

Transcript degradation and noise of small RNA-controlled genes in a switch activated network in *Escherichia coli*

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ABSTRACT

Post-transcriptional regulatory processes may change transcript levels and affect cell-to-cell variability or noise. We study small-RNA downregulation to elucidate its effects on noise in the iron homeostasis network of *Escherichia coli*. In this network, the small-RNA RyhB undergoes stoichiometric degradation with the transcripts of target genes in response to iron stress. Using single-molecule fluorescence *in situ* hybridization, we measured transcript numbers of the RyhB-regulated genes *sodB* and *fumA* in individual cells as a function of iron deprivation. We observed a monotonic increase of noise with iron stress but no evidence of theoretically predicted, enhanced stoichiometric fluctuations in transcript numbers, nor of bistable behavior in transcript distributions. Direct detection of RyhB in individual cells shows that its noise is much smaller than that of these two targets, when RyhB production is significant. A generalized two-state model of bursty transcription that neglects RyhB fluctuations describes quantitatively the dependence of noise and transcript distributions on iron deprivation, enabling extraction of *in vivo* RyhB-mediated transcript degradation rates. The transcripts' threshold-linear behavior indicates that the effective *in vivo* interaction strength between RyhB and its two target transcripts is comparable. Strikingly, the bacterial cell response exhibits Fur-dependent, switch-like activation instead of a graded response to iron deprivation.

INTRODUCTION

The remarkable capability of bacteria to adapt to variations in the external environment is achieved in large part through changes in gene expression mediated by small RNAs (sRNAs) (1–3). sRNAs play major roles in diverse cellular responses to environmental cues and stresses including iron deprivation, quorum sensing, oxidative stress, carbon metabolism and more (2,4). Most sRNAs downregulate the expression of their target genes by forming imperfect duplexes with target transcripts, through mediation by the Hfq chaperone protein. sRNA–mRNA complexes are then degraded by the RNase E degradosome, or transcript translation is inhibited by either the occlusion of the ribosome binding site or of a translational enhancer (5). In both cases, regulation is stoichiometric in character, in contrast with the catalytic regulation of gene expression by transcription factors (6).

Stoichiometric degradation gives rise to unique features such as a threshold-linear switching behavior between an efficiently silenced regime and an active regime. Silencing occurs when the rate of production of the sRNA exceeds that of its target mRNA, generating low levels of mRNA and protein. The active regime occurs when the production rate of the sRNA is smaller than that of the mRNA, leading to a linear increase of target mRNA levels (7,8). Stoichiometric degradation has also been predicted by theoretical models to give rise to an enhancement of intrinsic noise of transcripts and proteins of sRNA target genes, at the crossover between the silenced and active regimes, i.e. when the production rates of the sRNA and an mRNA target are comparable (7,9,10). Intrinsic noise is the cell-to-cell variation in gene expression of genetically identical cells (11–16), stemming from the stochastic nature of biochemical processes such as transcription and translation (17). Enhanced stoi-

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chiometric fluctuations are different from those due to small numbers, which are expected to dominate noise as sRNA production increases deeper into the silenced regime.

The iron homeostasis network of *E. coli*, which includes the sRNA RyhB (18), is an attractive model system to characterize sRNA regulation and test theoretical predictions. This network is controlled by the master ferric uptake regulator (Fur), a transcription factor that senses the intracellular concentration of free iron. In its iron-bound form, Fur represses the expression of many genes involved in iron acquisition, storage, oxidative stress and more. When iron is limiting, Fur becomes iron-free, relieving the repression of iron homeostasis genes. A specific target of Fur repression is *ryhB*, encoding an sRNA that coordinately inhibits the production of some twenty iron-using proteins (4), by targeting the degradation of their mRNAs (19,20). RyhB was reported to inhibit Fur production in a mixed double-negative feedback loop configuration (21), suggesting the possibility of bistable behavior similar to that found in double-negative feedback motifs involving two transcription factors (22,23). Bistability in a mixed double-negative feedback loop involving a sRNA and a transcription factor has been observed in a stochastic numerical analysis, but only in a limited region of parameter space (24).

Recently, the effects of sRNA regulation on protein noise have been studied within the framework of the iron homeostasis network of *E. coli* (25). It was found that the total protein noise from two RyhB targets is robust over a wide range of RyhB production rates, despite large reductions in the mean concentrations of both proteins. Furthermore, it was shown that this robustness is due to the dominance of extrinsic noise sources, e.g. fluctuations in the number of global factors such as RNA polymerases, ribosomes and metabolites, which affect the expression of all genes in a cell equally (14,15,26). It was suggested that extrinsic noise dominated protein fluctuations, and prevented the observation of intrinsic enhanced fluctuations in the expression of target gene proteins. No evidence for bistable behavior was observed in the distributions of protein numbers of the two RyhB target genes studied (25).

Here, we report the results of a comprehensive study of the effects of regulation by an sRNA on mRNA transcript noise, using the iron homeostasis network of *E. coli* as a model system. The transcript level is appropriate to measure this noise because it responds directly to sRNA regulation, and unlike the protein level, is less prone to be masked by global extrinsic noise (27). We focused on the coupled degradation of RyhB sRNA and its target mRNAs *sodB*, which codes for iron-superoxide dismutase, and *fumA*, which codes for fumarase A, one of three fumarase isozymes participating in the TCA cycle. We measured numbers of individual mRNA molecules in individual cells using single-molecule fluorescence *in situ* hybridization (sm-FISH) (26–28), and looked for evidence of enhanced stoichiometric fluctuations in the noise of target transcripts. By measuring the fluorescence of RyhB tiled with a small amount of labeled oligomers in individual cells, we characterized sRNA noise as well and its dependence on iron levels. Furthermore, we tested whether a two-state model of transcription that has been highly successful in describing the statistics of transcript copy number (27,29–31), can be

generalized by adding RyhB-mediated degradation, to account for the dependence of transcript noise and distributions on iron deprivation. We also tested whether the network exhibits bistability as iron deprivation is varied, and the extent to which threshold-linear behavior is observed in the wild-type. Previous demonstrations of threshold-linear behavior were carried out by controlling RyhB production from an inducible promoter on plasmids (8). As we will show below, our measurements also reveal that far from exhibiting a graded response to changes in iron deprivation by external iron availability, RyhB regulation becomes activated in a switch-like fashion within a very narrow range of iron concentrations.

MATERIALS AND METHODS

Growth media and conditions

Bacterial strains were propagated in LB medium at 37°C overnight from a single colony inoculum. For experiments, cultures were diluted 1:100 into fresh LB medium and grown to OD₆₀₀ of 0.2–0.4 before the indicated treatment. Cells were then diluted 1:100 into LB medium with different concentrations of DTPA or arabinose as indicated, and allowed to grow for 3 h to OD₆₀₀ of 0.4–0.6.

Bacterial strains and plasmids

Strains carrying mutations $\Delta sodB$, $\Delta fumA$, $\Delta sufC$, $\Delta shiA$ and $\Delta fepB$ are part of the Keio Collection, Keio University, Japan, (32) and are in the background of the derivative of MG1655 called BW25113. All other strains used here are derived directly from MG1655. EM1238 carries $\Delta ryhB$ (33); TM338 is *rne-Flag-cat* and TM528 is its mutated derivative *rne701-Flag-cat* (34); IRE109 carries the arabinose-inducible RyhB and IRE111 the arabinose-inducible YFP; NC499 is an RNAase III mutant strain; HL1188 is an Hfq mutant from S. Gottesman (see Supplementary Materials and Methods). More details of all strains are presented in Supplementary Table S1.

Measurements of promoter activity

MG1655 (wild-type) bacterial cells bearing plasmids with a promoter fusion *pryhB*-YFP, *psodB*-YFP and *pfumA*-YFP (25,35) were grown as indicated above. Cell suspensions were then transferred to a plate reader, in order to measure their fluorescence intensity F_t every $\delta = 4$ min, for a period of 1.5 h. The fluorescence intensity was corrected by subtracting the background of samples of wild-type bacterial cells, also measured throughout the same period. For each time point the promoter activity was calculated as $PA_t = (F_{t+\delta} - F_t)/(\delta \cdot OD_t)$. The final promoter activity PA is an average over all the values of the time series PA_t , taking into account only points within the log phase of growth.

