Supplementary Materials for

The Innate Growth Bistability and Fitness Landscapes of Antibiotic-Resistant Bacteria

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Published 29 November 2013, Science 342, 1237435 (2013)
DOI: 10.1126/science.1237435

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for

The innate growth bistability and fitness landscapes of antibiotic resistant bacteria

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Supplementary Materials

SUPPLEMENTARY METHODS

Strain construction

All of the strains used in this study are derived from *Escherichia coli* K12 strain MG1655 and described in table S1. The region containing *lacI*, *lacZ* and *lacY* genes was deleted using the method of Datsenko and Wanner (71). Construction of strain Lac2 (also known as EQ37, in which *lacI* and *lacY* were deleted and *lacZ* is driven by P_{Ltet-O1}) is described in Scott *et al* (16). DNA oligos used for construct and strain preparation are described in table S4. For all experiments, strains stocks were stored at -80 °C. Plates were streaked with stock strains within two weeks of use and stored at 4 °C.

Construction of a new chromosomal *lacZ* reporter strain. The synthetic and Crp-independent P_{Llac-O1} promoter (-60 to +47, relative to the transcriptional start site) was amplified from pZE12 (72) and substituted for P_{Ltet-O1} in pKD13_{Ltet-O1} (16), yielding pKD13_{Llac-O1}. Using the method of Datsenko and Wanner (71), the promoter plus the upstream Km\(^r\) gene (Km\(^r\):P_{Llac-O1}) in pKD13- P_{Llac-O1} was integrated into the chromosome to replace the *lacI* gene and the native *lac* promoter (including the 5’ UTR of *lacZ*) of MG1655 deleted for *lacY* (57). The resultant strain, in which *lacZ* is driven by P_{Llac-O1} and both *lacI* and *lacY* are deleted, is named as EQ5.

Chromosomal incorporation of resistance markers. The P_{Llac-O1} promoter described above was substituted for P_{Ltet-O1} in pKDT_{Ltet-O1} (57), yielding pKDT_{Llac-O1}. The cat structural gene, encoding the protein responsible for chloramphenicol resistance, was amplified from pZA31 (72) using forward primer Cat-Kpn-F and reverse Cat-Bam-R. The PCR products were gel purified, digested with KpnI and BamHI, then ligated into the same sites of pKDT_P_{Llac-O1}, yielding pKDT_P_{Llac-O1}-cat. Present in this plasmid, the construct Km\(^r\):rrnBT:P_{Llac-O1}-cat was amplified using primers Placcat-P1 and Placcat-P2. The Placcat-P1 contains a 51 bp region that is homologous to the intergenic region (272\(^{nd}\) to 222\(^{nd}\) nucleotides relative to the start codon of *ycaD*) between *ycaC* and *ycaD* while Placcat-P2 contains a 50 bp region that is reverse complemented to the *ycaC/ycaD* intergenic region located between 116\(^{th}\) and 67\(^{th}\) nucleotides relative to the start codon of *ycaD*. The PCR products were gel purified and then electroporated into MG1655 cells that carried pKD46 (71) expressing the λ-Red recombinase. The cells were incubated with shaking at 37 °C for 1 hour and then applied onto LB + Km agar plates. The plates were incubated at 30 °C overnight. The Km\(^r\) colonies were verified for the substitution for the *ycaC/ycaD* intergenic region between 222\(^{nd}\) and 116\(^{st}\) nucleotides relative to the start codon of *ycaD* by colony PCR and subsequently by sequencing. The Km\(^r\):rrnBT:P_{Llac-O1}-cat construct was moved to background strain EQ4 with P1 phage transduction (73), and colonies were selected for resistance to Km and Cm on agar plates.
The Km\textsuperscript{r} gene was flipped out using the standard method \((71)\). The resultant strain is named EQ75, also known as Cat1 in the main text.

To make chromosomal cat driven by P\textsubscript{Ltet-O1} \((72)\), the same cat gene as above was cloned into \textit{KpnI/BamHI} sites downstream of P\textsubscript{Ltet-O1} in pKDT\_P\textsubscript{Ltet-O1} \((57)\), yielding pKDT\_P\textsubscript{Ltet-O1}\_cat. The cassette “Km\textsuperscript{r}:rrnBT:P\textsubscript{Llac-O1}\_cat” from this plasmid was incorporated into the same \textit{ycaC/ycaD} site as described above. This strain is named EQ92, also referred to as Cat2 in the main text.

As expected, the cell with the chromosomal P\textsubscript{Llac-O1}\_cat was highly resistant to Cm. To vary Cm resistance levels using site-directed mutagenesis, we modified the region containing the ribosome binding site (RBS) in P\textsubscript{Llac-O1} by inserting 4 nucleotides (TCCT) immediately downstream of RBS or/and changing one or two nucleotides in RBS (see table S3). The modified P\textsubscript{Llac-O1} promoters (containing various altered RBS) together with cat were moved to the chromosome at the same location as described above. The resulting strains are referred to as Cat3 through Cat6.

To make the chromosomal tetA driven by P\textsubscript{Llac-O1}, the tetA gene was cloned from the transposon Tn10 (also known as tetA(B) \((74)\)) and subsequently substituted for cat in pKDT\_P\textsubscript{Llac-O1}\_cat (see above), yielding pKDT\_P\textsubscript{Llac-O1}\_tetA. Similarly as described above, The cassette “Km\textsuperscript{r}:rrnBT:P\textsubscript{Llac-O1}\_tetA” was integrated into the \textit{ycaC/ycaD} locus.

**Chromosomal deletion of motA and chromosomal integration of P\textsubscript{Ltet-O1}\_gfp.** The motA gene deletion in strain JW1879-2 \((E.\ coli\ Genetic\ Stock\ Center,\ Yale\ Univ.)\), in which a Km resistance (Km\textsuperscript{r}) gene is substituted for the motA gene, was transferred to EQ4 and Cat1 by P1 transduction \((73)\), and Km\textsuperscript{r} was then removed as described above to yield EQ4m and Cat1m (table S1).

To make the chromosomal GFP driven by P\textsubscript{Ltet-O1}, a \textit{gfpmut3b} structural gene \((75)\) was cloned downstream of P\textsubscript{Ltet-O1} in pKD13\_P\textsubscript{Ltet-O1} \((57)\), yielding pKD13P\textsubscript{Ltet-O1}\_gfp. The construct “Km\textsuperscript{r}:rrnBT:P\textsubscript{Llac-O1}\_gfp” present in this plasmid was integrated into the \textit{intS/yfdG} intergenic region of the MG1655 chromosome as previously described \((76)\). The “Km\textsuperscript{r}:rrnBT: P\textsubscript{Llac-O1}\_gfp” construct was then P1-transduced into the Cat1m background to create strain GCat1m.

**Microfluidic device fabrication**
Microfluidic devices were fabricated by molding silicone elastomer (PDMS Slygard 184, Dow Corning) to the master molds \((28)\), consisted of two layers of cross-linked epoxy (Su-8s, MicroChem) patterned by negative phototransparency masks (FineLine Imaging, CO) on silicon wafers (fig. S4A). The layer for growth chambers (~1.5 µm) was deposited first using SU-8 2002 and processed according to the manufacturer’s manual. On top of the first layer, the second layer for channels (~25 µm) was made using SU-8 2025 (side view).
PDMS was mixed 1:15 ratio of catalyst and resin, poured to the master mold, degassed for 30 min and cured in an 80 °C oven for 90 min. After the elastomer was peeled off the mold, inlet and outlet holes were punched. Then, the device was boiled in 1% HCl solution for 1 hour, attached to No.1 microscope cover glass (pre-cleaned with ethanol) and baked in the 80 °C oven overnight for maximum binding.

**CAT assays**

We used two separate CAT assays in this study, the radiometric assay and the spectrophotometric CAT assay as described below. The spectrophotometric assay was used to obtain an estimate of the absolute CAT activity per OD₆₀₀ in strain Cat1 (table S2) and Cat2, and to quantify the CAT activities of strain Cat1 when growth was inhibited by tetracycline (Fig. 3C). We used the radiometric assay (performed on Cat1 and Cat3-Cat6) to determine the CAT activities of strains Cat3-Cat6 relative to Cat1. Relative CAT activities of all strains Cat1-Cat6 are given in the bottom of Fig. 4B.

