Negative feedback increases information transmission, enabling bacteria to discriminate sublethal antibiotic concentrations

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In the cell, noise constrains information transmission through signaling pathways and regulatory networks. There is growing evidence that the channel capacity of cellular pathways is limited to a few bits, questioning whether cells quantify external stimuli or rely on threshold detection and binary on/off decisions. Here, using fluorescence microscopy and information theory, we analyzed the ability of the transcriptional regulator TetR to sense and quantify the antibiotic tetracycline. The results showed that noise filtering by negative feedback increased information transmission up to 2 bits, generating a graded response able to discriminate different antibiotic concentrations. This response matched the antibiotic subinhibitory selection window, suggesting that information transmission through TetR is optimized to quantify sublethal antibiotic levels. Noise filtering by negative feedback may thus boost the discriminative power of cellular sensors, enabling signal quantification.

INTRODUCTION

Environmental sensing is essential for cell survival, yet noise in signal reception and gene expression constrains the ability of signaling pathways to reliably convey information (1, 2). At the signal reception level, the stochastic nature of diffusion and binding imposes a limit on the ability of cell receptors to quantify the concentration of diffusible molecules (1, 3, 4). At the signal transduction level, gene expression noise generates fluctuations in the copy number of the RNAs and proteins responsible for transducing the signal (5). Signal transduction pathways are thus inherently noisy, and the ability of the cell to control its internal sensory machinery is itself limited (2).

Noisy pathways are subject to an upper limit on the amount of information they can transmit, a quantity known as the channel capacity. Determining the channel capacity of signaling pathways and regulatory networks is thus essential to evaluate the sensing ability and regulatory power of the cell (6). The amount of information encoded by a noisy channel can be quantified in terms of Shannon’s mutual information (MI), and it is usually measured in bits. Recent experiments have shown that signaling pathways exhibit low MI, with channel capacities of approximately 1 bit (7–11). The physiological significance of these observations has been a matter of debate. Because 1 bit is the amount of information required for a binary switch, Cheong et al. (7) proposed that 1-bit information transmission reflected a decision-making process fundamentally limited to on/off responses. Under this interpretation, 1 bit is indicative of a sensor with two possible states, able to detect a signal but unable to quantify it. However, the same amount of information (1 bit) may be transmitted by a graded sensor suffering from an associated inference error (12). Hence, observing 1 bit does not necessarily imply a digital controller. The ultimate question is whether signal transduction mechanisms are accurate enough to quantify external signals or whether molecular noise is so pervasive that the cell has no chance but to rely on threshold detection and on/off responses (Fig. 1A) (13). Answering this question has been complicated by the entangled nature of eukaryotic regulation (14). Here, we investigate information transmission by the prokaryotic transcriptional repressor TetR, a regulator of tetracycline (Tc) resistance, and a widely used workhorse for controlled gene expression in synthetic gene networks.

TetR is the prototype of a large family of one-component signal transduction systems involved in responses against small molecules, such as antibiotics, quorum-sensing molecules, and metabolites. TetR regulates the expression of TetA, a protein conferring Tc resistance. Expression of TetA is deleterious to the host cell; hence, its production is tightly regulated and only occurs when Tc is present in the environment (15). Although the molecular mechanisms involved in TetR regulation have been thoroughly described (Fig. 1B), whether this prototypic circuit responds to the antibiotic in a graded fashion or as an all-or-none switch is currently unknown. Moreover, in recent years, it has become clear that subinhibitory antibiotic concentrations can exert selective pressure and elicit a variety of cellular responses (16). Determining what range of antibiotic concentrations is actively monitored by the cell, and with how much precision, might help us understand the overall impact of antibiotic pressure on microbial communities.

Using fluorescence microscopy, single-cell analysis, and information theory, we measured information transmission in the TetR sensory-regulatory circuit. We analyzed the influence of network architecture on information transmission and determined the overall ability of the system to quantify and respond to the signal (Tc). Our results show that TetR is able to convey up to 2 bits of information, being able to discriminate antibiotic levels below the minimal inhibitory concentration (MIC). In its natural setting, TetR engages in a negative feedback loop (NFL) that increases information transmission through the signaling pathway, enabling signal quantification.