Half-life of *sodB* mRNA using quantitative real time PCR (qRT-PCR)

Bacterial cells were grown as indicated above. RNA lifetime measurements were carried out following previously

published methods (27). Details about modifications and the subsequent qRT-PCR are given in the Supplementary Methods.

Single-molecule fluorescence *in situ* hybridization (smFISH)

Bacterial cells were grown and treated as indicated above. Single-molecule fluorescence *in situ* hybridization (smFISH) experiments were performed following previously published procedures with the indicated modifications provided here and in the Supplementary Methods (27). Images were taken on a Nikon Eclipse –Ti-E microscope controlled by the NIS-Elements software using a $100 \times$ N.A 1.45 oil immersion phase contrast objective lens (Nikon plan-apochromat $100 \times 1.45 \lambda$) and an Andor iXon X3 EM-CCD camera. All the filters are from Chroma. The filters used were ET-534/30 \times 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 2 s integration time of each slice. Each sample was imaged at multiple locations to get a total of at least 500 cells.

Direct detection of RyhB

RyhB was measured directly in individual cells by fluorescence *in situ* hybridization (FISH). Experiments were performed as described for smFISH through the fluorescence emitted by the labeled oligomer probes when hybridized to the complementary RyhB transcripts. The short length of RyhB (90 nt long) allows hybridization to only three labeled oligomer probes, preventing counting of fluorescent loci as representing individual molecules as in smFISH. In spite of this, hybridization of RyhB to labeled oligomer probes can be exploited to detect RyhB directly by the mean fluorescence density in cells. Analysis of cell fluorescence was carried out following the same procedures as when analyzing fluorescently-labeled proteins (25).

Image and smFISH data analysis

Spot recognition was performed as described in the Supplementary Methods. Cell recognition is based on the MicrobeTracker suite (36), which was integrated into our custom Matlab program. To quantify the fluorescence intensity of each mRNA spot and correct for variation in cell labeling the following procedure was used: the average intensity from a diffraction-limited spot was measured. We then subtracted from the intensity of each spot in a bacterium the median of the intensity values of the pixels in this bacterium which do not contain a fluorescent spot. The variation of the mRNA numbers between experiments is $\sim 20\%$ which is comparable to the errors in So *et al* (27). The discrete nature of the intensity of spatially separable fluorescence spots in our experiments is illustrated in Supplementary Figure S1. To estimate the amount of false positive transcript counts, we analyzed images of *sodB* and *fumA* transcripts in Δ *sodB* and Δ *fumA* strains respectively, using the same parameter values as in the analysis of images in the wild type strain. We obtained 0.04 ± 0.02 *sodB* transcript molecules and 0.05 ± 0.03 *fumA* transcript molecules, values which are ~ 25 and ~ 10 times smaller than the respective

smallest mean transcript numbers obtained in the wild type strain at large DTPA concentrations. A similar procedure was carried out in the case of *shiA* and *fepB* transcripts. While it is customary to normalize smFISH data to a single gene copy (27,37), we have refrained from doing so with our data, since both *sodB* and *fumA*, at 37.36 and 36.31 centisomes, respectively, are quite close to the terminus region at 34 centisomes, and therefore their copy number is not expected to vary much with growth rate (38). In contrast, the *ryhB* gene, at 77.14 centisomes, is much closer to the *oriC* region (84.57 centisomes), and therefore variations in the number of copies with growth rate contribute significantly to the levels of RyhB.

Generalized two-state model of transcription regulation

A two-state model has been shown to give a satisfactory description of the stochastic properties of transcription, in particular bursting (27,29,37). According to this model, promoters can be either in an *on* state from which mRNA molecules are produced with a rate k_{tx} , or in an *off* state, in which transcription does not occur. The rates of transition from the *on* to the *off* states and vice versa are k_{off} and k_{on} , respectively. Additionally, mRNA degradation can occur at a rate k_{deg} . When the half-life of mRNA molecules is larger than the average time in which the promoter is active ($k_{off} \gg k_{deg}$), the stationary solution for the distribution of mRNA molecules as function of the kinetic parameters is Equation (1), the negative binomial distribution (29,31):

$$P(m) = \frac{\Gamma(m+r)p^r(1-p)^m}{m!\Gamma(m)} \quad (1)$$

where m is the number of mRNA molecules, $r = k_{on}/k_{deg}$, $p = k_{off}/(k_{off} + k_{tx})$ and $\Gamma(m)$ is the Gamma function. When $k_{on} \gg k_{off}$, i.e. when the gene is primarily in the *on* state, the negative binomial distribution reduces to a Poisson distribution. The mean μ and variance σ^2 of $P(m)$ are given respectively by $\mu = rp/(1-p)$ and $\sigma^2 = r(1-p)/p^2$, from which the noise can be evaluated:

$$\frac{\sigma^2}{\mu^2} = \frac{1}{r(1-p)} \quad (2-a)$$

or alternatively as:

$$\frac{\sigma^2}{\mu^2} = \frac{1}{\mu p} \quad (2-b)$$

The difference between Poissonian noise and this expression is the additional factor of $1/p$ in Equation (2-b).

Fitting a threshold-linear dependence to mean transcript numbers data

To carry out fits of Equation (4) for the mean transcript number per cell m as function of the promoter activity of RyhB $\alpha_s = \alpha_{RyhB}$, with λ as the only fitting parameter, to the experimental data of μ_{sodB} and μ_{fumA} , we regarded the transcript production rates α_{sodB} , α_{fumA} of both genes as fixed, and estimated their values from their respective degradation rates β_{sodB} and β_{fumA} , under iron-rich conditions in which the production of RyhB can be neglected. The measured

half-lives of both transcripts, 6.8 min for *sodB* (see Results) and 4 min for *fumA* (39), yield $\beta_{\text{sodB}} \simeq 0.102 \text{ min}^{-1}$ and $\beta_{\text{fumA}} \simeq 0.173 \text{ min}^{-1}$. The lifetimes of both transcripts lie in the range of long values measured under iron-rich conditions in a genome-wide study of mRNA degradation in this *E. coli* strain (40). From Equation (4), we obtain $m = \alpha_m/\beta_m$ for $\alpha_s = 0$, and therefore $\alpha_{\text{sodB}} \simeq 0.694 \text{ min}^{-1}$ and $\alpha_{\text{fumA}} \simeq 0.412 \text{ min}^{-1}$. We note that while α_{sodB} and α_{fumA} have been observed to decrease and increase respectively with the extent of iron deprivation (25), these variations are small over the regime in which fitting the present data with Equation (4) is relevant. The values of α_{RyhB} for different concentrations of DTPA were obtained from a fit of a Hill function to the measured promoter activity of *RyhB* shown in Figure 2A. Finally, the value of $\beta_s = \beta_{\text{RyhB}} \simeq 0.02 \text{ min}^{-1}$ was deduced from previous measurements (19,41).

RESULTS

Effects of iron deprivation on *RyhB* target transcript distributions

To characterize the effect of post-transcriptional regulation by *RyhB* on the cell–cell transcript variability of one of its main targets, *sodB*, we carried out smFISH measurements in individual cells, for different levels of iron deprivation. Fluorescence and phase contrast images of wild-type *E. coli* cells exposed to different concentrations of the cell-impermeable iron chelator diethylene triamine penta-acetic acid (DTPA) are shown in Figure 1A. Cells were fixed at least 3 h after exposure to DTPA, in order to allow adaptation to iron-poor conditions and avoid gene expression oscillations that result upon sudden exposure to DTPA (35). Spots within cells in the fluorescence images correspond to either single or few *sodB* transcripts, and the determination of the transcript number in each cell is carried out by quantifying localized fluorescence (see Supplementary Methods and Supplementary Figure S1), following well established procedures (42). Distributions of *sodB* transcript number per cell for four representative concentrations of DTPA that lead to different levels of *RyhB* production (43) are shown in Figure 1B. We carried out similar measurements of transcripts of *fumA*, another gene whose transcripts undergo stoichiometric degradation with *RyhB*. Distributions of *fumA* transcripts for four representative values of DTPA concentration are shown Figure 1C. Two features of these distributions are noteworthy. First, their respective means shift toward smaller values as the extent of iron deprivation increases. Second, each distribution corresponding to a given value of DTPA concentration is unimodal and similar to those characterizing mRNAs that are not sRNA targets (26), in spite of reports of a mixed double-negative feedback loop configuration involving *RyhB* and *Fur* (21). Such motifs may lead to bistability (3).