**Sample collection and cell permeabilization for CAT assays.** We collected samples (at OD₆₀₀ ~ 0.15 – 0.4) from cultures growing in minimal medium after at least 3 generations of exponential growth and stored them at -80 ºC for assays the following day. After thawing samples we added an equal amount of extraction buffer (0.2 M Tris, pH 8.0, 10 mM EDTA, and for the radiometric assay, an additional 40 mM β-mercaptoethanol). We then added toluene to 1% by volume, vortexed vigorously for 15 seconds (or 10 seconds for radiometric assay) and allowed cultures to incubate at room temperature for 15 minutes before performing the assays as described below (either spectrophotometric or radiometric). To identify the range in our assay over which CAT activity was proportional to CAT protein levels, we diluted samples with various volumes of permeabilized wild type cells (EQ4) and assayed activity. CAT activities reported in this study are within the linear dose-response window as identified above.

**Spectrophotometric CAT assays.** We used essentially the method of Shaw (77). The assay was performed by aspirating 15 µL permeabilized sample into 135 µL reaction buffer inside the cuvette of a Thermo Scientific Genesys 20 spectrophotometer preheated to 37 ºC. The standard reaction buffer contained 150 µL of 4 mg/ml 5,5'-dithiobis-(2-nitro- benzoic acid) suspended in Tris/HCl, pH 8.0, 60 µL each of 20 mM Acetyl-CoA and 5 mM Cm, and 1.08 mL deionized H₂O. We recorded absorbance A₄₁₂ every 6 seconds using VISIONlite software (Thermo Scientific) between minutes 4-8 of the reaction (absorbance increased linearly during the reaction for at least 10 minutes). Reported activity was taken as the slope of the absorbance over time (minus the slope of a blank sample lacking CAT), divided by the final OD₆₀₀ of cells in the reaction volume (U/OD₆₀₀). To obtain the in vitro activity estimate for strain Cat1 (table S2), we performed a total of 7 assays from samples collected from two replicate cultures on different days with fresh media and reagents to obtain a value repeatable to within ~10%.

**Radiometric CAT assays.** Radiometric CAT assay was performed essentially as described by Seed and Sheen (78). Briefly, after samples thawed they were diluted to
OD_{600} = 0.32 (Bio-Rad SmartSpec Plus) by the medium used for the culture. The assay was performed by adding 50 µL permeabilized sample to 75 µL reaction mixture preincubated at 37 °C for 5 min to give final concentrations of 100 mM Tris/HCl, pH8.0, 100 µM [\textsuperscript{3}H] chloramphenicol (25 µCi/µmol), 240 µM butyryl CoA, 1% (vol/vol) ethanol. After incubation at 37°C for 30 min, the reaction was terminated by addition of 250 µL xylene, and mixed vigorously by vortexing. After centrifugation, 80% of the xylene phase was extracted twice with an equal volume of 10 mM Tris/HCl, pH7.5, 1 mM EDTA. The remaining organic phase was counted in a scintillation counter.

### β-Galactosidase assay

Chloramphenicol-sensitive strains Lac1 (EQ5) and Lac2 (EQ37) express LacZ constitutively from the chromosome (table S1). We grew these strains in M63 minimal media + 0.4% glucose in presence and absence of Cm inside 20 mm diameter glass tubes, as described above in the section “batch culture growth.” Experimental culture tubes contained up to 10 µM chloramphenicol to retard growth. During exponential growth (with and without Cm), four to six samples (~200 µL) were collected from each growing culture (OD_{600} = 0.1 to ~0.8), fast frozen on dry ice and stored at -80°C prior to the assay. β-Galactosidase activity was measured at 37°C by the traditional Miller method (79). The LacZ activity per volume of cell culture (OD) was obtained as the slope of the plot of LacZ activity levels (ΔOD_{420}/min/ml) versus OD_{600} to give Miller Units used in Fig. 3C.
Supplementary Discussion
THEORETICAL ANALYSIS

We present here in detail our mathematical model for antibiotic-resistant growth, including the origins of Eqs. [1]-[4] which define the model in the main text, and describe how growth bistability and plateau-shaped fitness landscapes emerge from these equations. In doing so we outline the details upon which these phenomena depend, in both a qualitative and quantitative sense (e.g., the dependence of MIC and MCC predicted by physical parameters in the model). A key concept in the model is the "growth-mediated feedback" that arises due to global effects of cellular growth state on gene expression. In particular, the feedback describes the way that growth rate affects expression of proteins offering antibiotic resistance.

We will not restrict our discussion to the behavior of a specific antibiotic or resistance enzyme, but we will apply the general model to the specific antibiotics and respective resistance mechanisms used in our experiments (chloramphenicol, tetracycline, and minocycline). We will examine translation inhibiting antibiotics and resistance conferred by enzymes that modify the antibiotic (inactivation), or pump the drug out of the cell and away from its target (efflux). We will begin with the simplest equations that realize our basic assumptions and then add complexity to these models when necessary to better describe physiological conditions in experiments; in many cases, we will see that the simplest descriptions capture the experimental findings without further elaboration. Whenever possible, we will also provide independent estimates of the parameters used in the model.

1. Modeling cell growth in the presence of antibiotics, for both antibiotic-sensitive and resistant cells

1.1 Effects of antibiotics on translation rate

Whether bacteriostatic or bactericidal, sub-inhibitory concentrations of translation-inhibiting antibiotics often allow exponential growth in bacteria, albeit at reduced rates (80–82). We concern ourselves here with this simple exponential growth, and model the effect of the antibiotic as reducing the average translation rate $\gamma$, due to the direct binding between the ribosome ($R_b$) and antibiotic ($a$) (22). We assume that the concentration of antibiotic-bound ribosomes $[R_b \cdot a]$ reaches equilibrium rapidly, characterized by a dissociation constant, $K_D$. Then the fraction of unaffected ribosomes is

$$\frac{[R_b]}{[R_b]_{\text{total}}} = \frac{1}{1+[a]_{\text{int}} / K_D}, \quad \text{[S1]}$$

where $[R_b]$ and $[R_b]_{\text{total}} \equiv [R_b] + [a \cdot R_b]$ are respectively the unaffected and total
ribosome concentration, and \([a]_{\text{int}}\) is the internal antibiotic concentration. Here we have described equilibrium antibiotic-ribosome binding without cooperativity for simplicity, but this form is in fact the correct one for the antibiotics used in this work Cm, Tc, and Mn (22, 83). Assuming that the drug-bound ribosomes do not translate at all, the total translation rate in a cell is given by \(\gamma_0 \cdot [Rb] = \gamma \cdot [Rb]_{\text{total}}\), where

\[
\gamma \equiv \frac{\gamma_0}{1 + [a]_{\text{int}} / K_D}
\]

[S2]

is referred to as the average translation rate, and \(\gamma_0\) is the translation rate in the absence of drug. (NB, here and throughout this work, a subscript of "0" indicates a value in the absence of drug).

1.2 The relation between translation rate and growth rate, and the emergence of the half-inhibition concentration

Under steady exponential growth, recent work by Scott et al. (16) has established a Monod-like expression for the growth rate,

\[
\lambda = \lambda_{\text{max}} \frac{v}{\gamma_0 + v}
\]

[S3]

where \(\gamma\) is the drug-dependent average translation rate as defined in Eq. [S2], \(v\) is related to the rate of nutrient utilization and is fixed for a given saturating nutrient medium, and \(\lambda_{\text{max}}\) is the maximum growth rate achievable with a theoretically best possible nutrient (see fig. S9). In drug-free medium, the above becomes

\[
\lambda_0 = \lambda_{0\text{max}} \frac{v}{\gamma_0 + v}
\]

[S4]

where \(\lambda_{0\text{max}} \approx 2.85 \text{ hr}^{-1}\) (16). Taking the ratio of Eqs. [S3] and [S4], and using Eq. [S2] for \(\gamma / \gamma_0\) yields

\[
\lambda = \frac{\lambda_0}{1 + [a]_{\text{int}} / I_{50}}
\]

[S5]

where

\[
I_{50} = K_D \cdot \lambda_{0\text{max}} / \lambda_0,
\]

[S6]

is the predicted half-inhibition concentration, the internal antibiotic concentration that reduces growth rate by 50%.