RESULTS

TetR is a classical representative of one-component systems, the most prevalent pathways for signal transduction in prokaryotes (17). Like other one-component sensors, TetR has a dual role, acting simultaneously as a signal receptor (binding Tc) and as a signal transducer (regulating gene expression). TetR represses the expression of the
Tc resistance protein TetA by binding two operators (tetO1 and tetO2) located in the pA promoter (Fig. 1B). Upon binding to Tc, TetR suffers a conformational change that prevents recognition of its cognate operators. Thus, information about the signal (Tc) is conveyed to pA expression levels by the amount of inducer-free TetR. To quantify the MI between inducer-free TetR and pA expression levels, we needed the output conditional distribution $P(pA|\text{TetR})$, that is, the distribution of pA expression levels produced in response to a gradient of inducer-free TetR. For this purpose, we generated a translational fusion between TetR and the red fluorescent protein mKate2, and a transcriptional fusion between pA and the green fluorescent protein (GFP) (Fig. 1C). This way, we could simultaneously monitor, in single cells, input TetR mKate2 and output pA::GFP levels (Fig. 1C).

TetR mKate2 is functional and does not increase noise in the signaling pathway

The translational fusion TetR mKate2 was fully functional and able to repress transcription from the pA promoter (fig. S2). The functionality of the FP fusion, however, did not guarantee a physiological behavior of the protein, because FP fusions are known to cause distribution artifacts. As shown by Landgraf et al. (18), FPs are “sticky” and may induce the formation of transitory aggregates when fused to other proteins. These aggregates are often functional, yet they decrease the effective number of protein particles that binomially partition between daughter cells during cell division. This increases the segregation error and the noise experienced by downstream processes (18). Because information transmission is hampered by noise, this effect may result in an overall reduction in the observed channel capacity. To check for this possibility, we performed a fluctuation after cell division test (18). Briefly, this test consists in measuring the fluctuations exhibited by daughter cells, immediately after cell division, in a downstream process affected by the FP fusion (fig. S2). In our case, we compared the fluorescence produced by pA::GFP between daughter cells. We compared the results between strains that contained TetR mKate2 (RRG112) and cells containing wt TetR (RRG28). As shown in fig. S2, we did not detect any substantial increase in pA::GFP fluctuations due to the mKate2 fusion. We concluded, therefore, that the translational fusion was functional and did not cause any apparent increase in the overall noise levels of the signaling pathway.

Measuring MI between TetR mKate2 and pA

Measuring the MI between TetR mKate2 and its target promoter pA required sampling pA responses against the widest possible range of TetR mKate2 inputs (19). To this end, we cloned tetR mKate2 under the control of the inducible promoter pBAD (Fig. 1) (20). This allowed us to generate a continuous input gradient, which spanned approximately 100-fold (figs. S3 and S4). To determine whether the plasmidic or chromosomal location of the circuit had any effect on information transmission, we cloned our experimental system in three different genomic settings (Fig. 1, D to F). Because plasmid copy number is also subject to random fluctuations (21), different genomic settings allowed us to determine the impact of different noise regimes.
Fig. 2. Extracting MI values from scatterplot data. Measuring entropy and MI from scatterplot data. (A) To measure MI, scatterplot data showing input and output values (upper left panel) must be transformed into a two-dimensional probability density plot (upper right panel). To do so, we used Grenander’s method of sieves, as discussed in Supplementary Results. The middle and bottom panels of the figure show the application of two sieves of different bin sizes. The middle panel shows a sieve with a low number of bins, while the bottom panel shows a sieve with a high number of bins. On the right of the figure, the probability density maps result from applying different sieves on the same scatterplot map. (B) Scatterplot of 9000, log-distributed, random input/output pairs used to test the performance of discretization methods. Because this dataset was randomly generated, discretization should result into 0 bits of MI. (C) MI values extracted from scatterplots from (B) using the discretization methods indicated in the figure. Blue, direct \( H_{\text{ML}} \) estimation; green, direct \( H_{\text{ML}} \) estimation under jackknife correction; red, \( H_{\text{CML}} \) estimation (maximum likelihood with Miller’s correction). As shown in the figure, Miller’s correction was required to prevent information inflation due to increasing bin number. (D) Datasets of different sizes generated to test the effect of undersampling on MI estimation. From the experimental dataset shown in Fig. 1D, we randomly drew 2000, 1000, 500, and 250 cells, generating the four datasets shown in the figure. (E) Effect of sampling size on MI estimation with and without jackknife correction. On the left panel, lines indicate the MI estimation for different bin sizes (x axis), retrieved when applying \( H_{\text{CML}} \) estimation with (green lines) and without (blue lines) jackknife correction. On the right panel, MI estimations for datasets shown in (D) with (green bars) and without (blue bars) jackknife correction. As detailed in Supplementary Results, jackknife correction provided only a marginal improvement of less than 1% of the MI.
on TetR (fig. S5). A gradient of TetR\textsubscript{mkate2} concentrations was then generated by differentially inducing pBAD, and input and output levels were measured by fluorescence microscopy (Fig. 1, D to F).

