experimental error.
Figure S11  Effect of an RNase E mutant strain in which co-degradation is blocked on RyhB production. Typical snapshots of fluorescence from undegraded intracellular RyhB, tiled with labeled oligonucleotides. Panel A: background strain (TM338). Panel B: in an RNase E mutant strain (TM528). Both images were taken at iron-poor conditions (75 μM DTPA).
Figure S12 Average histograms of RyhB fluorescence density for three DTPA concentrations. Fluorescence density is defined as fluorescence measured in individual cells, normalized by their area. Cells were grown, treated with DTPA and RyhB was labeled as described in Materials and Methods (main text). The average histograms correspond to RyhB distributions measured at DTPA concentrations below (top), above (bottom) and within the switching region (middle) (see Figure 4B). The average histograms represent bin-by-bin means of histograms measured in three independent experimental runs, while errors bars represent standard errors. Prior to calculating histograms within each run, the average background fluorescence measured in a ΔryhB strain was subtracted from the fluorescence in the wild-type strain.
Figure S13  Half-life of sodB transcripts for two DTPA concentrations obtained from qRT-PCR experiments. Relative level of sodB transcripts as a function of time after addition of rifampicin, measured using qRT-PCR for 0 µM and 75 µM DTPA. Two independent repeats were performed in each case (red and blue circles). The corresponding red and blue lines represent linear fits to the data, the inverse of the slopes yielding the corresponding half-life. The shown values of the half-life $t_{1/2}$ represent an average over the corresponding two experiments in the plot while errors represent standard errors.
Figure S14  Effects of iron deprivation on cell growth. Mean doubling time of cells as function of DTPA concentration. *E. coli* cell cultures (MG1655 strain) were grown overnight at 37°C in LB medium, then diluted 1:100 into fresh LB and allowed to reach an OD$_{600}$ of 0.3-0.4. Cells were then further diluted 1:80 in LB containing various DTPA concentrations under the same growth conditions at 37°C as in the smFISH experiments and after 2.5 hours the optical density (OD$_{600}$) was measured as a function of time in a 96-well plate. Plates were maintained at 37°C with shaking throughout the experiments. The data represent an average over two experimental runs carried out under the same conditions. Error bars denote standard errors. Each experiment was repeated at least twice on different days.
### Supplemental Tables

#### Table S1 Strains used in the study

| Strain Name | Strain #  | Mutations                  | Reference |
|-------------|-----------|----------------------------|-----------|
| WT-Keio     | BW25113   | sodB734(del)::kan          | (2)       |
| ΔsodB       | JW1648    | sodB734(del)::kan          | (2)       |
| ΔfumA       | JW1604    | fumA760(del)::kan          | (2)       |
| ΔsufS       | JW1670    | sufS757(del)::kan          | (2)       |
| ΔshiA       | JW1962    | shia755(del)::kan          | (2)       |
| ΔfepB       | JW0584    | fepB730(del)::kan          | (2)       |
| WT          | MG1655    |                            | (3)       |
| ΔryhB       | EM1238    | ryhB::cat                  | (4)       |
| ΔryhB       | IRE-106   | ΔaraFGH ΔaraE ryhB<>cat    | New strain|
| ΔaraFGH ΔaraE ryhB<>cat | IRE-109 | PBAD–ryhB                  | New strain|
| PBAD–ryhB   | IRE-110   | ΔaraFGH ΔaraE ryhB<>cat PBAD–ryhB | New strain|
| PBAD–yfp    | IRE-111   | ΔaraFGH ΔaraE ryhB<>cat PBAD–yfp | New strain|
| rne-Flag-cat| TM338     | rne-Flag-cat,              | (5)       |
| rne701-Flag-cat| TM528 | rne701-Flag-cat            | (5)       |
| rnc14       | NC499     | rnc14::miniTn10            | New strain|
| Δhfq        | HL1188    | hfq::cat                   | New strain|
| Gene | ryhB | sodB | sufC |
|------|------|------|------|
| Probe | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye |
|      | Or |      |      |
|      | Stellaris® FISH Probes, Custom Assay with Quasar® 670 Dye |      |      |
| Probe sequences (5’ to 3’) | CTTTCAGGTTTCTCCGCGAGG AGCAATGTGAGCAATGTCG CCGCTGGCTAAGTAATACT | GTGACAGGTAATTCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCTGACCGFGTAAATCTTTA CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
|      | TGCAGGTTATCCGAATGAC AGAGCATCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | GACCAGTCTTTAGCATATGG TTCCGCAGAAATGTGCGGTG TGCCGTAGTGATCTGCTAG | CGCAGGATAGCTTTATCTTC GAAGGACCTCGAGGGCTTAA |
| Gene | fumA | fepB | shiA |
|------|------|------|------|
| **Probe** | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye | Stellaris® FISH Probes, Custom Assay with CAL Fluor® Red 590 Dye |
| **Probe sequences** | GCTGGTTAGCAGGTAATACT ATTCAAGATACGCTATGCTGGT CCAACAGTTAACGCTTTCCG AATGAAATGACGATGGAAACA TTTTCTACATTTTGCTGGA TACTAGCGCGACTGGAATCAC ACCACCAACTTTCGCTGTA AATAGTGGGGAAGGCTAT TACCAAGGTAAGGCAATTTA | CCTGGTAAATAGAAGGGGCTG AACTCGCGCATATTCTCGT ACGGCTGTCAGTAACTCG ATGCTGCCTTCCAGTGTATG CTGCTGGTGAACATACGTTG ATGCTGCCTTCCAGTGTATG ATGCTGCCTTCCAGTGTATG | ATCGGAGCGAGTGAGATGA CCGGCGGCACTTTACGAAAG ATCATACAGTCGACGACGG CGGCTGTCAGTAACTCG ATGCTGCCTTCCAGTGTATG ATGCTGCCTTCCAGTGTATG ATGCTGCCTTCCAGTGTATG |
## Primers and probes used for qRT-PCR