Reports of in vitro measurements of \(K_D\) for Cm-ribosome binding in \(E. coli\) range from 0.6 to 2.7 \(\mu\text{M}\) (22, 84). For wild-type \(E. coli\) (EQ4) cells growing in glucose minimal media, \(\lambda_0 = 0.67 \text{ hr}^{-1}\). Eq. [S6] thus predicts \(I_{50}\) to be of several micromolar \((2.8 \leq [I_{50}] \leq 12 \mu\text{M})\) during Cm-inhibited growth, as long as the dominant mode of
action for the antibiotic is to inhibit translation via ribosomal binding.

1.3 The relation between intra- and extra- cellular antibiotic concentrations in the absence antibiotic resistance expression

Inverting Eq. [S5] leads to the result that the relative doubling time \( \frac{\tau}{\tau_0} = \frac{\lambda_0}{\lambda} \) increases linearly with the drug concentration (see Eq. [4] in Fig. 3 of the main text). The observed linear relationship in Fig. 3D is consistent with this prediction if the antibiotic concentration in the medium, denoted by \([a]_{\text{ext}}\), is proportional to \([a]_{\text{int}}\) over some range. The latter is expected generally if antibiotic enters the cell via passive diffusion, with an influx rate

\[
J_{\text{influx}} = \kappa \cdot ([a]_{\text{ext}} - [a]_{\text{int}})
\]

where \(\kappa\) is the membrane permeability, and exits the cell via native low-affinity efflux

\[
J_{\text{efflux}} \approx V_{\text{native}}^{\text{max}} \cdot [a]_{\text{int}} / K_{\text{native}}
\]

assuming Michaelis kinetics for efflux. Note that low-affinity efflux is common in antibiotic-susceptible cells and this endogenous efflux is not associated with clinically relevant antibiotic resistance (41). The passive uptake of antibiotics into cells is well-established for Cm (41), Tc (85), and many others (86). Balancing the influx and efflux in the steady state (and neglecting dilution due to cell growth1), we obtain the linear relation

\[
[a]_{\text{ext}} = [a]_{\text{int}} \cdot \left(1 + \frac{V_{\text{native}}^{\text{max}}}{\kappa \cdot K_{\text{native}}}\right),
\]

which leads to an effective half-inhibition concentration \( I_{50} \equiv I_{50} \cdot \left(1 + \frac{V_{\text{native}}^{\text{max}}}{\kappa \cdot K_{\text{native}}}\right) \), describing external antibiotic concentration when growth rate is reduced by half. Since endogenous efflux of antibiotics is weak for cells grown in minimal media (87), we expect \( V_{\text{native}}^{\text{max}} / (\kappa \cdot K_{\text{native}}) < 1 \) for Cm (41) and tetracyclines (87, 88) in our experiments. The data of Fig. 3D give a value of 5.5 µM Cm for the half-inhibition concentration. This is in the range estimated above for \( I_{50} \), suggesting that efflux may indeed be negligible for Cm at low concentrations in minimal medium.

1.4 The relation between intra- and extra- cellular antibiotic concentrations in the presence of antibiotic resistance expression

A key qualitative result in this section is captured by Eq. [S12], from which we will see that the relation between intra- and extracellular antibiotic concentration is marked by two distinct regimes in the presence of an enzyme offering antibiotic resistance. We

1 Typical influx rates are much larger than growth rate, at least ~200-fold higher for drugs used in this study (See (88) and table S2)). However, when dilution due to growth is non-negligible, and in the absence of efflux, the dilution may have important consequences for bacterial growth (54).
assume an enzyme $E$ is expressed at a concentration $[E]$ in the steady state, with the effect of reducing the intracellular pool of the active antibiotic. Whether the protein functions as an efflux pump (as in Tc-TetA system) or as enzyme inactivating the drug (as in the Cm-CAT system studied in the main text), we assume the enzymatic process can be simply described by Michaelis-Menten kinetics, with the rate of antibiotic removal given by

$$J_{\text{removal}} = \frac{V_{\text{max}}}{1 + K_m / [a]_{\text{int}}}$$

with affinity $K_m$, and $V_{\text{max}} \equiv k_{\text{cat}} \cdot [E]$. Assuming that the antibiotic enters the cell via passive diffusion (Eq. [S7]), the balance of influx and removal leads to the following relation between the external and internal drug concentrations:

$$[a]_{\text{ext}} = [a]_{\text{int}} + \frac{V_{\text{max}} / \kappa}{1 + K_m / [a]_{\text{int}}}.$$  \[S11\]

This relation is sketched in Fig. 3B, and the origin of the approximate threshold-linear form is motivated in its caption. Mathematically, the two regimes are simply

$$[a]_{\text{int}} \approx \begin{cases} 
[a]_{\text{ext}} / \left(1 + V_{\text{max}} / \kappa K_m\right) & \text{for } [a]_{\text{ext}} \ll V_{\text{max}} / \kappa \\
[a]_{\text{ext}} - V_{\text{max}} / \kappa & \text{for } [a]_{\text{ext}} \gg V_{\text{max}} / \kappa \end{cases}$$

\[S12\]

This threshold-linear behavior, when combined with the growth-mediated feedback as described below, provides the mechanism for bistability in antibiotic resistance. An important dimensionless parameter that arises whenever passive influx is balanced by Michaelis-Menten efflux kinetics is $V_{\text{max}} / (\kappa K_m)$, a term that was previously found useful in describing antibiotic resistance (89), albeit in a different context. Strong effect of drug relief is realized for $V_{\text{max}} / K_m \gg \kappa$, where $[a]_{\text{int}} \ll [a]_{\text{ext}}$ for a broad regime of drug concentrations $[a]_{\text{ext}} \ll V_{\text{max}} / \kappa$.

We mention as an aside that if growth-mediated feedback did not play a role, then Eqs. [S11] and [S5] would determine growth rate. With $V_{\text{max}}$ as a constant, the growth rate is expected to decrease smoothly with increasing external antibiotic concentrations, similar to the response of wild-type cells. We present the results of this no-feedback model fig. S17 (dashed lines) to compare to growth rate data from our CAT-expressing strains. The behavior of the no-feedback model cannot account for the observed abrupt drops in growth rate even at a qualitative level.

1.5 The effect of growth-mediated feedback in cells expressing antibiotic resistance

The above relation between the internal and external drug concentration assumes a constant expression level of the enzyme providing drug resistance, i.e., constant $V_{\text{max}}$, as the external drug concentration is varied. However, sub-inhibitory concentrations of
antibiotics generically lead to altered expression of proteins (12, 13, 16, 17, 43, 90) unless very specific regulatory mechanisms such as negative feedback control are involved (57). In this study, we investigate the unregulated (or “constitutive”) expression of resistance mechanisms, because (i) in many instances the expression of drug resistance is known not to involve specific regulatory mechanisms (25), and (ii) de novo evolution of novel drug resistance mechanisms is likely to proceed without drug-specific regulation.