Determining MI between TetR\textsubscript{mkate2} and pA also required transforming input/output scatterplots into probability density maps \[P(\text{input},\text{output})\]. For that purpose, a mesh of variable size was applied to bin experimental data (Fig. 2A). Probabilities were then obtained using a maximum likelihood estimator \([H_{ML}]\) \((22, 23)\). As shown in Fig. 2A, decreasing the bin size should, in theory, improve our estimation of \(P(\text{input},\text{output})\). However, this procedure is known to artificially inflate MI as the number of bins is increased. To avoid this problem, we introduced Miller’s correction in our maximum likelihood estimator \([H_{CML}]\) \((22, 23)\). We checked that this procedure prevented artifactual MI inflation by applying it to a random distribution with a domain equivalent to that of our experimental results (Fig. 2B). This random distribution should show 0 bits of MI, but direct \(H_{ML}\) estimation generated MI inflation (Fig. 2C). In contrast, \(H_{CML}\) correction prevented this problem, yielding the expected MI. When applied to experimental data, \(H_{CML}\) approaches an asymptotic maximum that is taken as the true MI (Fig. 2E) \((9)\). As described by Hansen and O’Shea \((9)\), the error in MI estimation can be obtained from the SD of MI values in the asymptotic region. All our MI measurements showed SDs between 0.01 and 0.03 bits. Another common problem in determining MI from experimental data arises from undersampling, which reduces our ability to correctly assign probabilities to each bin. Jackknife resampling can be used to correct for this problem \((7, 9, 23)\). However, our experimental sample sizes (>2000 cells in all cases) were well beyond the undersampled regime (Fig. 2D). A complete description of the methods used for MI estimation can be found in Supplementary Results.

Input/output scatterplots were transformed into probability density maps, and from these maps, MI values were obtained as described before. The results showed MI values close to 1 bit, regardless of genomic location (Fig. 3, A to C). Because the input distribution obtained from pBAD was not uniform across the overall induction range, we analyzed the output distribution (pA::GFP levels) under equiprobable inputs. For this purpose, from the datasets shown in Fig. 1 (D to F), we computationally generated uniform TetR\textsubscript{mkate2} distributions (Fig. 3, D to F). The resulting MI under equiprobable inputs \((\text{MI}_U)\) yielded 0.980 ± 0.02 bits for the one-plasmid system (RRS113), 1.06 ± 0.01 bits for the two-plasmid system (RRS112), and 0.88 ± 0.01 bits for the chromosomal insertion (RRS247). Thus,