To confirm that the behavior of *RyhB* target transcripts we observed is indeed due to *RyhB*-induced degradation and components of the degradosome, we studied the effects of iron deprivation in various mutant strains (Supplementary Figure S2). As expected, we found that the mean transcript numbers of *sodB* and *fumA* are rather insensitive to iron deprivation when experiments were conducted in a

ΔryhB mutant relative to the wild type strain (Supplementary Figure S2A). Similar measurements with a Δhfq mutant also display a weaker decay in mean transcript number with iron deprivation, when compared to the wild type strain (Supplementary Figure S2B). Control experiments with a strain in which RNase E function is impaired by truncation of the last 360 amino acids of the C-terminal domain (34) show that the degradation of two *RyhB* targets is visibly reduced with iron deprivation (Supplementary Figure S2C). Note that the three mutants in which the function of components of the degradosome was impaired, display larger values of the mean transcript number relative to the wild type strain. Finally, we also conducted similar experiments with a mutant in which the RNase III function was impaired and found that under our experimental conditions, the behavior of *RyhB* target transcripts was similar to the wild type (Supplementary Figure S2D), in agreement with previous findings (19).

Switch-like activation of the network with increasing iron deprivation

Our initial analysis of the effect of iron levels on the distributions of *sodB* and *fumA* mRNAs per cell revealed two types of behavior: one type characterized by a peak at a nonzero number of transcripts (at about 3–4 *sodB* mRNAs, as illustrated by the top histograms in Figure 1B; 1–2 *fumA* mRNAs, as illustrated by the top histograms in Figure 1C), and the other type having a peak at zero transcripts per cell (bottom histograms in Figure 1B and C). To monitor how the behavior of transcript distributions changes between these two types in more detail, we carried out measurements of transcript distributions with a finer resolution in DTPA concentration. Surprisingly, we found that the transition between the two types of *sodB* mRNA distributions occurred sharply, as illustrated by a plot of the mean *sodB* transcript number μ as a function of DTPA concentration (Figure 2A): μ decreases sharply from 4.9 ± 0.6 transcripts per cell to 0.9 ± 0.2 , within a very narrow window of DTPA concentrations. A fit of a Hill function to the data, also shown in Figure 2A, yields an effective Hill coefficient of $n = 15 \pm 6$, a value that underscores the sharpness of the transition. This switch-like behavior of the mean *sodB* transcript number as a function of iron deprivation by addition of DTPA was confirmed using a qRT-PCR technique on total extracted *sodB* mRNA (Supplementary Figure S3). Switch-like behavior is also displayed by *fumA* transcripts as function of DTPA concentration (Figure 2B). To find out whether the switch-like change in μ is also observed upstream of *sodB*, we measured the promoter activity of *ryhB* in cell ensembles bearing plasmids with a promoter fusion, *pryhB*–YFP (Materials and Methods section). The promoter activity of *ryhB*, fitted with a Hill function is also shown in Figure 2A. Clearly, *RyhB* production also displays switch-like activation that mirrors the trend in the mean *sodB* transcript number, indicating that the origin for the switch lies upstream of *ryhB* transcription, i.e. in its regulation. Addition of iron to cells grown in M9 medium also resulted in a sharp reduction in the promoter activity of *ryhB*, and switch-like behavior of the mean *sodB* transcript number, with a Hill coefficient $n = 6 \pm 2$ (Supplementary Figure

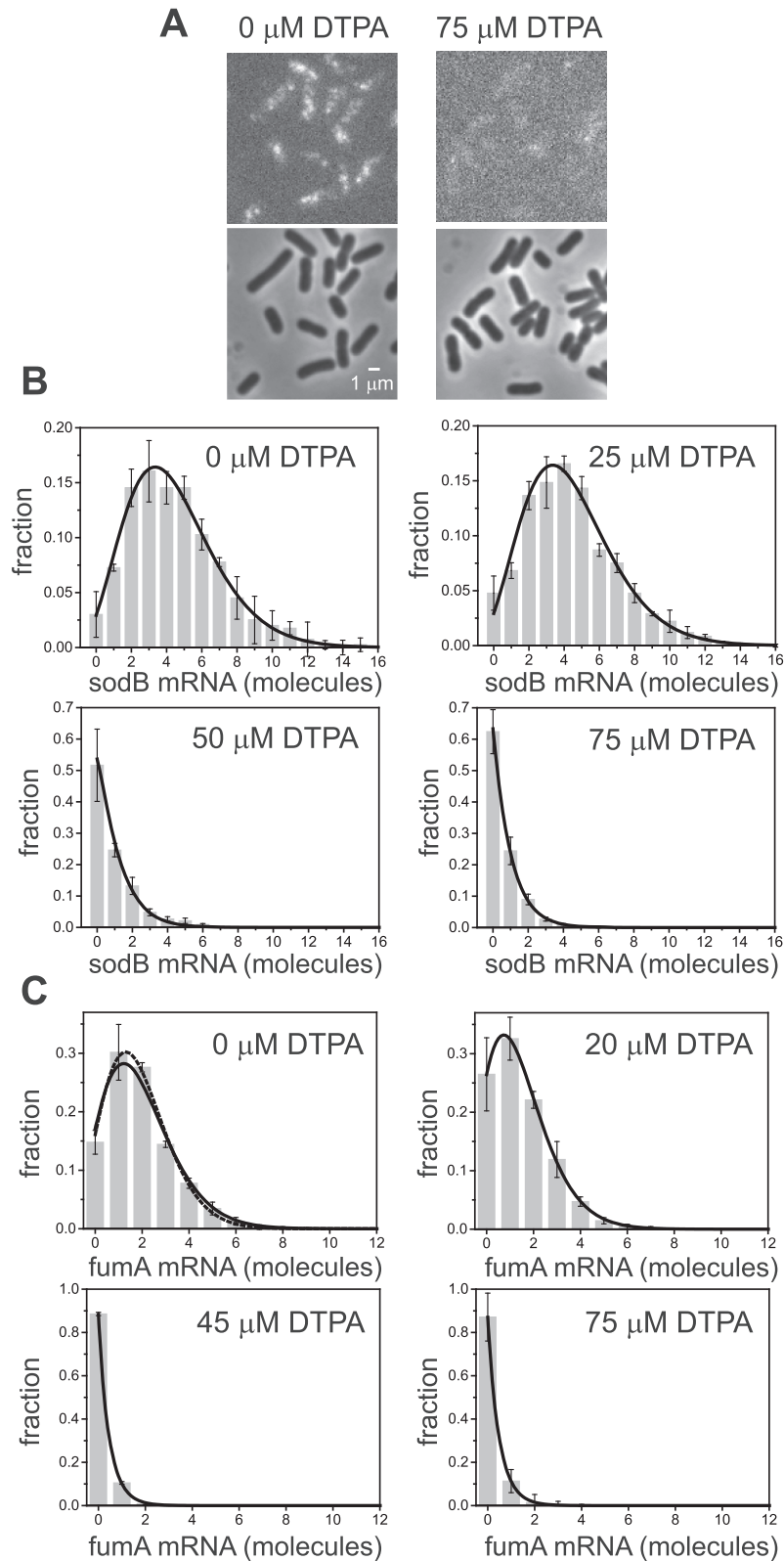


Figure 1. Effects of iron deprivation on *sodB* and *fumA* transcripts visualized using single-molecule FISH (smFISH). (A) Top: typical images showing labeled *sodB* transcripts under iron-rich and iron-poor conditions. Bottom: phase contrast images of the same fields of view. (B) Distributions of *sodB* transcript number in individual cells, for the indicated concentrations of the iron chelator DTPA. The distributions represent an average over four experimental repeats while error bars represent standard errors. The solid lines are fits to the experimental data using the negative binomial distribution (Equation 1). (C) Distributions of *fumA* transcript number in individual cells, for the indicated concentrations of the iron chelator DTPA. The distributions represent an average over three experimental repeats while error bars represent standard errors. The solid and dashed lines are fits to the experimental data using the negative binomial Equation (1) and the Poisson distributions, respectively.