| Oligo name   | Sequence                                      |
|--------------|-----------------------------------------------|
| *sodB* F′ primer | ACC GCG TTT GAA GGT AAA TCA                   |
| *sodB* R′ primer | GAC CTG AGC TGC GTT GTT GA                     |
| *rpsL* F′ primer | AAG CGT CCG TTC TAC CAG GTT                   |
| *rpsL* R′ primer | AAC CAA CGC GCT CGA TGA                       |
| *sodB* probe  | 6-FAM-TTA TTC GCA GCT CTG AAG G-MGB           |
| *rpsL* probe  | 6-FAM-TTG TCG CTG ACA GCC G-TAMRA             |
**Supplemental Materials and Methods**

Expression of the arabinose *pBAD* operon following induction with limiting levels of arabinose results in some cells being fully induced and others remaining un-induced for *araBAD* operon transcription. Induction by arabinose is autocatalytic and the first cells to take up some arabinose are induced to express transport functions AraFGH and AraE allowing the uptake of more arabinose (6). This all-or-nothing induction of the *pBAD* promoter is prevented by using a strain deficient in both *araE* *araFGH* as well as the catabolic functions (*araBAD*). Morgan-Kiss *et al.* demonstrated that in such an *ara* mutant strain, a particular LacY protein mutant could perform facilitated diffusion of arabinose resulting in homogeneous and graded expression of the *ara* promoter over extended incubation times at sub-saturating levels of arabinose (7).

**Deletion of the arabinose transport genes *araFGH* and *araE*.** The *araFGH* cluster was deleted by a two-step selection/counter-selection process using recombineering methods (8). We started with strain IRE101, which is MG1655 with the pSIM18 hygromycin resistant plasmid used to express the Red recombineering functions (9). First, the *tetA-sacB* cassette was inserted by recombineering methods into IRE101 to replace *araFGH* using selection for tetracycline resistance conferred by the *tetA* gene and sensitivity to growth on LB sucrose conferred by the *sacB* gene, thereby generating the strain IRE102. Next, recombination with oligo RY17 and selection for sucrose resistance, created the clean deletion of the *araFGH* cluster from IRE102 and removal of the *tetA-sacB* cassette. RY17 contains the sequence 5’GTG GGA AAA AAC GCT AAA TTG TTG CAG AAA AAA GCA / TTG TCT TTG GTA CCC ATG CGG GAT GTC TTC TTT TTA 3’ where the / defines the deletion endpoints to each side of the *araFGH* segment in the new strain IRE103.

The *araE* gene was deleted from IRE103 in a similar two-step selection/counter-selection process using recombineering methods (8). First, the *tetA-sacB* cassette was inserted to replace *araE* using selection for tetracycline resistance conferred by the *tetA* gene. Recombinants were tested for sensitivity to growth on sucrose conferred by the *sacB* gene, thereby generating the strain IRE104. Next, recombination with oligo RY12 and selection for sucrose resistance, created a clean deletion of the *araE* gene from IRE103 and removal of the *tetA-sacB* cassette. RY12 contains the sequence 5’AAT GTT CAG CGC AGT GTA GAG CCA GAA CGT ACC GGC / TTC AAA TTA AGT TGA ATT ATT GAG ATT ATT ATT AAC 3’ where the / defines the deletion endpoints to each side of the *araE* gene in the new strain IRE105.