![Fig. 3. Determining the MI between TetR\textsubscript{mkate2} and pA. (A to C) MI values (top) and density maps (bottom) for the different genomic constructions. Input/output scatterplots were transformed into density plots by applying a mesh of variable bin sizes, as described in the main text. Upper chart shows MI values (y axis) as a function of the number of bins (x axis). Asymptotic MI values were taken as the lower bound of the MI. Density maps correspond to bins sizes yielding maximum MI. (D to F) Probability density maps for equiprobable inputs. Upper charts shows TetR\textsubscript{mkate} input distributions. Lower charts show joint probability density maps \(P(\text{TetR}\textsubscript{mkate}, \text{pA}\textcolon\text{GFP})\). Probability densities follow the color bars on the right.](image-url)
the results indicated that overall MI\textsubscript{Tc} levels were similar for the three genomic locations (Fig. 3, D to F). It is noteworthy, however, that the chromosomal construction displayed slightly lower MI\textsubscript{Tc}, despite showing lower output noise levels (fig. S5). This can be explained by the smaller TetR\textsubscript{mkate2} induction range exhibited by this construction (compare x axis in Fig. 3). This difference in induction ranges is most likely due to the higher copy number achieved by plasmid constructions (which contain pSC101 replication origins) compared to the chromosome (21).

Once we obtained \( P(\text{input, output}) \) density maps, we could assess whether pA responses corresponded to an on/off binary switch or, alternatively, to an analog sensor able to quantify the signal. Judging from MI\textsubscript{Tc} values and the probability density maps shown in Fig. 3, the answer was actually neither. Despite conveying 1-bit MI, the response could not be characterized as a pure on/off switch, because intermediate TetR\textsubscript{mkate2} concentrations generated intermediate pA levels. These intermediate values, however, were so noisy that the ability of the sensor to quantify the signal was severely compromised. Just moving 15% away from the on/off saturating pA values, MI\textsubscript{Tc} levels in RRS113 dropped from 1.06 ± 0.01 bits to 0.28 ± 0.03 bits. Thus, the results indicated that the TetR/pA circuit could not be categorized as a strict digital switch or an effective analog sensor.

**Measuring MI between Tc, TetR\textsubscript{mkate2}, and pA**

A gradient of TetR, artificially induced from the pBAD promoter, was able to convey 1 bit of information to its target promoter. Total TetR concentration, however, is not the natural input of the TetR/TetA circuit. Rather, the inducer of the system, Tc, binds TetR, inactivating it and preventing repression of pA. Therefore, in its natural setting, pA responds to changes in the levels of free TetR rather than to changes in its total concentration. Although the same TetR gradient can be generated by Tc inactivation or by TetR synthesis/degradation, theory predicted that each of these mechanisms would exhibit different noise levels (Supplementary Calculations). Noise constrains information transmission (2); thus, MI values may depend on the way the input gradient is generated. To test this possibility, we measured MI\textsubscript{Tc} between Tc (input) and pA::GFP (output). To this end, TetR\textsubscript{mkate2} was induced to saturating levels in RRS113 cells. These cells were then subjected to a Tc gradient, imaged, and analyzed as described in Materials and Methods. From the \( P(\text{input, output}) \) density maps, we could assess whether pR responses corresponded to an on/off binary switch or, alternatively, to an analog sensor able to quantify the signal. Judging from MI\textsubscript{Tc} values and the probability density maps shown in Fig. 3, the answer was actually neither. Despite conveying 1-bit MI, the response could not be characterized as a pure on/off switch, because intermediate TetR\textsubscript{mkate2} concentrations generated intermediate pA levels. These intermediate values, however, were so noisy that the ability of the sensor to quantify the signal was severely compromised. Just moving 15% away from the on/off saturating pA values, MI\textsubscript{Tc} levels in RRS113 dropped from 1.06 ± 0.01 bits to 0.28 ± 0.03 bits. Thus, the results indicated that the TetR/pA circuit could not be categorized as a strict digital switch or an effective analog sensor.