The expression levels of unregulated proteins have been shown to depend linearly on the growth rate under sub-inhibitory doses of translation-inhibiting antibiotics (16). This is seen in Fig. 3C for constitutively expressed LacZ and CAT under Cm- and Tc- inhibited growth, respectively. The $V_{\text{max}}$ of the enzyme under sub-inhibitory growth can hence be written in terms of the growth rate $\lambda$, and the $V_{\text{max}}$ and growth rate in the absence of drug, $V_0$ and $\lambda_0$ respectively, as

$$V_{\text{max}} = V_0 \cdot \frac{\lambda}{\lambda_0} \quad \text{[S13]}$$

Combining Eq. [S13] with [S11] then gives the full expression relating $[a]_{\text{ext}}$ and $[a]_{\text{int}}$ at different growth rates $\lambda$:

$$[a]_{\text{ext}} = [a]_{\text{int}} + \frac{\lambda}{\lambda_0} \cdot \frac{V_0}{\kappa} \cdot \frac{V_0 / \kappa}{1 + K_m / [a]_{\text{int}}}.$$ \[S14\]

Eliminating $\lambda / \lambda_0$ using Eq. [S5] gives

$$[a]_{\text{ext}} = [a]_{\text{int}} + \frac{V_0}{\kappa} \cdot \frac{1}{1 + [a]_{\text{int}} / I_{50}} \cdot \frac{1}{1 + K_m / [a]_{\text{int}}}.$$ \[S15\]

Eq. [S15] can be inverted to predict the internal drug concentration for every external drug concentration, given the chemical properties of the drug ($K_m$, $\kappa$, and $I_{50}$), and the drug resistance expression system that defines $V_0$. The value of $[a]_{\text{int}}$ can subsequently be used (via Eq. [S5]) to predict the relative growth rate ($\lambda / \lambda_0$) for the given external drug concentration.

2. Model predictions and their sensitivity to biophysical parameters

Since the relative growth rate ($\lambda / \lambda_0$) only depends on $[a]_{\text{int}} / I_{50}$, and $I_{50}$ itself does not depend on the concentrations of the drug or drug resistance enzyme (see Eq. [S6]), we shall rewrite Eq. [S15] in terms of $a_i = [a]_{\text{int}} / I_{50}$ and $a_e = [a]_{\text{ext}} / I_{50}$, with

$$a_e = a_i \cdot \left(1 + \frac{1}{1 + a_i} \cdot \frac{\rho}{1 + \sigma \cdot a_i}\right) \quad \text{[S16]}$$
Here, we have introduced two dimensionless parameters: the 'resistance efficacy'

$$\rho \equiv \frac{V_0}{(\kappa \cdot K_m)}$$  \[S17\]

which indicates the efficiency of the drug-resistance enzyme in unit of drug influx, and

the 'saturability'

$$\sigma \equiv \frac{I_{50}}{K_m}$$  \[S18\]

which characterizes the extent to which the activity of the drug-resistance enzyme is saturated at growth-inhibiting concentrations of the antibiotic. The relative growth rate,

$$\frac{\lambda}{\lambda_0} = \frac{1}{1+a_i}$$  \[S19\]

is now completely specified by $a_e$, $\rho$, and $\sigma$.

For a given strain growing in a medium with a particular antibiotic and expressing some degree of resistance to that antibiotic, in vitro parameters uniquely determine the values of $\sigma$ and $\rho$; see tables S2, S5 for biochemical parameters of the specific systems studied in this work (Cm-CAT, Tc-TetA, Mn-TetA). In the main text we describe how experimentally tuning the parameter $\rho$ affects cell growth. Here we characterize the generic behavior of the system, i.e., the dependence of cell growth at different external concentrations as defined by Eq. [S16], with $\sigma$ and $\rho$ as arbitrary parameters.

2.1 Criteria for the coexistence of multiple growth phases

2.1.1 Origin of coexistence

Plotting the relation between $a_e$ and $a_i$ as defined by Eq. [S16] for a range of $\rho$ values, we see that for sufficiently large $\rho$, there exists a range of external antibiotic concentration $a_e$ for which multiple $a_i$ values correspond to a single $a_e$ for a given $\rho$ (fig. S18A). Since each internal concentration $a_i$ corresponds to a unique growth rate according to Eq. [S19], multiple solutions here correspond to the coexistence of multiple growth phases. As indicated in fig. S18B for a fixed set of $\sigma$ and $\rho$, the appearance of the local extrema $a_e^{\text{min}}$ and $a_e^{\text{max}}$ define the boundaries of external drug concentrations within which multiple growth phases coexist (for a particular strain/condition that correspond to the chosen values of $\sigma$ and $\rho$).

The two local extrema divide the solution of Eq. [S16] into three branches of internal antibiotic concentration: low, high, and intermediate, for $a_i < a_i^*$, $a_i > a_i^*$, and $a_i^* < a_i < a_i^*$, respectively, as indicated in fig. S18B, with the red curve segment indicating the intermediate region. Figs. S18C–D show the dependence of the growth rate $\lambda$ on $a_e$, by converting $a_i$ to $\lambda$ using Eq. [S19]. Comparing fig. S18B and 17D shows that the low-$a_i$ region corresponds to sub-inhibitory growth, and the high-$a_i$ region corresponds to growth inhibition, though the growth rate in this region is small but finite according to the
model. In the section below, we show that $a_{e}^{\text{max}}$ and $a_{e}^{\text{min}}$ correspond to the theoretical MIC and MCC respectively (Eqs. [S28] and [S39]) for parameter values relevant to this study. The intermediate-$a_i$ region is dynamically unstable (and hence irrelevant) according to deterministic kinetic analysis (see discussion surrounding Eq. [S44]).

2.1.2 Phase diagram and the onset of coexistence at the critical resistance efficacy

Fig. S18A shows that the local extrema disappear for sufficiently small $\rho$, i.e. for $\rho < \rho_c$; the disappearance of the phase of coexistence in the phase diagram of Fig. 4B reflects this transition for small $V_0$ (which controls the magnitude of $\rho$ in our experiments). The bifurcation point where the extrema just appears, commonly referred to as the “critical point”, can be obtained from the following defining conditions,

$$\frac{\partial a_e(a_i, \rho, \sigma)}{\partial a_i} = \frac{\partial^2 a_e(a_i, \rho, \sigma)}{\partial a_i^2} = 0$$ [S20]

and solving for the critical values $\rho_c$ and $a_{i,c}$ as a function of $\sigma$. We find the minimal resistance efficacy that produces phase coexistence to be

$$\rho_c(\sigma) = \left[\sigma^{1/3} + 1 + \sigma^{-1/3}\right]^3.$$ [S21]

The form of $\rho_c(\sigma)$ is plotted in fig. S19A, and the minimum at $\rho_c^{\text{min}} = 27$ when $\sigma = 1$ indicates that phase coexistence occurs most readily when $K_m \approx I_{50}$; i.e., relatively low values of resistance efficacy $\rho$ already give rise to coexistence. For the systems investigated in our experiments, $\sigma$ is of order one and coexistence is predicted to occur for $\rho \geq 29$ (see $I_{50}$ and $K_m$ values in tables S2, S5). For the Cm-CAT system, this corresponds to a critical value of $(V_0 / \kappa)_{\text{crit}} = 350 \mu$M based on the parameters in table S2.

2.1.3 Critical external antibiotic concentration

Similarly, solving Eq. [S21] for $a_{i,c}$ gives

$$a_{i,c} = \sigma^{-1/3} + \sigma^{-2/3}$$ [S22]

yielding the critical external concentration at the bifurcation point

$$a_e = (1 + \sigma^{-1/3})^3$$ [S23]

or

$$[a_e]_{\text{crit}} = \left(\sqrt[3]{I_{50}} + \sqrt[3]{K_m}\right)^3$$ [S24]

which takes the value $[\text{Cm}]_{\text{crit}} \approx 67 \mu$M for the Cm-CAT system (see table S2).

For $\rho < \rho_c(\sigma)$, there is a unique solution $a_i$ for each external concentration $a_e$, and a
homogeneous growth phase is expected. For $\rho > \rho_c(\sigma)$, $a_e^{\text{max}}$ and $a_e^{\text{min}}$ define the MIC and MCC respectively, as described above. We next describe their dependences on $\rho$, which is readily probed in experiments by changing the basal expression level of antibiotic resistance ($V_0$ in the model).