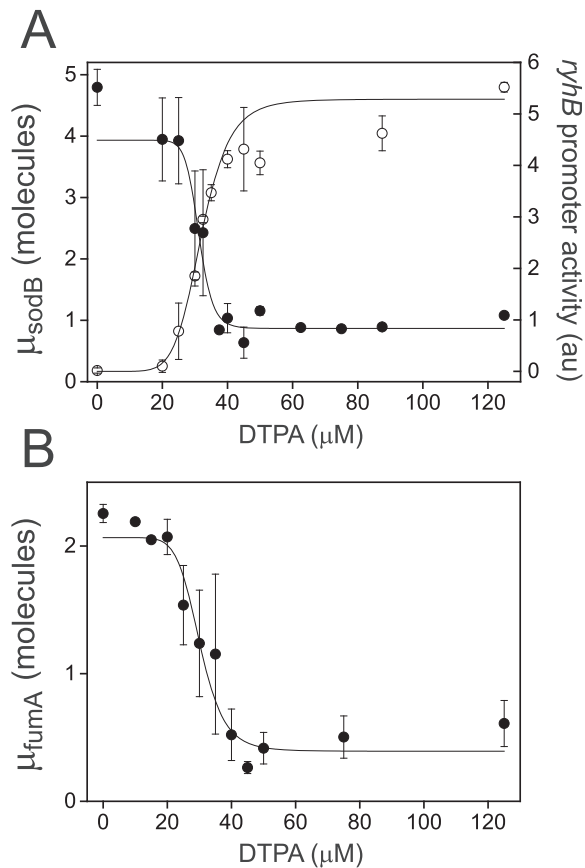


Figure 2. Switch-like behavior of the mean *sodB* and *fumA* transcript numbers upon iron deprivation. **(A)** Mean *sodB* transcript number μ_{sodB} in individual cells as function of the concentration of the cell-impermeable iron-chelator DTPA (black dots). Also plotted is the promoter activity of RyhB (empty circles), with reporter plasmids encoding a *pryhB-yfp* fusion. The black lines through both sets of data, are fits with Hill functions ($y = a + (b - a)(x^n / (k^n + x^n))$) with $a = 4.0 \pm 0.4$, $b = 0.9 \pm 0.1$, $k = 1.25 \pm 0.04$, $n = 15 \pm 6$ for *sodB* transcripts and $a = -0.15 \pm 0.23$, $b = 5.0 \pm 0.2$, $k = 1.25 \pm 0.03$ and $n = 11.0 \pm 2.6$ for *pryhB*. **(B)** Mean *fumA* transcript number μ_{fumA} in individual cells as function of the concentration of the cell-impermeable iron-chelator DTPA (black dots). Data were obtained from four 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 $a = 2.1 \pm 0.2$, $b = 0.4 \pm 0.1$, $k = 1.21 \pm 0.09$, $n = 8 \pm 4$.

S4). While this value of the Hill coefficient is smaller than when adding DTPA to LB ($n = 15 \pm 6$), the cooperative behavior is still significant. To test whether other genes whose transcription is regulated by Fur also display switch-like behavior independent of RyhB, we measured the mean transcript number of *sufC*, which like *ryhB* is directly repressed by Fur. Indeed, the mean *sufC* transcript number increases in a switch-like fashion over the same DTPA range as the mean *sodB* mRNA decreases (Supplementary Figure S5). In addition to stoichiometric degradation, RyhB can also down-regulate translation by blocking the ribosome binding site of a target transcript. We have tested the effects of RyhB on *fepB*, a target with which RyhB base-pairs in the region around the Shine-Dalgarno sequence (44). The observed increase in the mean transcript number is due to direct derepression by Fur, in agreement with previous ob-

servations (45) (Supplementary Figure S6). It is noteworthy that the mean transcript number of the *shiA* gene within the iron homeostasis network, whose transcripts' translation is activated by binding to RyhB, increases with iron deprivation (Supplementary Figure S6). This observation is consistent with the increase in *shiA* transcript stability due to its interaction with RyhB (46). As expected, expression from chromosomal *lac* promoters, which are outside the iron homeostasis network, is independent of iron deprivation (25).

Noise of stoichiometrically degraded target RyhB transcripts increases with iron deprivation

The above measurements of transcript distributions allow us to test experimentally for the existence of theoretically predicted enhanced intrinsic fluctuations of target transcripts that undergo stoichiometric degradation with RyhB. To this end, we quantified the cell-to-cell variability of *sodB* and *fumA* transcripts as a function of iron deprivation by calculating the transcript noise, defined by the nondimensional ratio σ^2/μ^2 , where σ is the standard deviation and μ the mean of mRNA numbers per cell. Different values of μ were obtained by varying the DTPA concentration that cells were exposed to. Data were obtained from seven independent experimental runs in the case of *sodB* transcripts, color-coded to illustrate reproducibility, with four to seven different concentrations of DTPA per run, covering different regimes. Plots of the transcript noise of *sodB* and *fumA* as a function of their respective means μ are shown in Figure 3A and B. Transcript noise increases monotonically in both cases as the average number of transcripts decreases and as RyhB transcription increases. This contrasts sharply with measurements of SodB and FumA protein noise, which remain constant over a wide range of RyhB production rates (25). It is noteworthy that there is no signature of a theoretically predicted peak in transcript noise within the range of values of μ of our experiments (7,10). A plot of the ratios of the promoter activities of RyhB and those of *sodB* or *fumA*, as a function of the concentration of DTPA, shown in Supplementary Figure S7, verifies that our measurements access the regime in which enhanced stochastic fluctuations are expected, i.e. when the production rates of RyhB and either of the two genes are comparable (10).

We reasoned that the narrowness of the window of DTPA concentrations spanning the switching region, over which RyhB production changes from values close to leaking production on one hand, to high production on the other (Figure 2A), may preclude the observation of enhanced fluctuations in transcript noise. In spite of repeated attempts with increasing resolution in changes of iron deprivation, the noise was observed to increase monotonously with iron stress for both *sodB* and *fumA* transcripts, as shown in Figure 3. We therefore constructed a mutant strain in which RyhB production is independent of iron deprivation and controlled from an inducible arabinose promoter. The dose response of this promoter fused to YFP, shown in Supplementary Figure S8, exhibits graded behavior. The effects of RyhB induction on the *sodB* mean transcript number in this strain are shown in Supplementary Figure S9. The measurements of *sodB* transcript noise in this strain as function of