**A negative feedback on TetR increases information transmission**

Tc induction generated lower pA output noise levels, yielding higher MI values. These findings motivated us to study the influence of network architecture, a well-known factor determining noise propagation in transcriptional circuits (10). At its native location in the Tn10 transposon, TetR is transcribed from pR, a promoter overlapping pA and oriented in the opposite direction (Fig. 4E). Expression from pR is repressed by TetR; thus, TetR engages in an NFL. This NFL implies that the effect of Tc on the circuit is twofold. On the one hand, Tc decreases the levels of free TetR, thus reducing its ability to repress pA transcription. On the other hand, lower levels of free TetR also increase pR transcription, increasing total TetR concentration. To analyze the overall impact of this NFL on the ability of TetR to discriminate Tc, we built the experimental system shown in Fig. 4E. In this construction, TetR\textsubscript{mkate2} was cloned under its cognate promoter pR, while GFP remained under the control of pA. E. coli cells carrying this construct (RRS129) were subjected to a gradient of Tc concentrations, and output pA::GFP levels were measured by fluorescence microscopy, as described in Materials and Methods. Input/output scatterplots (Fig. 4, E and I) were transformed into \( P(\text{input, output}) \) density maps, as described before. The results demonstrated that, under the same Tc gradient, the NFL circuit yielded MI\textsubscript{Tc} = 1.96 ± 0.01 bits, while the NFL-free circuit produced 1.54 ± 0.02 bits. This increase in information transmission was accompanied by a substantial decrease in output noise (Fig. 5A). Noise filtering is a well-known property of NFLs, and it has been shown that some degree of feedback tends to improve information transmission in signaling pathways (24, 25). However this noise-filtering power comes at the cost of reducing the system’s sensitivity to the input (26). This trade-off between input sensitivity and noise suppression implies that the adaptive value of NFLs in signal reception cannot be taken for granted. To shed light on this question, we followed Tkáčik et al. (24) and analyzed theoretically how information transmission in a sensory circuit depended on the presence of a negative feedback. This required assuming that the overall circuit behavior can be approximated by a Gaussian channel (10, 19, 24, 27).

Although noise in bacterial transcription and translation is generally non-Gaussian, this approximation has provided insights in qualitative agreement with experimental results (10, 28). An analysis of the theoretical channel capacity of an ideal one-component sensory circuit, with and without feedback, is shown in Supplementary Calculations. The results revealed that a noncooperative negative feedback \((n = 1)\) operating on the sensor (TetR, in this case) achieved higher channel capacities than its unregulated counterpart. Moreover, a noncooperative feedback was also observed to overperform NFLs displaying cooperativity \((n ≥ 2)\). Thus, theory predicted that a negative feedback on TetR, with \( n = 1 \), was optimal for MI transmission.

We then determined the apparent cooperativity of the NFL feedback operating on TetR. For this purpose, we measured the average responses of pR and pA promoters to Tc induction. The average response by pA showed a sigmoidal curve, with apparent cooperativity \( n = 2 \). In contrast, the response from pR was observed to be hyperbolic, with \( n = 1 \). Therefore, TetR engages in a noncooperative feedback, the strategy predicted by theory to yield the best MI transmission (Fig. 5B). The results shown in Fig. 5B were also in full accordance with previous in vitro data showing that (i) TetR affinity for tetO1 is 50% lower than for tetO2 and (ii) repression of pA requires binding to tetO1 and tetO2, while pR responds to tetO1 only (29).

**TetR discriminates Tc concentrations in the subinhibitory range**

Differences in information transmission and dynamic responses by pR and pA promoters translated into different sensing precision levels.
TetR mkate2 levels produced from pR could be divided into two regions (H1 and H2 in Fig. 5C), which discriminated between Tc concentrations above and below 0.01 μg/ml. Meanwhile, GFP levels produced from pA showed four discriminative quartiles, each one corresponding to a different Tc concentration range (Q1 to Q4 in Fig. 5C). Analysis of the impact of growing Tc concentrations on bacterial growth rates revealed a correlation between pA output levels and the antibiotic effect (Fig. 5E). Tc concentrations without detectable effects on bacterial growth did not elicit responses from pR/pA. Increasing concentrations above this threshold, however, pushed cells into higher pR/pA expression levels (Fig. 5D). The results demonstrated that the 2-bit MI displayed by the system allowed the discrimination of four Tc concentration ranges, corresponding to four different levels of antibiotic toxicity. Using rate-distortion methods, we used the conditional output distribution (Fig. 4K) to back-calculate the MI theoretical maximum (Supplementary Results). This theoretical maximum corresponds to the channel capacity of the pathway. The results showed that the TetR/TetA circuit uses 90% of its channel capacity, the system being optimized to discriminate antibiotic concentrations well below the MIC, between 5 and 25 nM (Fig. 5F).