**Deletion of the *ryhB* gene to generate strain IRE106.** In the strain IRE105 (MG1655 Δ*araFGH ΔaraE /pSIM18), the native *ryhB* gene was replaced with a *cat* cassette (*ryhB<>cat*) conferring chloramphenicol resistance by the process of recombineering. The *cat* cassette was amplified by PCR with primers carrying 50 nucleotide long regions at their 5’ ends homologous to each flank of the *ryhB* gene. RY18 and RY19 are the two oligo primers used to generate the *cat* cassette with 50bp flanking homologies. RY18: 5’ TTT
GCA AAA AGT GTT GGA CAA GTG CGA ATG AGA ATG ATT ATT ATT GTC TCT GTG ACG GAA GAT CAC TTC G and RY19: 5' TAA CGA ACA CAA GCA CTC CCG TGG ATA AAT TGA GAA CGA AAG ATC AAA AAA CCA GCA ATA GAC ATA AGC G where each underlined sequence is the primer segment for cat cassette amplification. The region upstream of RY18 and RY19 is the homologous segment upstream and downstream respectively of the rhyB gene.

**Insertion of rhyB under araBAD promoter control.** Starting with the strain IRE106 (MG1655 ΔaraFGH ΔaraE rhyB<cat strain carrying pSIM18), we inserted the rhyB gene under transcriptional control of the araBAD promoter in a precise way so that the native rhyB 5’ RNA end is in the position of the native pBAD 5’ RNA start and at the same time removed the entire pBAD RNA coding region for the BAD genes and replacement with the rhyB gene. First, phage P1 was grown on the strain XTL298 that carries the tetA-sacB cassette in place of the araBAD genes with the amp gene also present just beyond sacB as described previously (8). This cassette was transduced by P1 into IRE106 so as to generate IRE107 (MG1655 ΔaraFGH ΔaraE rhyB<cat araBAD< tetA-sacB amp). Next the rhyB gene was amplified to encode from the first nucleotide of RyhB RNA through the 3’ end of the RNA by PCR using primers carrying 50 nucleotide long regions at their 5’ ends. One homologous to the regions just upstream of the ara RNA start and the second homologous to the DNA beyond the tetA-sacB counter-selection cassette. RY21 and RY20 are the two oligo primers used to amplify rhyB with their respective flanking homologies. RY21: 5’ AGC GGA TCC TAC CTG ACG CTT TTT ATC GCA ACT CTC TAC TGT TTC TCC ATG CGA TCA GGA AGA CCC TCG C and RY20: 5’ GCT TGA GTA TAG CCT GGT TTC GTT TGA TTG GCT GTG GTT TTA TAC AGT CAG ATT TCG TCC TTT TTA AGG TGG TTA TTT ACA C where the underlined sequence corresponds to sequences priming amplification of the rhyB gene. This rhyB DNA cassette with flanking homologies was used to replace the tetA-sacB counter-selection cassette and fuse RyhB expression to the pBAD promoter by recombineering and selection of sucrose resistant recombinants. The final strain IRE108 is MG1655 ΔaraFGH ΔaraE rhyB<cat P_BAD – rhyB in which the pSIM18 plasmid (whose replication is temperature sensitive) has been eliminated by continued growth at 42 degrees (9).

**LacY permease mutant expressed from the plasmid placY*A177C.** Plasmid placY*A177C constitutively expresses a permease capable of facilitating diffusion of arabinose into cells at sub-saturating levels (5). This plasmid was transformed into the IRE108 strain (MG1655 ΔaraFGH ΔaraE rhyB<cat P_BAD – rhyB) to generate strain IRE109.