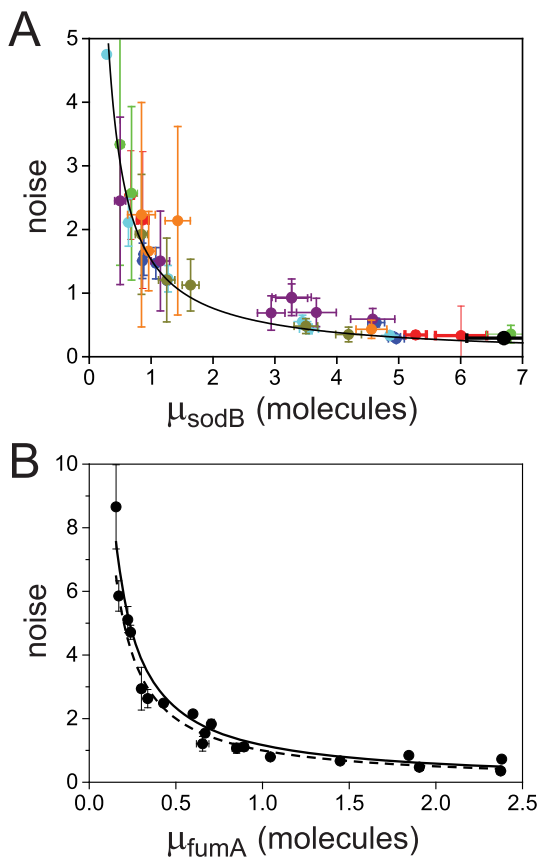


Figure 3. Noise of *sodB* and *fumA* transcripts as a function of their respective mean transcript number per cell. **(A)** Noise of *sodB* transcripts measured by the ratio σ^2/μ^2 , where σ is the standard deviation and μ the mean of mRNA numbers per cell. The data were obtained from seven independent smFISH runs in which cells were exposed to different concentrations of DTPA for 3 h prior fixation, each color representing one experimental run, to illustrate the reproducibility of the data. Error bars were determined from 1000 bootstrap samples. The solid line is a fit to the experimental data using the expression for noise Equation (2-b) obtained from the negative binomial distribution Equation (1). The noise versus mean in a *RyhB*-deleted strain is shown with a black circle. Error bars are the standard error from three independent experiments. **(B)** Transcript noise as a function of μ_{fumA} the mean number of *fumA* mRNA molecules per cell. Noise is defined by the ratio σ^2/μ^2 , where σ_{fumA} is the standard deviation of *fumA* mRNA molecules. The data were obtained from two independent smFISH runs shown separately, in which cells were exposed to different concentrations of DTPA for 3 h prior fixation. Error bars were determined from 1000 bootstrap samples. The solid line is a fit to the experimental data using the expression for noise Equation (2-b) obtained from the negative binomial distribution Equation (1), while the dashed line represents the noise of a Poisson distribution $\sigma_{fumA}^2/\mu_{fumA}^2 = 1/\mu_{fumA}$.

the *sodB* transcript number (Supplementary Figure S10), display fully monotonic behavior and largely overlap with those measured in the wild-type strain as function of DTPA concentration. There is no discernible evidence of a peak in noise due to enhanced stoichiometric fluctuations.

Direct measurement of RyhB and its noise as function of iron deprivation

Considerable insights can be obtained by monitoring *RyhB* and its cell-cell variability by detecting *RyhB* directly in individual cells as a function of iron deprivation, using

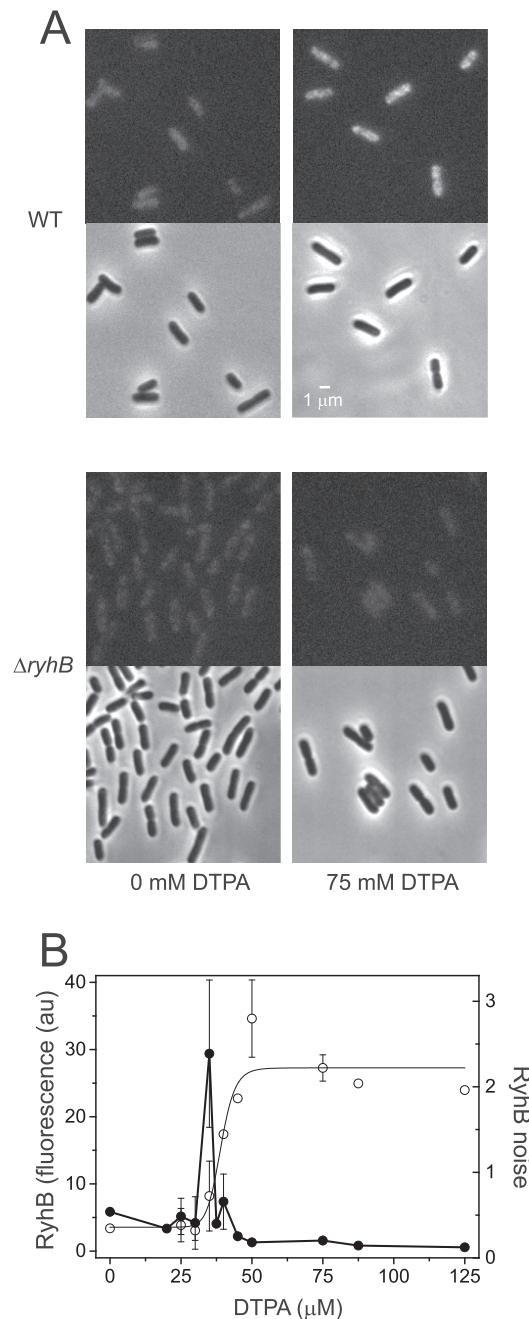


Figure 4. Direct measurement of *RyhB* in individual cells and its noise. **(A)** Top: typical snapshots of fluorescence from undegraded intracellular *RyhB*, tiled with labeled oligonucleotides under iron-rich (0 μ M DTPA) and iron-poor (75 μ M DTPA) conditions, in wild-type cells. The corresponding phase contrast images of the same fields of view are appended below each frame. Bottom: control images obtained in a Δ *ryhB* background under the same conditions. **(B)** Empty circles: Switch-like behavior of the fluorescence density per cell from undegraded intracellular *RyhB*, tiled with labeled oligonucleotides, as function of DTPA concentration. The density is defined as the total cell fluorescence divided by the cell area. The line is a fit with a Hill function with $a = 4 \pm 3$, $b = 27 \pm 2$, $k = 1.56 \pm 0.07$ and $n = 16.0 \pm 9$. Full circles: *RyhB* noise $(\sigma_f/\mu_f)^2$, where σ_f and μ_f are the standard deviation and mean of the fluorescence density of undegraded intracellular labeled *RyhB*. Values of the fluorescence density and of the noise represent means over three independent experiments and the error bars represent standard errors. Crosses: control measurements of *RyhB* noise measured in an RNase E mutant strain in which stoichiometric degradation is abrogated, at 0 and 75 μ M DTPA.

FISH methods. FISH techniques are not used here in single-molecule modality, since RyhB molecules are tiled by maximum three oligomeric probes and therefore their intensity is close to that of the background. Furthermore, individual RyhB molecules cannot easily be resolved given their large number (Materials and Methods). Typical snapshots of cells displaying fluorescence from labeled RyhB under rich and poor iron conditions (0 and 75 μM DTPA, respectively) are shown in Figure 4A, for both the wild-type strain and a control ΔryhB mutant. Two features are noteworthy: first, RyhB production clearly increases with iron deprivation and second, a comparison with the control strain indicates that RyhB production can also be detected under iron-rich conditions. Consistent with this, *sodB* and *fumA* mean transcript numbers are higher in the ΔryhB mutant, as well as in other mutants in which components of the degradosome are impaired (Supplementary Figure S2). The mean fluorescence density of RyhB—defined as the total cell fluorescence divided by the cell's area, averaged over the cell population—is shown in Figure 4B. The behavior of the mean fluorescence density of RyhB recapitulates the switch-like characteristics of the RyhB promoter activity. To estimate what fraction of the total RyhB production is detected in these experiments, we compared the mean fluorescence density of RyhB measured in an RNase E mutant strain in which stoichiometric degradation is abrogated, with the wild type, at 75 μM DTPA (Supplementary Figure S11). The ratio between the two is 1.8 ± 0.7 under conditions in which the production rate of RyhB is high, indicating that about half of the total RyhB produced is degraded with its targets and demonstrating that RyhB sRNA is not a limiting factor at low iron levels. We have also characterized the dependence of the noise in RyhB production as a function of DTPA concentration. A plot of RyhB noise is shown in Figure 4B, while distributions of RyhB in individual cells for selected DTPA concentrations are shown in Supplementary Figure S12. Significantly, a narrow, strong peak in RyhB noise is observed, exposing strong fluctuations in RyhB concentration within the switching regime. It is also noteworthy that RyhB noise is 5–10 times smaller than *sodB* transcript noise above the switch, a regime in which RyhB production is large. The noise of both RyhB and *sodB* are comparable below the switch. Control measurements of RyhB noise in an RNase E mutant strain in which stoichiometric degradation is abrogated, at 0 and 75 μM DTPA also display small noise levels.