**DISCUSSION**

Information transmission through biochemical networks is essential for cell survival, yet experimental measurements have revealed channel capacities around 1 bit (7, 8, 12). While these values do not imply that cells necessarily rely on binary computation (12), they nevertheless represent low channel capacities, poorly suited for accurate signal quantification (30). Eukaryotes may solve this problem by using alternative signaling strategies, such as encoding information in transcription factor dynamics (8) or relying on collective decision-making (31). In bacteria, however, these strategies have not been
described, and with a few remarkable exceptions (25, 32, 33), the information capabilities of prokaryotic signaling pathways remain largely unexplored. Signal transduction in prokaryotes is mostly undertaken by one-component response regulators, proteins able to bind to specific molecules and modify gene expression accordingly (17). Here, we analyzed the ability of TetR, a canonical one-component regulator, to convey information about its cognate inducer, the antibiotic Tc.

The results showed that the TetR response against Tc was neither intrinsically digital nor analog but was heavily conditioned by the noise levels experienced by the signaling pathway. Noise filtering by negative feedback was key for the system to achieve a channel capacity of 2 bits. While this channel capacity may not represent a marked increase over levels reported for other pathways, it was enough for the cell to discriminate between different antibiotic concentrations. The overall ability of TetR to quantify the signal may seem poor. However, one must take into consideration that sensors and transducing molecules are generally present at low copy numbers and that transcription factors are usually found in the nanomolar range. At these concentrations, noise is expected to severely constrain channel capacities, with upper limits predicted to be around 3 bits (34). While it is possible for the cell to improve information transmission by increasing the concentration of sensors and transducing molecules, this strategy quickly leads to diminishing returns, demanding astronomical investments to boost signal detection (2, 34). It is thus likely that the sensing ability of TetR and other cellular sensors is primarily constrained by cellular economy.

The results showed that an NFL increased information transmission in TetR. Similarly, theory indicated that a noncooperative NFL operating on the sensor optimizes the channel capacity of one-component response regulators. If economy is a critical factor limiting sensing precision, then the advantage of an NFL may be twofold. It not only increases information transmission but also reduces the sensor concentration at the steady state when the signal is not present. An NFL operating on the sensor may thus increase information transmission while simultaneously reducing the metabolic burden imposed by the sensory pathway.

One of the key features of information transmission through the TetR circuit was its ability to discriminate antibiotic concentrations below the MIC (Fig. 5D). Sub-MIC antibiotic levels may not kill bacteria, but this does not imply that they are innocuous. Subinhibitory antibiotic concentrations may damage the cell, decreasing the growth rate of susceptible populations (Fig. 5E). The sub-MIC selection...
window comprises all antibiotic concentrations below the MIC that produce a fitness deficit big enough to promote the evolution of resistant strains (16). In E. coli, the minimal selective concentration for Tc was found to be 15 ng/ml (35). The minimal informative concentration in our experiments was 10 ng/ml. Thus, our results support the idea that TetR-mediated resistance is primarily aimed against sublethal antibiotic doses that directly impair cell growth. In the environment, antibiotics rarely achieve the concentrations used in the clinical setting, yet sub-MIC levels are frequent, as a result of widespread antibiotic contamination (36). Quantifying the signal also makes sense in the context of repeated exposure to subinhibitory levels of antibiotic. Concentrations over the MIC are lethal; thus, their effect on fitness is likely to be close to binary (dead/alive), depending on whether bacteria express resistance. In contrast, subinhibitory levels have an incremental effect on growth rates (Fig. 5E). Because TetA expression is known to be metabolically expensive (15), it may pay off for the cell to invest in a sensory system able to adjust resistance levels to the fitness deficit caused by different sub-MIC antibiotic concentrations. Testing this hypothesis is complicated by the need for precise methods to quantify the fitness distribution, in single cells, under sublethal antibiotic levels. If achievable, however, this analysis may shed light on the elusive relationship between the sensory power of the cell and its overall impact on cellular fitness.