**Insertion of yfp coding segment under the control of the araBAD promoter and the translation initiation codon of araB.** The strain IRE110 was made and used to generate a strain with yfp under control of pBAD. Strain IRE105 (MG1655 ΔaraFGH ΔaraE /pSIM18) was transduced by P1 to insert araBAD< tetA-sacB amp to be used for counterselection and replacement with yfp. The yfp gene was amplified from its ATG start codon through the stop codon using primer RY28 and RY29 that carry 50 nucleotide long regions at the primers 5’ ends. RY28 has homology to the region just upstream of the araB start codon, and RY29 has homology to the DNA beyond the tetA-sacB amp cassette. Recombination with these PCR products deletes the araBAD operon and precisely replaces all their open
reading frames with the open reading frames of \( yfp \) beginning at the ATG of \( araB \). Note that \( yfp \) and \( cfp \) are identical in sequence at the 5’ and 3’ ends of their coding sequences, and thus the same primers can be used for both genes. The \( Yfp \) expressing strain is IRE111 and is MG1655 \( \Delta araFGH \Delta araE \) ryhB<cat \( P_{BAD} - yfp \). \( RY28 \): 5’ ACT CTC TAC TGT TTC TCC ATA CCC GTT TTT TTG GAT AGG AGG ATG AAA CGA TGC GTA AAG GAG AAG AAC TTT TC, and \( RY29 \): 5’ GCT TGA GTA TAG CCT GGT TTC GTT TGA TTG GCT GTG GTT TTA TAC AGT CAT TAT TTG TAT AGT TCA TCC ATG CC where the underlined sequence corresponds to sequences priming amplification of the \( yfp \) gene. The pSIM18 plasmid whose replication is temperature sensitive was eliminated from both by continued growth at 42 degrees (9). The Ampicillin resistant plasmid placY*A177C, which constitutively expresses the LacY permease capable of facilitating diffusion of arabinose into cells at sub-saturating extracellular levels (7), was transformed into and is present in final IRE113 and IRE114 strains.

**RNAase III mutant \( rnc14 \) in MG1655.**
An RNAase III mutant containing a mini-Tn10 insertion in the \( rnc \) gene (\( rnc14 \)) has been described (10). The \( rnc14 \) mutation was moved by P1 transduction (12) to MG1655 and designated NC499. NC499 has the same phenotype as other \( rnc \) mutants (10).

**Hfq deletion mutant in MG1655.**
HL1188, from the Gottesman lab (S. Gottesman and H. Li) was made by moving \( hfq::cat \) into MG1655 by P1 transduction, and selecting for CmR from a strain TX3077 referenced in [http://www.ncbi.nlm.nih.gov/pubmed/9393714](http://www.ncbi.nlm.nih.gov/pubmed/9393714).

**Recombineering:** Recombineering methods and protocols performed are described (see: [http://redrecombineering.ncifcrf.gov/](http://redrecombineering.ncifcrf.gov/)) (11, 13). In strain construction of gene replacements by drug markers, there was a two-hour outgrowth, and then the cells were plated on appropriate antibiotic plates (30 \( \mu \)g/ml ampicillin, 10\( \mu \)g/ml chloramphenicol, 12.5 \( \mu \)g/ml tetracycline, 50\( \mu \)g/ml kanamycin).

**Sequencing and Analysis:** Final gene constructions made by recombineering protocols were sequenced. Sequencing was done by Leidos Biomedical Research, Inc. Sequencing results were analyzed with Sequencher version 4.8.

**Oligos and bacterial strains:** Gene replacements were made by dsDNA recombineering (13); construction details including final construct sequence will be supplied upon request. Multiple mutants were introduced by P1 transduction (14). Oligos were purchased from Integrated DNA Technologies (IDT) and were desalted but not further purified.
Single-molecule fluorescence *in situ* hybridization (smFISH): We followed already published procedures (15).

**Probe design and labeling:** DNA oligonucleotide probes were designed using the Stellaris RNA probe FISH designer (https://secure.biosearchtech.com/stellarisdesigner/). The Stellaris™ FISH probes were tagged with CAL Fluor® Red 590 dye or with Quasar® 670 dye. Sequences of smFISH probes are given in the Supplementary Table S2. The probes were dissolved in 10mM Tris-HCl 1mM EDTA (pH 8.0) to create a probe stock at a total oligo concentration of 25 μM. The tube was wrapped in aluminum foil and stored at -20 °C.