### 2.2 Interpretation and prediction of the Minimum Inhibitory Concentration (MIC)

To solve for the MIC, we look for the maximum of $a_e$ as defined by Eq. [S16]. The value of $a_i$ giving the maximum, $a_i^*$ (as illustrated in fig. S18B), is found to obey the following equation,

$$\sigma \cdot (a_i^*)^2 = 1 + (1 + a_i^*)^2 \cdot (1 + \sigma a_i^*)^2 / \rho.$$  \[S25\]

Solving for $a_i^*$ in powers of $1/\rho$ (and identifying $a_i^*$ as the smaller of the two solutions), we obtain

$$a_i^* \approx \frac{1 + b / \rho}{\sqrt{\sigma}}$$  \[S26\]

where $b = (1 + \sigma^{-1/2})^2 \cdot (1 + \sigma^{1/2})^2 / 2$, giving

$$a_e^{\text{max}} \approx \frac{\rho}{(1 + \sigma^{-1/2})^2} + \sigma^{-1/2} + \frac{1}{\rho} \left(1 + \sigma^{1/2}\right)^4 / 2\sigma^{3/2} + \ldots$$  \[S27\]

from Eq. [S16]. Since $\rho_c \geq 27$, then for $\rho > \rho_c(\sigma)$ where the maximum exists, the subleading terms in $1/\rho$ are negligible; see fig. S19B–C that illustrate this for the Cm-CAT system ($\sigma \approx 0.46$, table S2). Thus, the MIC is predicted to be

$$\text{MIC} = a_e^{\text{max}} \cdot I_{50} \approx \frac{V_0 / \kappa}{\left(1 + \sqrt{K_m / I_{50}}\right)^2} + \sqrt{K_m / I_{50}}$$  \[S28\]

which becomes for the Cm-CAT system

$$\text{MIC} \approx 0.16 \cdot V_0 / \kappa + 8 \ \mu\text{M}$$  \[S29\]

The approximate linear dependence of the MIC on the basal expression level of drug resistance is verified by the data (circled numbers) shown in Fig. 4B.

### 2.3 Abrupt drop in growth rate and the plateau-shaped fitness landscape

In fig. S19B, we show the accuracy of the approximation for $a_i^*$. Equation [S26] is seen to approximate the full solution very well until very close the critical point (compare the solid black and dashed red lines). In the limit $\rho \gg b$, which corresponds to $\rho \gg 4 \cdot \rho_c(\sigma)$ for the Cm-CAT system, the leading expression in Eq. [S26] (dashed black line)
already suffices. In this limit, Eq. [S16] is reduced to

\[ a_e = a_i \cdot \rho \cdot \frac{1}{1+a_i} \cdot \frac{1}{1+\sigma \cdot a_i}. \]  

[S31]

The growth rate at \( a_e^{\text{max}} \) becomes independent of \( \rho \), with

\[ \lambda^* = \frac{\lambda_0}{1+a_i} \approx \frac{\lambda_0}{1+\sqrt{K_m/I_{50}}}, \]  

[S32]

giving

\[ \lambda^* \approx 0.4 \cdot \lambda_0 \]  

[S33]

for the Cm-CAT system (fig. S19D). As there is no growth for concentrations above MIC, \( \lambda^* \) represents the size of the abrupt drop in the growth rate as the drug concentration is increased beyond the MIC. Growth rate data are consistent with the approximate independence of \( \lambda^* \) on the level of drug resistance activity \( V_0 \) as long as resistance efficacy \( V_0 / (\kappa \cdot K_m) \) is not too close to the critical point \( \rho_c \); see e.g., Figs. 4C and 5A.

2.4 Criteria for dormancy and the Minimum Coexistence Concentration (MCC)

To solve for the MCC, we look for the minimum of Eq. [S16], \( a_e^{\text{min}} \), which occurs at the internal concentration \( a_i = a_i^\dagger \) (see fig. S18B for illustration). It is also necessary to establish the conditions on parameters \( \rho \) and \( \sigma \) for which growth is effectively completely inhibited when \( a_i \geq a_i^\dagger \) (such that the system has a dormant state in addition to the growing state). We obtain \( a_i^\dagger \) as the larger solution to Eq. [S25]. In the limit \( [a]_{\text{int}} \gg K_m \), or equivalently \( a_i \gg \sigma^{-1} \), we find

\[ a_i^\dagger = \sqrt{\frac{\rho}{\sigma}} \left( 1 - \frac{\sigma^{1/2} + \sigma^{-1/2}}{\rho^{1/2}} \right) \]  

[S34]

to the first sub-leading order in \( \rho^{-1/2} \), with the value of the minimum given by

\[ a_e^{\text{min}} \approx \sqrt{\frac{\rho}{\sigma}} \left( 2 - \frac{\sigma^{1/2} + \sigma^{-1/2}}{\rho^{1/2}} \right) \]  

[S35]

This approximation is in good agreement with the full solution of \( a_e^{\text{min}} \) as shown in fig. 14.
S19E-F for $\sigma = 0.46$ (for the Cm-CAT system). To determine whether or not a dormant state exists for $a_e > a_e^{\text{min}}$, we find the predicted growth rate at this boundary ($a_i = a_i^\dagger$)

$$\lambda_i = \frac{\lambda_0}{1 + a_i} = \frac{\lambda_0}{\sqrt{\rho / \sigma - \sigma^{-1}}}$$  \[S36\]

The growth rate takes on low values, $\lambda_i / \lambda_0 < 0.1$ (practically undetectable), for antibiotic resistant strains in this work until very close to the critical point; see fig. S19G. When $\sigma > 2$ (and $\rho > \rho_c(\sigma)$ to ensure coexistence) this predicted growth rate can be appreciably greater than zero and is approximately

$$\frac{\lambda_i}{\lambda_0} \approx \sqrt{\frac{\sigma}{\rho}} = \sqrt{\frac{I_{50} \cdot \kappa}{V_0}}$$  \[S37\]

Equation [S37] suggests that growth rate along this “slow growth” branch can be increased arbitrarily by decreasing the ratio $\rho / \sigma$; however, bistability disappears for $\rho < \rho_c(\sigma) = (\sigma^{-1/3} + 1 + \sigma^{1/3})$ as in Eq. [S20]. The growth rate at $a_i = a_i^\dagger$ with $\rho = \rho_c(\sigma)$ gives the maximum value $\lambda_i$ may take at $a_e^{\text{min}}$:

$$\lambda_i^{\text{max}}(\sigma) = \frac{\lambda_0}{1 + \sigma^{-1/3} + \sigma^{-2/3}}$$  \[S38\]

(with $\rho = \rho_c(\sigma)$). This value is small (e.g. $\lambda_i^{\text{max}} \leq \lambda_0 / 3$) when $\sigma \leq 1$, or equivalently when $I_{50} \leq K_m$. The $I_{50}$ and $K_m$ parameters for the CAT and TetA systems used in this study (corresponding to $\sigma = 0.46$ and 0.4 respectively), give $\lambda_i^{\text{max}} / \lambda_0 \approx 0.25$ at the critical point such that cell growth in the ”high” $[a]_{\text{int}} > I_{50} \cdot a_e^{\text{min}}$ branch of drug concentration is virtually completely inhibited. Thus, the minimal coexistence concentration $a_e^{\text{min}}$ corresponds practically to the Minimal Coexistence Concentration (MCC), and $a_e^{\text{max}}$ corresponds to MIC as claimed above (Eq. [S28]). Using the approximate form Eq. [S35] and with $\rho / \sigma \gg 1$, the MCC is predicted to increase with the square root of drug resistance activity ($V_0$), as

$$\text{MCC} = a_e^{\text{min}} \cdot I_{50} \approx 2 \cdot \sqrt{I_{50} \cdot V_0 / \kappa} - (I_{50} + K_m)$$  \[S39\]

Additionally, we can write the MCC in terms of MIC for strains with high levels of resistance. In the limit $\text{MIC} \gg \sqrt{I_{50} \cdot K_m}$, MIC becomes approximately proportional to $V_0 / \kappa$ (to zero order in $\sqrt{I_{50} \cdot K_m / \text{MIC}}$, Eq. [S28]). Then with Eq. [S39] one finds a simple expression for MCC in terms of MIC

$$\text{MCC} \approx 2 \cdot \left(\sqrt{K_m + \sqrt{I_{50}} \cdot \sqrt{\text{MIC}}\right.$$  \[S40\]
Thus, it is possible to predict the range of concentrations over which growing and non-
growing cells coexist without explicitly knowing or fitting the value of parameter $V_0 / \kappa$.