**MATERIALS AND METHODS**

**Bacterial strains**

All bacterial strains used in this study were derivatives of E. coli K-12 and are listed in table S1.

**Plasmid construction**

Plasmid pRRG13 (fig. S1A, upper right side) was generated by inserting the tetA promoter region (pA) in front of the gfp gene in plasmid pUA66. Promoter pA was polymerase chain reaction (PCR) amplified from the Tn10 region of the natural plasmid R100 using oligonucleotides pTetA1XhoI and pTetA2BamHI. The resulting band was inserted in pUA66 using XhoI and BamHI restriction endonucleases. The TetRmkate2 fusion protein was generated by binding the N terminus of protein mKate2 to the C-terminal end of TetR through a flexible linker made of Ser-Gly-Gly-Gly-Gly peptide. For this purpose, we constructed plasmid pRRG54. Plasmid pRRG54 was built by Gibson assembly out of three DNA fragments. The first one was the PCR product of amplifying mkate2 with primers isomkatefusdir and isomkate_rev from plasmid pRAF22. The second one, containing the gfp gene, was amplified from pUA66 using primers pRRG74_GFPdir and pRRG74_GFPrev. Last, the third one contained the Tn10 region that includes tetR, pR, and pA. It was amplified from plasmid R100 using primers pRRG74_p tetArev and pRRG74_TetRrev.

**Strain generation**

Strain RRG112 was generated by transforming plasmids pRRG13 (Km') and pRRG62 (Amp') into E. coli BW27783 by electroporation. Strains RRS113 and RRS129 were generated by transforming plasmids pRRG63 (Km', Amp') and pRRG74 (Amp'), respectively, into E. coli BW27783 by electroporation. Strain RRS247 was generated by inserting the region comprising the araC to km genes (both included) from plasmid pRRG63 into E. coli chromosome. This way, strain RRS247 is the chromosomal counterpart of strain RRS113 (fig. S1A). The DNA fragment was PCR amplified from RRS113 using primers AraC_Wanner and Km_AraD_Wanner. These primers contained a homologous region to the ara operon such that the amplified fragment could be recombined into E. coli TB10 strain following the protocol described in (38). A P1 lysate was then prepared and transduced to strain BW27783 to yield strain RRS247.

**Culture conditions**

Unless otherwise stated, bacterial growth for DNA extraction and strain propagation was performed in LB, and cells were grown at 37°C, with orbital shaking and supplemented with appropriate antibiotic concentrations. Antibiotic concentrations used were as follows: ampicillin (Amp; 100 μg/ml), chloramphenicol (Cm; 25 μg/ml), and kanamycin (Km; 50 μg/ml). Arabinose induction for pBAD-regulated constructions used the concentrations shown in Supplementary Results and fig. S4, following the protocol described in (39).

**DNA purification**

Plasmid DNA was purified using the GeneJet Plasmid Miniprep Kit, following the vendor’s instructions. Total DNA was extracted using Bio-Rad's InstaGene Matrix, following the vendor’s instructions.