**Sample fixation and permeabilization:** Overnight cultures of the indicated *E. coli* strain deletions from the KEIO collection (2) and an MG1655 *E. coli* strain, were diluted 1:100 in 4 ml of LB medium. The culture was incubated at 37 °C with shaking. When OD$_{600}$ of the culture reached 0.3–0.4, the cells were pelleted by centrifugation (5 minutes, 4500×g, 4°C). The supernatant was removed and the cells were resuspended in 1 ml 1× PBS (Diethylpyrocarbonate (DEPC)-treated) and 4 ml freshly prepared 4.6% formaldehyde. The cells were then mixed on a rotator at room temperature for 30 minutes. The cells were pelleted by centrifugation (10 minutes, 1000×g, 4°C). The supernatant was removed and the cells were washed in 1 ml 1× PBS twice (i.e. re-suspended in 1 ml 1× PBS, centrifuged at 1000×g for 7 minutes, 4°C, and supernatant removed). The cells were resuspended in 300 μl (DEPC)-treated water, and then 350 μl of 100% ethanol was added and mixed twice to get to a final concentration of 70% ethanol. The cells were left at room temperature with mixing on a rotator for at least 1 hour (or alternatively, at 4 °C for at least an overnight) to permeabilize the cell membrane.

**Hybridization:** After permeabilization, cells were centrifuged (7 minutes, 750×g) and the supernatant was removed. The cells were resuspended in 1 ml of 20% wash solution (0.2 g of formamide in 1 ml of 2×SSC (DEPC treated) and the tube was left standing for a few minutes. An aliquot of 20% hybridization solution (10 ml of 20% hybridization solution filter sterilized which contains 1 g of dextran sulfate, 2 g of formamide, 10 mg of *E. coli* tRNA, 1 ml of 20× SSC, 40 μl of 50 mg/ml BSA and 100 μl of 200 mM ribonucleoside vanadyl complex) was warmed to room temperature and 50 μl was added to a microcentrifuge tube. The indicated Fish probes were added at final concentration of 250 μM to the hybridization solution and mixed well. The cells were then centrifuged (7 minutes, 600×g) and the supernatant was removed. The cells were resuspended in the hybridization solution with probes and left at 30 °C overnight. Hybridized samples could be stored at 4 °C.

**Washing:** 10 μl of hybridized sample was transferred to a microcentrifuge tube. The rest was stored at 4 °C. 1ml of 20% wash solution was added to the tube and mixed well. Cells were pelleted by centrifugation (7 minutes, 750×g) and the supernatant was removed. The cells were washed 2 more times in 1ml of 20% wash solution, centrifuged at 750×g for 7 minutes, and supernatant removed). The cells were resuspended in 25 μl of 2× SSC and imaged under the microscope.
**Transcript visualization**: An agarose gel pad (1.5%) in SSCx2 was made on a coverslip and 5 μl of sample was pipetted onto the pad and a #0 mm coverslip was placed on top of the agarose gel pad. The sample was imaged on a Nikon Eclipse –Ti-E microscope controlled by the Niss elements software using a 100x N.A 1.45 oil immersion phase contrast objective lens (Nikon plan-apochromat 100x 1.45 lambda) and an Andor iXon X3 EMCCD camera. All the filters are from Chroma. The filters used were ET-534/30x for excitation, 560lp as dichroic mirror and ET-585/40M for the emission. A phase contrast image was acquired followed by a z-stack of 13 slices and 250 nm spacing of fluorescent images with 2s integration time of each slice. Each sample was imaged at multiple locations to get a total of at least 500 cells.

**Spot recognition**: A spot recognition program developed in our lab, based on a version of the IDL ParticleTracking software (www.physics.emory.edu/~weeks/idl/) (16) adapted to Matlab by D. Blair and E. Dufresne (http://physics.georgetown.edu/matlab), and incorporated into our custom Matlab program (2), was used to automatically identify and quantify localized fluorescence signals. A Gaussian filter was first applied to smooth out noise, and spots were recognized by the presence of a local maximum in both x- and y-directions. This was done at each z-position in the stack of images, and each spot was quantified at the z-position where it had the highest fluorescence intensity (where the spot is in focus).

**Estimating mRNA numbers**: A fluorescence spot could consist of multiple mRNAs in close proximity. The integrated intensity arising from a single mRNA needed to be estimated for each smFISH experiment so that fluorescence intensities could be normalized to give the absolute number of mRNAs. The typical intensity of “false positives” in an experiment was first estimated from the histogram of individual spot intensities of a negative control (e.g.: E. coli strains ΔsodB or ΔryhB). Histograms of individual spot intensities from relatively low expression samples (exposed to 125 μM DTPA) were then examined. Because most spots in these samples were expected to contain a single mRNA, the first peak in the intensity histogram that emerged above the false positive range served as an estimate for the intensity of a single mRNA. The sum of intensities of all spots in each cell was then normalized by the single mRNA intensity to give the absolute number of mRNAs.