2.5 Stochasticity cannot generate the plateau-shaped fitness landscape
We examine here whether the stochastic expression of CAT from its constitutive
promoter, when combined with the nonlinear relation between external and internal Cm
(Fig. 3B), might be sufficient to explain the plateau-shaped fitness landscape without
invoking growth-mediated feedback. In this scenario, a small (stochastic) decrease in
CAT expression could lead to a large increase in intracellular antibiotic concentration,
thereby arresting growth. A similar model was previously used to describe the transition
between growing and dormant states in natural persistence (91), in which the stochastic
expression of a “toxin” gene (hipA) induced cells to enter a dormant state, with the
lifetime of the dormant state determined by HipA concentration in a threshold-linear
manner. We find that while such a model can in principle give rise to heterogeneity in
growth rates among a population, it cannot reproduce the abrupt drop in growth rate
observed at the MIC: the average growth rate is still expected to decrease smoothly as a
function of drug concentration.

This can be understood by a simple mathematical analysis: an abrupt drop in growth rate
at the MIC manifests itself as a divergence in the average doubling time of the population
$\frac{\partial \tau}{\partial [A]_{\text{ext}}} \rightarrow \infty$, which can be rewritten as $\frac{\partial \tau}{\partial [A]_{\text{ext}}} = \frac{\partial \tau}{\partial [A]_{\text{int}}} \cdot \frac{\partial [A]_{\text{int}}}{\partial [A]_{\text{ext}}}$. Because the relation
between doubling time and intracellular antibiotic concentration is continuous (Eq. [4],
leading to the result in Fig. 3D), the observation of an abrupt jump would require

$$\frac{\partial [A]_{\text{int}}}{\partial [A]_{\text{ext}}} \rightarrow \infty \quad [S41]$$

i.e., the divergence must come from a discontinuity in the relation between
$[A]_{\text{int}}$ and $[A]_{\text{ext}}$. However, the latter relation is fixed by the well-characterized kinetics
of the CAT enzyme, which is given by the solution to Eq. [S11] (red line in Fig. 3B), and
clearly lacks any discontinuity in the absence of growth-mediated feedback. Even in the
extreme case where the relation between $[A]_{\text{int}}$ and $[A]_{\text{ext}}$ is exactly of the threshold-
linear form, it is still continuous (dashed grey line of Fig. 3B); the derivative has a jump
but still is not divergent. Therefore, including stochastic variation in this model cannot
give rise to the abrupt drop in growth rate at the MIC, i.e., the plateau-shaped fitness
landscape.

2.6 Constitutive expression of drug resistance and other forms of growth-mediated
feedback
In our analysis we assumed constitutive expression of drug resistance. Aside from being
the simplest system to characterize, CAT is indeed most often expressed constitutively in

Gram- bacteria inside Tn9 and elsewhere (24). This scenario also occurs naturally in inte\nrons, in which many unregulated resistance genes are placed under the control of a single high-activity constitutive promoter (25). Resistance genes in resistant clinical isolates have also been found to be under the control of constitutive promoters (27). But for a protein whose expression is specifically regulated, e.g. at the transcriptional or translational level, the growth-rate dependence need not be linear as in Fig. 3C (17, 57).

In the simplest approximation one can modify Eq. [S13] to the form

\[ V_{\text{max}} = V_0 \cdot \left( \frac{\lambda}{\lambda_0} \right)^n, \quad n \geq 0. \]  

[S42]

Deriving the consequences of this relation is beyond the scope of the current work, but we mention here some key results: First, growth bistability and plateau-shaped fitness still occur for a broad range of the (\(\rho, \sigma\))-parameter space, but the criterion Eq. [S21] for the occurrence of the critical point depends on the value \(n\). With the modification in Eq. [S42], one can show that \(\sigma^\text{max}\) still increases linearly with resistance efficacy \(\rho\) for strong resistance, and hence approximately MIC \(\sim V_0 / \kappa\) as before, although the slope and intercept in Eq. [S28] will depend nontrivially on \(n\) also. Interestingly, linear dependence of the empirical MIC on \(V_0\) has also been reported for bacterial resistance to other antibiotics (89, 92). This is likely a result of the generic relationship between \([a]_{\text{int}}\) and \([a]_{\text{ext}}\), in which Michaelis-Menten kinetics of resistance activity (Eq. [S10]) are balanced by diffusive influx (Eq. [S7]). Then, assuming cell growth is inhibited at some characteristic internal drug concentration (corresponding empirically to \([a]_{\text{ext}}=\text{MIC}\)), simple flux balance dictates a linear relationship between this \([a]_{\text{int}}\) and MIC (Eq. [S10], \([a]_{\text{ext}} \gg V_0 / \kappa\)).

The MCC is more sensitive than MIC to the form of the feedback, with the result

\[ \text{MCC} \sim \left( \frac{V_0}{\kappa} \right)^{\frac{1}{n+1}} \]  

[S43]

For \(n > 1\), the feedback is more cooperative and the MCC is reduced. Thus, the onset of growth bistability occurs at lower antibiotic concentrations as the cooperativity in feedback \((n)\) is increased. Conversely, for weak feedback (in the limit \(n \to 0\)) the region of bistability shrinks as \(\text{MCC} \sim V_0 / \kappa\) approaches the MIC.

### 2.7 Stability analysis

Here we perform a stability analysis of the deterministic model whose steady states are captured by Eqs. [S16] and [S19]. Specifically, we address the qualitative question regarding the stability in the coexistence regime where multiple solutions are obtained. By themselves, Eqs. [S16] and [S19] provide no information as to whether cells can approach the steady state when antibiotic concentrations are transiently perturbed. However, we can address this kinetic question qualitatively by implementing a quasi-steady-state approximation in which the effect of drug on cell growth (Eq. [S19]) and the
growth-rate dependence of enzyme levels (Eq. [S13]) remain satisfied as internal antibiotic concentration changes. In this limit we write the rate of change of the internal concentration $a_i$ as

$$\frac{da_i}{dt} = \kappa \left( a_e - a_i \cdot \left( 1 + \rho \frac{1}{1+a_i + \sigma \cdot a_i} \right) \right) = f(a_i) \quad [S44]$$

This form is dictated by the effects of drug influx and efflux (or modification, degradation) on the internal drug concentration. Eq. [S16] gives the steady state condition of Eq. [S44], i.e., $f(a_i) = 0$.

We perform linear stability analysis with respect to changes in concentration by perturbing about the steady state solution $a_{i,s}$, defined by $f(a_{i,s}) = 0$. The expected response to small perturbations $\delta a_i = a_i - a_{i,s}$ is $\delta f = c \cdot \delta a_i$, with

$$c = \left. \frac{\partial f}{\partial a_i} \right|_{f=0} = -\kappa \cdot \left. \frac{\partial a_e(a_i; \rho, \sigma)}{\partial a_i} \right|_{a_i=a_{i,s}} \quad [S45]$$

and using Eq. [S16] for the dependence of $a_{i,s}$ on $a_e$. Stability requires $c < 0$ such that the system returns to the steady state for small perturbations away from the stationary solution. The sign of $c$ can be easily determined graphically, by inspecting the form of $a_e(a_i)$ as depicted in Fig. 17A-B: We see that $\partial a_e / \partial a_i < 0$ only in the intermediate (red) segment in between the two extrema. Thus, $c > 0$ and the solution $a_{i,s}$ in the intermediate range between $a_i^\ast$ and $a_i^\dagger$ is unstable, while everywhere else the solution is stable. This geometric statement can be readily proved algebraically for any values of the parameters for this system.

3. Generalizations of the model and applicability to tetracycline resistance: detailed descriptions of drug influx and efflux

In our analysis of antibiotic resistance above, we made several simplifying assumptions to describe the passive penetration of antibiotics into cells. Here we describe which assumptions may be relaxed without qualitatively changing central results described above. Doing so enables us to provide an independent estimate of the permeability parameter $\kappa$ to compare with the value fixed by results in the main text (table S2). We also demonstrate the applicability of the model to more complicated antibiotic resistance systems, using efflux of tetracyclines via TetA as a second model system.