Kinetic parameters within the framework of a two-state model

We tested the applicability of the generalized two-state model (Materials and Methods) to describe the variability of targets whose degradation rate is not a constant, but is modulated by an sRNA. We assumed a mean field approximation in which the stochasticity of sRNA levels is smaller than the intrinsic fluctuations in mRNA production, and that Hfq is not a limiting factor. Therefore, degradation is set by the mean number of sRNA molecules in the cells.

The experimental noise data are well fitted by Equation (2-b), taking a fixed value of $p = k_{\text{off}}/(k_{\text{off}} + k_{\text{tx}})$, where k_{tx} is the transcription rate and k_{off} is the rate of switching

back to the off state, when the promoter is active. Fits for the noise of *sodB* and *fumA* transcripts are shown in Figure 3A and B for different experiments in which the concentration of DTPA is varied. In the case of *sodB*, the fit yields ± 0.01 , from which we obtain $k_{\text{tx}}/k_{\text{off}} = 0.52 \pm 0.02$, indicating that transcription and the transition to an inactive promoter state occur on similar timescales, and that both processes are not appreciably affected by iron deprivation. This value of the ratio $k_{\text{tx}}/k_{\text{off}}$, falls within the range of variation expected from measurements carried out on seven different promoters (27). We checked that if the ratio $r = k_{\text{on}}/k_{\text{deg}}$ between the rates of gene activation k_{on} and degradation k_{deg} is taken as fixed, the fit to the data is quite poor, suggesting that k_{deg} is altered when the DTPA levels are modified, as expected. Additionally, the transcript histograms are well fitted by the negative binomial distribution (Equation 1) over the whole range of DTPA concentrations, as illustrated in Figure 1B, using r as the only fitting parameter and the value of P obtained above. Similarly, in the case of *fumA*, a fit of Equation (2-b) to the data yields $p = 0.85 \pm 0.05$, a value that is closer to $p = 1$ than for *sodB* transcripts, suggesting that the transcript noise of *fumA* has a more Poissonian character. While a Poisson distribution without fitting parameters describes the data well (Figure 3B), the negative binomial distribution is characterized by an extra degree of freedom that allows us to evaluate the RyhB-induced degradation rate. Fits of experimentally-measured *fumA* transcript histograms with negative binomial distributions are shown in Figure 1C.

Using the values of p for *sodB* and *fumA* transcripts, one can now extract the respective *in vivo* degradation rates and their dependence on the degree of iron deprivation. k_{deg} can be obtained from the relation $k_{\text{deg}} = k_{\text{on}}/r$ using Equation (2-a). To determine k_{on} in the case of *sodB*, we carried out measurements of the half-life of *sodB* transcripts for 0 and 75 μM DTPA using qRT-PCR and a transcription inhibitor (Materials and Methods). These concentrations of DTPA are representative of low and high expression levels of RyhB. The values we obtained, $k_{\text{deg}} = 1.7 \pm 0.2 \times 10^{-3} \text{ s}^{-1}$ and $8.1 \pm 0.3 \times 10^{-3} \text{ s}^{-1}$, respectively (plotted in Figure 5A), are in good agreement with the values obtained from our analysis based on the two-state model when choosing $k_{\text{on}} = 0.01 \text{ s}^{-1}$, a value that falls well within the range of values measured for different promoters in *E. coli*, which can display approximately a ten-fold variation (21). The dependence of k_{deg} on the *sodB* mean transcript number μ is shown in Figure 5A. The degradation rate increases about 8-fold relative to the value under iron-rich conditions, as the mean transcript number decreases over the full range of values of DTPA we tested. From the k_{deg} values obtained from the qRT-PCR measurements we calculated the corresponding half-lives under these conditions: $t_{1/2} = 6.74 \pm 0.7 \text{ min}$ and $t_{1/2} = 1.43 \pm 0.1 \text{ min}$, respectively (Supplementary Figure S13). These values are in agreement with half-life measurements of *sodB* transcripts reported previously (19). Recently the association/dissociation of the SgrS sRNA interacting with two of its targets has been studied using super-resolution microscopy and mean-field kinetic equations, which neglect any fluctuations including bursty transcription and possible limitation of Hfq (47). The degradation rates obtained

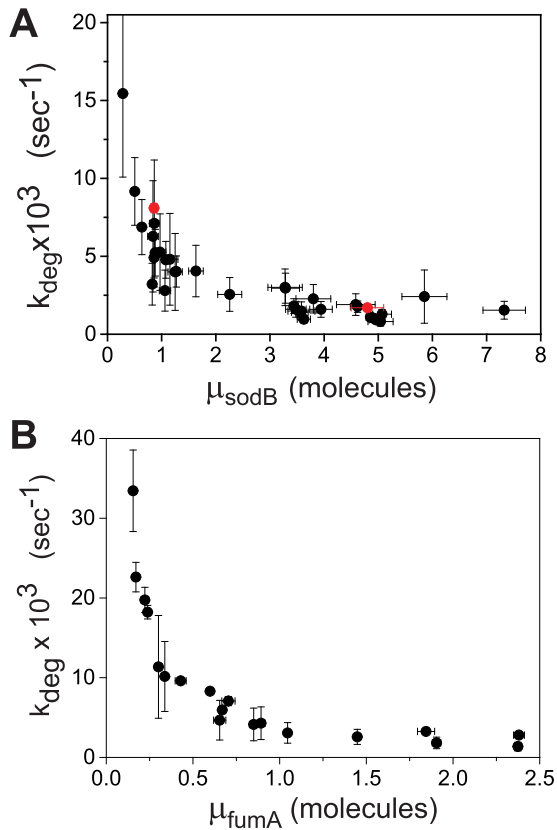


Figure 5. Degradation rates k_{deg} of *sodB* and *fumA* transcripts calculated from an analysis of noise data with a generalized two-state model of bursty transcription. (A) Degradation rate k_{deg} of *sodB* mRNA as function of the mean *sodB* transcript number. The values of k_{deg} for *sodB* transcripts were obtained from Equation (2-a), with the value of $p = 0.66 \pm 0.01$ obtained from a fit to the noise (Figure 3A) and $k_{\text{on}} = 0.01 \text{ s}^{-1}$. Error bars were determined from propagation of the noise data (Figure 3A), via Equation (2-a). Red points represent independent measurements of k_{deg} using qRT-PCR at 0 and 75 μM DTPA (Supplementary Figure S13). (B) Degradation rate k_{deg} of *fumA* mRNA as function of the mean *fumA* transcript number. The values of k_{deg} were obtained from Equation (2-a) using the value of $p = 0.86 \pm 0.05$ obtained from a fit to the noise data (Figure 3B) and $k_{\text{on}} = 0.027 \text{ s}^{-1}$. Error bars were determined from 1000 bootstrap samples of the noise data and propagated via Equation (2-a).

in that study are comparable to those we observed under conditions in which RyhB production is high (e.g. at 75 μM DTPA).

To check what the degradation rate of *sodB* transcripts is in the absence of RyhB production, we carried out sm-FISH measurements of *sodB* transcripts in a *ryhB*-deleted strain under iron-rich conditions. We obtained a mean *sodB* transcript number of 6.7 ± 0.6 , within the range of values obtained for the RyhB wild type strain under iron-rich conditions (4.9 ± 0.6 transcripts per cell). The noise σ^2/μ^2 as a function of mean for this strain is shown in Figure 3A. The data point falls within experimental error on the same curve as the data for the wild-type strain, allowing us to use the same parameters as for the wild-type in the determination of k_{deg} using the two-state model. We obtained $k_{\text{deg}} = 1.0 \pm 0.1 \times 10^{-3} \text{ s}^{-1}$ ($t_{1/2} = 11.5 \pm 1.2 \text{ min}$), in fair agreement with values for the wild type under iron-rich conditions, consistent with the literature (27). The slightly higher degradation

rate values for the wild type could reflect RyhB promoter leakage.