**Growth rate determination**

Growth rates were determined by measuring the absorbance in a Victor3 (PerkinElmer) microplate reader. Cells were grown in M9 medium supplemented with casamino acids [0.2% (w/v)] as nitrogen source and glycerol or glucose (0.5%) as carbon source. Cells were pregrown in flasks at 37°C overnight and then subjected to a 1:1000-fold dilution in fresh medium. A total of 150 μl of these diluted cultures were added to the wells of a 96-well microtiter plate (Deltalab). Absorbance values at 600 nm were taken every 7 min. These absorbance values were background subtracted and transformed into OD600 (optical density at 600 nm) values by using a calibrating curve obtained with a Shimadzu UV-1603 spectrophotometer. Growth rates
Cells were grown at 37°C in M9 medium (Fluka-Sigma M9 salts supplemented with 100 μM CaCl2, 20 mM MgSO4, 0.2% casamino acids, and 0.5% glucose or glycerol). Appropriate concentrations of arabinose/glucose/glycerol/Tc were added according to the experimental requirements. Overnight saturated cultures were diluted 1:1000 in the same medium and incubated at 37°C for 2 to 3 hours until they reached OD600 = 0.1 to 0.2. Thus, cells were grown for approximately 16 generations in the same growth medium before their analysis by fluorescence microscopy. A total of 2 μl of cells were placed onto agarose pads containing the same growth medium used for preimaging growth. Agarose pads were generated by stacking an adhesive frame (Frame-Seal Incubation Chamber, Bio-Rad) onto a microscope slide. In the cavity formed by the adhesive frame, a volume of 200 μl of hot M9 medium + 1.5% agarose was added. Another microscope slide was placed over the frame immediately after pouring M9 + agarose to obtain a uniform and leveled gel. M9 medium used to generate agar pads was supplemented with the appropriate arabinose/glucose/glycerol/Tc concentrations. Pads were allowed to cool down for 30 min at room temperature, and then, using a sterile scalpel, a square of about 0.25 cm² of the agarose gel was cut. The rest of the agarose pad was removed, and 2 μl of the appropriate culture was placed on top of the pad. The culture droplet was allowed to dry for approximately 5 min, and then, a coverslip was placed over the frame, sealing it carefully. Sealed pads were transferred to the slide holder of a Leica AF6500 inverted epifluorescence microscope, inside an environmental chamber that was kept at 37°C along the course of the experiment. Cells were allowed to adapt to the pad and temperature for 30 min before we started to image. Images were acquired with ×630 magnification using HCX PL S-APo 63×1.3 oil objective. We acquired images in phase contrast and in the green and red fluorescence channels. Filters used for fluorescence images were 562/40-nm excitation and 641/75-nm emission for mKate2 and 482/18-nm excitation and 520/28-nm emission for GFP. A Leica EL6000 external light engine, equipped with a mercury vapor lamp (HXP Short Arc Lamp, Osram), was used for fluorescence excitation. All images were obtained using the excitation lamp at maximum power. Images were acquired using a 12-bit Andor iXon885 high-speed camera, without binning. For red fluorescence images, we used two exposure times (500 ms and 2 s) in all images taken. For the green fluorescence channel, variable exposure times were applied (10 ms, 100 ms, and 1 s). To avoid fluorescence bleaching from previous exposures, snapshots were taken at least four fields of view away from each other. Linearity of the fluorescence emission with exposure time was checked using a set of predefined fluorescence beads (Rainbow Fluorescent Particle Slide, Spherotech) (fig. S1B).

Supplementary materials for this article is available at http://advances.sciencemag.org/cgi/content/full/4/11/eaat5771/DC1

Supplementary Results

Supplementary Calculations

Fig. S1. Genetic constructions used in this work and signal calibration.

Fig. S2. Validation of TetR<sub>max</sub>, translational fusion.

Fig. S3. Induction of TetR<sub>max</sub> from pBAD.

Fig. S4. TetR<sub>max</sub>, induction histograms are gamma distributed.

Fig. S5. Average responses and noise levels in arabinose-inducible constructions.

Fig. S6. Output distributions and MI for equiprobable, arabinose-induced inputs.

Fig. S7. Influence of feedback cooperativity on MI.

Table S1. Bacterial strains and bacteriophages used in this work.

Table S2. Oligonucleotides used in this work.

Table S3. Bacterial plasmids used in this work.

Table S4. List of reagents used in this work.

Table S5. Image data depository.

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