3.1 Effective membrane permeability

First we must appreciate that permeability ($\kappa$) in the sense used above depends on both the specific permeability ($\pi$) of antibiotic through the membrane, and the surface area ($A$) of the cell membrane such that the rate of molecules passing through the membrane is
given by
\[ j_{\text{influx}} = \pi \cdot A \cdot (\{a\}_\text{ext} - \{a\}_\text{int}) \] [S46]

with \( \pi \) in units of distance per time (lower-case \( j \) denotes flux of molecules rather than flux of concentration described by \( J \) in Fig. 3B and Eq. [S7]). One recovers
\[ J_{\text{influx}} = \kappa \cdot (\{a\}_\text{ext} - \{a\}_\text{int}) \],
the rate at which intracellular concentration increases in Eq. [S7], with the definition
\[ \kappa \equiv \pi \cdot \frac{A}{\Omega} \] ,
[S47]

taking \( \Omega \) to be the volume of the cell.

3.1.1 A single effective membrane permeability can describe the combined permeability of both the inner- and outer- membranes

The simplest justification for the use of Eq. [S45] or [S7] is for antibiotics that enter the cell passively by diffusion. However, reality is more complex. Gram’ bacteria possess both outer- and inner-membranes, each with their own characteristic permeabilities that in general depend on the levels of porins and low affinity efflux pumps (93, 94). Despite these complexities, we show below that in many cases, including for the Cm-CAT system, Eq. [S7] or [S46] is a reasonably effective model of drug influx at steady state.

Allowing for separate permeability rates of outer and inner membranes (OM and IM), we rewrite the steady-state \( (J_{\text{influx}} = J_{\text{removal}}) \) as
\[ 0 = \kappa_{\text{IM}} \cdot (\{a\}_\text{per} - \{a\}_\text{int}) - V_0 \cdot \frac{1}{1 + \{a\}_\text{int} / I_{50}} \cdot \frac{1}{1 + K_m / \{a\}_\text{int}}, \] [S48]

where \( \{a\}_\text{per} \) is the concentration of antibiotic inside the periplasmic cavity and \( \kappa_{\text{IM}} \) is the permeability of the inner membrane. Provided that the resistance enzyme removes antibiotic from the cell completely (e.g. not merely expulsion into the periplasm), we can equate the flux of number of molecules across each membrane at steady state as follows,
\[ j_{\text{influx}} = j_{\text{OM}} = j_{\text{IM}} \Rightarrow\]
\[ \pi_{\text{OM}} \cdot A_{\text{OM}} \cdot (\{a\}_\text{ext} - \{a\}_\text{per}) = \pi_{\text{IM}} \cdot A_{\text{IM}} \cdot (\{a\}_\text{per} - \{a\}_\text{int}) \] [S49]

Using \( j_{\text{OM}} = j_{\text{IM}} \), one eliminates \( \{a\}_\text{per} \) from the above Eq. [S48] and defines an effective permeability constant
\[ \tilde{\kappa} = \frac{\pi_{\text{IM}}}{\Omega_{\text{cyl}}} \cdot \frac{A_{\text{IM}} \cdot \pi_{\text{IM}}}{A_{\text{OM}} \cdot \pi_{\text{OM}}} \] [S50]

to recover the results of Eqs. [S15], [S16], and [S17], with \( \kappa \to \tilde{\kappa} \). If the resistance mechanism in consideration expels antibiotic into periplasm, as is the case for TetA-mediated efflux of tetracyclines (88), one obtains an even simpler expression:
\[ \kappa' \equiv \kappa_{IM} = \pi_{IM} \cdot \frac{A_{IM}}{\Omega_{cyt}}, \]  

a result independent of outer membrane permeability at steady state. These results connect our simple permeability parameter \( \kappa \) to the biophysical properties of the cell and motivate our claim that antibiotic influx can be treated according to Eq. [S7]. Henceforth, we drop the tilde notation with the understanding that \( \kappa \) is used as a “catch-all” parameter in the sense above.

### 3.2 The effect of endogenous drug efflux and drug chelation on tetracycline resistance

We formulated our model with the goal of describing effects of antibiotics on bacterial growth in the presence of a single high affinity efflux- or enzyme-mediated drug resistance mechanism. But as described near Eq. [S7], cells often additionally possess low-affinity native efflux mechanisms that potentially complicate our simple description. Additionally, some antibiotics may be charged by chelators under physiological conditions, ultimately affecting permeation rate and equilibrium distributions of the drug across cell membranes (85). We briefly treat the effects of endogenous efflux and chelation here, emphasizing that key results do not differ qualitatively from those derived above.

#### 3.2.1 Modeling drug influx in the presence of endogenous drug efflux and drug chelation

We begin by incorporating the effects of chelation or endogenous efflux into our description of passive influx of drug into the cell. We assume the presence of low-affinity efflux or chelation alters the equilibrium concentration of internal drug by a factor of \( \eta \) such that \([a]_{\text{int}} = \eta \cdot [a]_{\text{ext}}\) at equilibrium. When \( \eta \) characterizes low-affinity efflux, comparison with Eq. [S9] reveals that \( \eta = \left(1 + \frac{V_{\text{native}}^{\max}}{\kappa \cdot K_{\text{native}}} \right)^{-1} \), and \( \eta \leq 1 \). If the parameter characterizes chelation, then \( \eta \) may take any positive value, indicating that Donnan potential may act to either increase (\( \eta > 1 \)) or decrease (\( \eta < 1 \)) internal drug concentration relative to extracellular concentration. One can thus describe the influx by modifying Eq. [S7] to the form

\[ J_{\text{influx}} = \kappa \cdot ([a]_{\text{ext}} - [a]_{\text{int}} / \eta). \]  

Yet another layer of detail is that equilibrium may be shifted (as described above) separately across the inner and outer membranes, giving two shift factors: \( \eta_{\text{per}} \) and \( \eta_{\text{cyt}} \). For example, an endogenous efflux pump could expel antibiotic from the cytoplasm into the periplasm (described by \( \eta_{\text{cyt}} \)), and this would have no effect on the equilibrium distribution of the drug between the periplasm and external medium (described by \( \eta_{\text{per}} \)).

#### 3.2.2 Effects of endogenous drug efflux and drug chelation on phase coexistence,
MIC, and MCC
Without providing the algebraic details here, we affirm that one can recover the steady state relation Eq. [S16] by balancing influx across cell membranes with a high-affinity efflux term $J_{\text{removal}}$, and then rescaling variables judiciously. All subsequent derivations proceed as before, and one can show that in analogy to Eq. [S21],

$$\rho_c(\sigma, \eta_{\text{per}}, \eta_{\text{cyt}}) = \left(\sigma^{-1/3} + 1 + \sigma^{1/3}\right)^3$$

[S53]

And in place of Eq. [S28] we have

$$\text{MIC} \approx \frac{V_0 / \kappa_\eta}{\left(1 + \sqrt{K_m / I_{50}}\right)} + \sqrt{K_m \cdot I_{50}} / \eta_{\text{per}} \cdot \eta_{\text{cyt}} + \ldots$$

[S54]

with high-affinity efflux described by $V_0$ as before, and $\kappa_\eta$ as effective permeability that generally depends on $\eta_{\text{per}}$ and $\eta_{\text{cyt}}$. We showed above that effective permeability $\kappa$ is defined by flux balance and depends on details of efflux (Eqs. [S50], [S51]) and $\kappa_\eta$ must be defined in the same manner. The essential qualitative results of the model have not changed; though this analysis shows that, for example, endogenous low-affinity efflux ($0 < \eta < 1$) would shift the critical point to a higher value (delaying the onset of bistability), but would not affect the linear relation between MIC and $V_0$. And in analogy to Eq. [S40], the theoretical MCC is given

$$\text{MCC} \approx 2 \cdot \left(\sqrt{K_m} + \sqrt{I_{50}}\right) \cdot \sqrt{\frac{\text{MIC}}{\eta_{\text{cyt}} \cdot \eta_{\text{per}}}}$$

[S55]

Chloramphenicol is electrically uncharged at physiological pH and believed to move passively across the cell membranes (41), and we can regard $\eta_{\text{per}} = \eta_{\text{cyt}} \approx 1$.