The degradation rate of *fumA* transcripts for different levels of iron deprivation can be evaluated following a similar procedure. Using $k_{\text{on}} = 0.01 \text{ s}^{-1}$ as for *sodB* transcripts yields $t_{1/2} \sim 6 \text{ min}$ for the half-life of *fumA* transcripts under iron-rich conditions, a value which is higher than measured values by $\sim 50\%$ (39). To match the measured half-life of *fumA* transcripts ($\sim 4 \text{ min}$), a higher value, $k_{\text{on}} = 0.027 \text{ s}^{-1}$ is needed. Using the latter yields the results plotted in Figure 5B. The RyhB-dependent degradation rate of *fumA* transcripts is higher than for *sodB* transcripts.

Estimate of the *in vivo* effective interaction strength between RyhB and its target *sodB* and *fumA* transcripts

A simpler mean-field kinetic model for sRNA-mediated silencing that neglects bursty mRNA production and fluctuations leads to the characteristic threshold-linear switching behavior of stoichiometric regulation (8). According to this model, cellular numbers of an sRNA (s) and a target mRNA (m) obey the following rate equations:

$$\frac{dm}{dt} = \alpha_m - \beta_m m - kms \quad (3)$$

$$\frac{ds}{dt} = \alpha_s - \beta_s s - kms$$

where α_m and α_s are the effective – constant – production rates of the target and sRNA, respectively, β_m and β_s the respective degradation rates, and the second order kinetic constant k is a measure of the effective strength of the sRNA–mRNA interaction. Under steady-state conditions, the solution to Equation (3) is:

$$m = \frac{1}{2\beta_m} \left[(\alpha_m - \alpha_s - \lambda) + \sqrt{(\alpha_m - \alpha_s - \lambda)^2 + 4\alpha_m\lambda} \right] \quad (4)$$

where $\lambda = \beta_m\beta_s/k$. We have plotted the mean transcript per cell μ_{sodB} and μ_{fumA} as a function of the promoter activity $\alpha_s = \alpha_{\text{RyhB}}$ in Figure 6. Fits of Equation (4) to both sets of data with λ as a fitting parameter and fixed values of the respective promoter activities α_m (Materials and Methods) are shown in Figure 6. From the values of λ derived from the best fits we obtain $k_{\text{sodB}} = 2.0 \pm 0.3 (\times 10^{-3} \text{ min}^{-1})$ and $k_{\text{fumA}} = 4.3 \pm 1.7 (\times 10^{-3} \text{ min}^{-1})$.

DISCUSSION

In the present work, we have studied how the post-transcriptional downregulation of gene expression by the sRNA RyhB affects various aspects of the cellular response to graded changes in iron availability. Our experiments were carried out at the RNA level, where RyhB naturally exerts its effects. At the RNA level, fluctuations are dominated by intrinsic noise sources (27), in contrast to the dominance of extrinsic fluctuations when protein levels are assessed (25,26). We discuss below the impact of RyhB on various aspects of the cellular response to iron scarcity.

Our analysis of the dependence of transcript distributions on iron availability shows that the two-state model, which

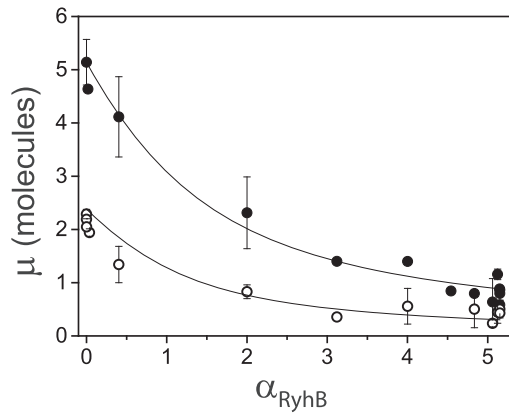


Figure 6. Threshold-linear switching and *in vivo* effective interaction strength between RyhB and its target *sodB* and *fumA* transcripts. Mean transcript per cell μ_{sodB} (full circles) and μ_{fumA} (empty circles) as function of α_{RyhB} , the promoter activity of RyhB (Figure 2A). The lines represent best fits of Equation (4) to both sets of data, with either $\lambda = \beta_{sodB}\beta_{RyhB}/k_{sodB}$ or $\lambda = \beta_{fumA}\beta_{RyhB}/k_{fumA}$ as a fitting parameters, where β_m are the degradation rates of either of the two gene transcripts, β_{RyhB} that of RyhB, and k is the effective strength of interaction between RyhB and either of the transcripts *sodB* or *fumA*. Error bars represent standard errors over at least three experimental runs.

has been used successfully to describe bursts of transcription created by infrequent initiations by RNA polymerase in individual cells (27), can be generalized to include RyhB-mediated transcript degradation. Remarkably, this analysis demonstrates that the resulting negative binomial distribution describes well the experimentally measured transcript distributions over the whole range of DTPA concentrations by changing only one parameter, namely the *in vivo* rate of degradation k_{deg} . RyhB increases the degradation rates of *sodB* and *fumA* mRNA, by a factor of five or more when comparing iron-rich to iron-poor conditions.

Analysis of mean transcript numbers of both *sodB* and *fumA* as the levels of iron deprivation vary, in terms of a mean-field threshold-linear model behavior has allowed us to estimate k , the effective *in vivo* strength of interaction between RyhB and each of its two targets *sodB* and *fumA* transcripts. Remarkably, the value of k_{fumA} is about twice larger than that of k_{sodB} . While it has been considered that *sodB* mRNA is the target whose expression RyhB affects most strongly (20), the relative reductions we observe in the abundances of *sodB* and *fumA* transcripts as function of iron availability are comparable. Moreover they are similar to those observed in other experiments (48,49), and at the level of the respective proteins (17). It is noteworthy that the predicted hybridization energies between RyhB and either *sodB* or *fumA* transcripts differ by only $\sim 10\%$, including corrections due to molecular unfolding (50,51). We stress however that in addition to these factors, k includes contributions from Hfq-mediated binding and the degradosome (52), as well as effects due to possible different spatial colocalizations of *sodB* or *fumA* transcripts with RyhB (47).

Transcript distributions for both genes are unimodal for the range of DTPA concentrations in our experiments. While iron deprivation induces the expression of RyhB through derepression by Fur, RyhB was reported to down-regulate Fur, albeit indirectly (21), thereby establishing a

double-negative feedback loop. A double-negative feedback motif may give rise to bistable behavior in the case of two transcription factors negatively regulating each other (22), suggesting similar behavior could be observed for the mixed sRNA-transcription factor motif (24,53). The notion of bimodal behavior has also been considered in the case of the sRNA MicF, which forms a double negative feedback loop with the global transcription factor Lrp (54). The unimodal character of the distributions we observe does not support this notion. Consistent with our observations, Gillespie simulations of a mixed loop involving a transcription factor and an sRNA show a limited region in parameter space where such a loop exhibits bistability (24). While the absence of bistability does not contradict the existence of a mixed double-negative feedback loop involving Fur and RyhB, we note that other biochemical studies have failed to reproduce this motif (48).

The predominance of extrinsic noise sources in controlling cell-cell variations at the protein level was suggested in our earlier work as a possible factor masking the theoretically predicted enhanced fluctuations in the crossover regime, which separates the strongly repressed, from the expressing regime characteristic of intrinsic noise in stoichiometrically dominated systems (25). Extrinsic noise stems from cell-cell variations in factors such as RNA polymerases, ribosomes and metabolites that affect the expression of all genes equally. In contrast, intrinsic noise originates from the stochastic nature of biochemical processes such as transcription and translation, and manifests itself as differences in expression between two copies of the same promoter, each labeled with a different reporter, within the same cell (14,55). The present experiments, conducted at the transcript level in which intrinsic noise is dominant, reveal no evidence of a peak in noise due to enhanced stoichiometric fluctuations, within a range of RyhB production rates in which the mean transcript number decreases for almost its full span. We surmised that the very narrow range of iron concentrations over which switch-like activation of RyhB production takes place in the wild-type strain prevented us from observing enhanced near-critical fluctuations, and therefore constructed a strain in which RyhB production is graded, by expressing it from an inducible promoter. No enhanced fluctuations were observed in this strain either. A number of factors, which we now discuss, may account for their absence: the possible limiting nature of Hfq (56,57); the presence of auxiliary weak RyhB targets (6), the presence of highly expressed 3' external transcribed spacer (3' ETS) fragments of pre-tRNAs (58) and finally, inefficient binding and degradation of RyhB with its targets (8,10).