**Quantitative reverse transcription polymerase chain reaction (qRT-PCR) RNA lifetime measurement**: RNA lifetime measurements were carried out following the previously published methods (15). Cells were diluted from overnight cultures and grown in a 25 ml volume, under the same growth conditions with or without DTPA as in smFISH experiments. When OD_{600} reached ~0.3, the culture was separated into two halves, each half being transferred to a new culture flask, and grown at the same conditions for 10-15 minutes. 1.5 ml of culture was then extracted from each flask and mixed with 3 ml of Qiagen Rnaprotect Bacterial reagent to stabilize cellular RNA (see Total RNA extraction below for details). These samples were treated as t = 0 samples. Rifampicin was at t=0 added to one of the remaining cultures, at a final concentration of 500 μg/ml, to inhibit
transcription. The culture without rifampicin served as a control. 1.5 ml of culture was extracted from each flask at 0, 7.5, 15, 22.5 and 30 min for 0 µM DTPA and at 0, 2, 4, 8, 10 min for 75 µM DTPA after addition of rifampicin and mixed with 3 ml of Qiagen RNAprotect Bacterial reagent to stabilize cellular RNA. Subsequent total RNA extraction and qRT-PCR were performed as described previously in this section. Target RNA level of the rifampicin-treated sample as a function of time was fitted to an exponential, \( \ln y = Ax+B \) and RNA lifetime (in minutes) estimated as \( \tau_{\text{RNA}} = -1/A \).

**Total RNA extraction:** Two volumes of RNAprotect Bacteria Reagent (Qiagen) were added to bacterial cell cultures that reached OD_{600} = 0.3-0.4. Total RNA was then extracted using the RNeasy Mini Kit with the optional DNase I digestion step (RNase-free DNase Set). The eluted from the final step was passed through the spin column one additional time to increase total RNA concentration. 2 µl of each total RNA sample diluted 10-fold in 10 mM Tris·Cl, pH 7.5 was measured on a Nanodrop. The purity and concentration of the total RNA samples were estimated from the absorbance spectrum between 220 and 320 nm. The samples were treated with DNasel for 30 minutes at 30 °C (using a DNA-free: DNase treatment & removal kit).

**Reverse transcription:** The RNA was then reverse transcribed to get the corresponding cDNA. Reverse transcription was performed immediately after the total RNA extraction. The cDNA was synthesized using High Capacity cDNA Reverse transcription kit following the manufacturer’s protocols. The optional RNase inhibitor was used in the reaction. The 2X RT master mix was prepared by mixing 2.0 µL of 10X RT buffer, 0.8 µL of 25 X dNTP mix (100 mM), 2.0 µL of 10X random primers, 1.0 mL of MultiScribe™ Reverse Transcriptase, 1.0 µL of RNase Inhibitor and 3.2 µL of RNAase-DNAse free PCR water to make a final volume of 10 µL total reaction. The thermal reaction was performed as follows: 25 °C for 10 minutes; 37 °C for 120 minutes; 85 °C for 5 minutes; 4 °C for storage. The cDNA samples were then diluted to a final concentration corresponding to a RNA amount of 20 ng. The cDNA samples were stored at -20 °C until further use.

**qRT-PCR conditions:** Amplification and detection of DNA by real-time PCR were performed with the ABI-PRISM 7700 Sequence Detection System (Applied Biosystems) using optical grade 96-well plates. Negative control of ΔsodB and RNAse DNase free PCR water were used for amplification of sodB transcript and water was used as control in order to ensure the reactions were devoid of any cDNA contamination. The PCR reaction was performed in a total volume of 10 µl using: 2 µL of cDNA sample corresponding to 20 ng RNA, 5 µL of TaqMan® Fast Universal PCR Master Mix 2x, 0.5 µL TaqMan® probe, 0.5 µL of 10 µM forward primer, 0.5 µL of 10 µM reverse primer, 1.5 µL of RNAase-DNAse free PCR water. Sequences of probes and primers used for qPCR are listed in Table S2. The reactions were incubated in a plate at 95°C for 20s, followed by 40 cycles of 95°C for 1s and 60°C for 20s. All reactions were run in duplicate samples. Data analysis made use of Sequence Detection Software version supplied by Applied Biosystems. The relative level of sodB transcript was calculated using a comparative Ct method (the threshold cycle (CT) is defined as the fractional cycle number at which the fluorescence passes the fixed
threshold.) and the housekeeping \textit{rpsL} gene which encodes the 30S ribosomal protein S12 was used as an endogenous control.
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