Previous calculations of enhanced fluctuations did not take any of these factors into account, except for inefficient binding (7,9,10). Hfq limitation may be significant under certain conditions (56,57), leading to a reduction in target transcript-RyhB coupling and thus target degradation, thereby limiting the range of possible fluctuations for given production rates of RyhB and its targets transcripts. The fact that *sodB* transcript numbers do not decay to zero at high concentrations of DTPA for which undegraded RyhB is directly detected, is consistent with Hfq limitation. Calculations indicate that the presence of auxiliary weak targets diminishes the correlation between sRNA numbers and

those of main transcripts, reducing significantly fluctuations (6). The significant numbers of predicted RyhB targets that remain to be validated (18,44,50,51), precludes experiments in which mutations in both RyhB and *sodB* would make the latter a sole target and thus provide a more stringent test of this notion (38). Lastly, numerical calculations indicate that the strength of enhanced fluctuations depends strongly on k (see Equation 3), the effective strength of the sRNA–mRNA interaction (10). As k grows, enhanced fluctuations are larger and at the same time, threshold-linear switching becomes sharper (59). The rounded behavior of the threshold-linear response of both *sodB* and *fumA* transcripts we observed (Figure 5) yields instead moderate values k for both transcripts. This suggests that enhanced fluctuations may not be large enough to be observable. It has recently been reported that the abundant 3' ETS^{leuZ} fragment of the *glyW-cysT-leuZ* polycistronic tRNA precursor base pairs with RyhB, sequestering the latter and thus acting as a sponge that may buffer noise (58). Disruption of this pairing increases RyhB activity significantly, even under iron-rich conditions. Interestingly, Hfq mediates the interaction between the polycistronic tRNA transcript and RyhB (58,60). Finally, the presence of auxiliary targets, together with the switch-like production of RyhB, contrive on one hand to attenuate the presence of enhanced fluctuations in the principal RyhB target, and on the other hand, minimize conditions under which these fluctuations could appear. This may confer an evolutionary advantage in populations of wild-type cells, ensuring that all cells activate the iron homeostasis network in response to iron stress. Effects of auxiliary targets on noise have also been proposed in the case of microRNAs in eukaryotes (61).

Iron, as an essential trace element, is known to affect cell growth in *E. coli* through several pathways (62). Consistently, the cell doubling time increases with iron deprivation in our experiments (Supplementary Figure S14). Within the iron homeostasis network, RyhB inhibits cell growth in a RelA-dependent fashion under limiting iron conditions (63). It was proposed that RelA, a central regulator of the bacterial stringent response, carries out this function by enhancing the amount of the protein chaperone Hfq that facilitates the association of sRNAs with their target transcripts. These experiments, as others in the past (20), typically used extreme conditions of iron abundance: either media rich in iron, or media in which large concentrations of iron chelators were added.

Our experiments challenging cells with high-resolution changes in external iron availability either by the addition of an iron-selective chelator to a rich medium (LB) or iron to a minimal medium (M9) have revealed that far from showing a graded response, the iron homeostasis network exhibits switch-like activation. This is observed most clearly both in the change of the mean transcript number of RyhB targets and in RyhB's promoter activity as function of the extent of iron limitation. The apparent difference in cooperativity quantified by the Hill coefficients in both growth conditions may derive from global physiological differences caused by the growth media, since in addition to iron, these media differ in many other components (62). It is likely that the presence of multiple binding sites of Fur on the RyhB promoter that allow Fur to interact cooperatively (64), to-

gether with the polymerization of Fur on DNA substrates (64,65), may be behind the switch-like activation we observe. Consistently with this notion, when RyhB is produced from a Fur-independent promoter e.g. as in the strain with arabinose-induced production, no switch-like behavior is observed. Furthermore, the promoter of the *suf* operon contains a single predicted Fur binding site (66). Cooperative binding of proteins on DNA substrates occurs in other contexts in which precision is desired, e.g. in the case of Bicoid, providing a mechanism for threshold-dependent gene activation in the *Drosophila* embryo, allowing this transcription factor to generate sharp boundaries of gene expression (67); another example is furnished by the octamerization of the lambda repressor in the lysogenic induction switch (23). Switch-like activation of the iron homeostasis network may on one hand buffer small fluctuations in iron availability, reducing the cost of the considerable metabolic remodeling involved network activation and on the other, ensure full activation when iron deprivation becomes significant. Switch-like activation may be important in the case of pathogens, in which Fur plays a key role sensing the limited iron environment in the host, regulating many virulence factors and altering gene expression to promote pathogen survival and transmission (68).

The experiments in which we tile RyhB molecules with labeled oligomeric probes to study directly the statistics of RyhB production as a function of iron deprivation reveal a number of important features. First, the observed increase in fluorescence as a function of DTPA concentration suggests that RyhB is produced in sufficiently large amounts so as not to be a limiting factor. The promoter of RyhB is indeed known to be strong (69). Consistent with this and when the RyhB promoter is induced by iron deprivation, experiments in which RNase E was truncated and lost its ability to interact with Hfq (34), yield nearly twice the amount of RyhB compared to the strain with wild-type RNase E. This leads us to suggest that the low noise levels of RyhB we measure are unrelated to any subsequent degradation of RyhB and are due primarily to transcriptional sources. Second, the peak of RyhB noise captures sensitively the cell-cell fluctuations in RyhB availability within the switching region. Third, the fact that noise of RyhB is 5–10 times smaller than the transcript noise of *sodB* in the regime in which RyhB production is significant, justifies our approach of neglecting bursts in RyhB production and regarding RyhB in an effective way by enhancing the degradation rate of RyhB targets in the two-state model. It is important to bear in mind that the fluorescence measurements report only on the RyhB subpopulation of molecules that have not undergone mutual degradation with their targets, and that are not blocked to binding by FISH probes, e.g. by RyhB sponges (58).

Our methods to obtain *in vivo* kinetic parameters of interest of sRNA function are based on measurable changes of target transcript numbers in response to changes in sRNA production, and can therefore be applicable to other sRNAs that undergo stoichiometric degradation with their targets, e.g. SgrS and MicC (70). However they will be of limited applicability in cases in which the number of transcripts does not change such as when an sRNA activates gene expres-

sion, e.g. RyhB acting on *shiA* transcripts (46), or when an sRNA acts catalytically (71–73).

To sum up, our study has shown that the transcript noise of target genes that undergo stoichiometric degradation with an sRNA increases in a fashion consistent with intrinsic small-number fluctuations and bursty transcription, allowing the generalization of a two-state model to include sRNA-induced transcript degradation as a nonfluctuating parameter. The generalized two-state model describes our data quantitatively over a large range of iron levels, allowing us to determine the *in vivo* degradation rate of RyhB's target transcripts. Notably, the degradation rate is the only parameter that needs to be varied in order to describe quantitatively transcript distributions over a large range of levels of iron availability. The absence of enhanced cell–cell fluctuations, a hallmark of stoichiometric degradation, is consistent with the coordinate action of RyhB on a multiplicity of weak, auxiliary targets, with possible Hfq limitation, imperfect binding of RyhB to its targets and with the presence of RNA sponges sequestering RyhB. These factors provide efficient mechanisms for noise suppression. Lastly, our study has revealed that activation of the iron homeostasis network in *E. coli* by chelation of external iron is switch-like, enabling the bacterial cell to filter out small iron fluctuations and to ensure metabolic remodeling only when iron stress becomes significant.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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