3.2.3 Modeling tetracycline efflux by TetA in the presence of Tc-chelation
Tc is chelated by Mg$^{2+}$ ions that strongly affect its partition, with $\eta_{\text{cyt}} \approx \eta_{\text{per}} \approx 2$. The equilibrium cytoplasmic tetracycline concentration is thus four-fold higher than the external concentration (88). In spite of this effect, we are justified in determining $V_0/\kappa_{\text{Tc}}$ using Eq. [S28] rather than Eq. [S54] (table S5), because with MIC $\approx 700$ µM (fig. S11A) and $\sqrt{K_m \cdot I_{50}} \approx 6.5$ µM (table S5), we have MIC $\gg \sqrt{K_m \cdot I_{50}}$ and Eq. [S54] is essentially equivalent to Eq. [S28] with the definition $\kappa_{\text{Tc}} = \kappa \cdot \eta_{\text{per}}$ (for TetA efflux, flux balance analysis reveals $\kappa_{\eta} = \eta_{\text{per}} \cdot \kappa$, independent of $\eta_{\text{cyt}}$).

Because $\eta_{\text{cyt}} \approx \eta_{\text{per}} \approx 2$ for Tc, Eq. [S55] predicts MCC values for the drug-resistant strain Ta1 to be lower (relative to MIC) than that predicted by Eq. [S40]. However, high levels of Tc (and other antibiotics) have been shown to depolarize cell membranes (95, 96), so
the shift factors likely have growth rate-dependence for which $\eta \to 1$ when $Tc$ concentrations are high enough to completely inhibit growth. Thus, MCC in these systems may take some intermediate value between that given by Eq. [S40] and that by Eq. [S55]. Nonetheless, the plots in fig. S11A-B depict the region of bistability predicted by the analysis above with $\eta_{cyt} = \eta_{per} = 2$ for all antibiotic concentrations.

### 3.3 Independent estimates of the cell's permeability to chloramphenicol

To determine whether our prediction of $\kappa \sim 1.5$ s$^{-1}$ for chloramphenicol (based on MIC and CAT activity of Cat1, table S2) is reasonable, we estimate here the specific permeability of $E. coli$’s inner- and outer-membranes (IM, OM) to Cm using primarily the method outlined by Thanassi et al. (88), which is based on Cm hydrophobicity, charge, etc. Then we compose an effective permeability constant from the specific permeabilities with Eq. [S50].

#### 3.3.1 Inner membrane permeability

Collander (97) demonstrated that a molecule’s specific permeability $\pi$ through a typical lipid bilayer membrane (similar in composition to the inner membrane in $E. coli$) is related to the molecular weight (MW) and the partition coefficient $c$ of the molecule in ether-water such that $\log(c^{1.3}) \sim \log(\pi \cdot MW^{1.5})$ at 20ºC. The apparent partition coefficient of Cm in ether-water for $2.0 \leq pH \leq 9.0$ is given (98) at 25ºC as $c_{25}=3$. However, to use Collander’s relation and obtain permeability, we must correct for the temperature dependence of partition coefficients (99) $\pi(T) \approx e^{-E_a/RT}$. We take Pramer’s (100) value of activation energy $E_a \sim 10^4$ cal/mol, based on the diffusion of Cm into Nitella cytoplasm; this number is comparable to values for other antibiotics passing solely through lipid bilayer (88, 99). This correction gives $c_{20} \approx 2.2$. Collander’s data then suggest that for molecules with MW and partition coefficient similar to those of Cm, the permeability falls in the range $10^{-5} \leq \pi \leq 10^{-4}$ cm/s . To obtain the specific permeability at 37º C, we again correct for temperature to obtain

$$2.6 \times 10^{-3} \lesssim \pi_{i,M} \lesssim 2.6 \times 10^{-4} \text{ cm/s .}$$  

#### 3.3.2 Outer membrane permeability

Estimating outer membrane permeability is less direct and we must appeal to permeabilities of other compounds. Chloramphenicol diffusion through OM porins is much faster than through lipopolysaccharide, and loss of porin production significantly decreases the ability of Cm to inhibit growth (93, 101). Cm diffuses through PhoE, OmpC, and OmpF porins; molecular weight, charge, size, and hydrophobicity have all been found to influence the permeability of antibiotics through these OM porins (93, 101). For the lack of direct data on chloramphenicol permeability, we use the permeabilities of two drugs with similar size and total charge (tetracycline and cephalaxin) through the OmpF porin as reference points. By combining the data due to Nikaido et al. (102, 103), we obtain absolute permeability rates of various compounds
through porin-containing membrane as a function of hydrophobicity, with hydrophobicity given by the logarithm of the partition coefficient in n-octanol. The partition coefficient of Cm in n-octanol is reported \( (104) \) as \( c_{n-oct} = 12 \). Electrically neutral compounds of similar size and hydrophobicity as Cm (like cephalixin), have permeability through OmpF of \( \sim 4 \times 10^{-4} \text{ cm/s} \) \( (103) \). Thanassi et al. \( (88) \) give a permeability of the bulkier tetracycline through these porins \( \sim 10^{-5} \text{ cm/s} \). We therefore estimate \( 10^{-5} \leq \pi_{O.M.} \leq 10^{-4} \text{ cm/s} \) for chloramphenicol.

### 3.3.3 Comparing model predictions with estimated permeability

Then using Eq. [S50] we combine these specific inner- and outer-membrane permeabilities and cytoplasmic volume \( \Omega_{cyt} \) to compute the effective permeability for both membranes together

\[
0.37 \leq \kappa \leq 7.5 \text{ s}^{-1}
\]  

with \( A_i = 105 \text{ cm}^2/\text{mg}, A_o = 132 \text{ cm}^2/\text{mg}, \) and \( \Omega_{cyt} = 0.0024 \text{ cm}^3/\text{mg} \) (all mass units are dry weight of cells) \( (88) \), although we use these parameter values only as an order of magnitude estimate, and in particular the \textit{in vitro} upper bound may be somewhat higher. In table S2 we see that our predicted value of permeability \( \kappa_{Cat1} \sim 1.5 \text{ s}^{-1} \) for strain Cat1 is within this estimated range. Similarly, with \( V_0 \) fixed for all CAT-expressing strains (Fig. 4B, bottom), the parameter fitting of all CAT-expressing strains for which growth rates were measured (colored lines, fig. S17), gives an average permeability for all strains \( \kappa_{all} \sim 1.5 \pm 0.14 \text{ s}^{-1} \) (SD).

Because the estimated range for \( \kappa \) is based solely on physical properties like the hydrophobicity, and on permeability of cells away from steady-state growth, we offer the following caveat: In our simple model (Eq. [S15]), we did not explicitly include the effects of low-affinity native efflux and porin regulation; therefore the permeability parameter \( \kappa \) is an effective one including such effects. (For similar reasons, we acknowledged near Eq. [S9] that our fit value of the half-inhibition constant \( I_{50} \) might actually represent an effective \( \bar{I}_{50} \)). Therefore, we expect this \( \kappa \), representing an effective permeability of the cell at steady state (from table S2), to be lower than a value estimated on physical properties alone.
Fig. S1: Positive feedback in drug resistance. A proposed positive feedback loop linking antibiotic drug, cell growth, and the expression of drug resistance. The positive link (green arrow) between drug-inhibited growth and gene expression is expected from global growth-mediated effects even for unregulated proteins (16).
**Fig. S2: The apparent MIC and viability of Cat1 cells in chloramphenicol from plate assays.** Typical results for the growth of Cm-resistant (Cat1) cells on LB agar plates containing various concentrations of Cm (shown at the bottom). Cells were diluted from log phase batch cultures lacking Cm, and were spread onto agar at densities indicated on the left of each panel. Plates were incubated overnight for ~18 hours at 37°C; see Supplementary Methods for details. 

A, To determine the apparent MIC, referred to as \( \text{MIC}_{\text{plate}} \) and defined as the Cm concentration above which visible colonies formed at a fraction of less than \( 10^{-4} \) per inoculum after 18 hours of incubation, the plates are plated at very high densities. The plating results lead to the value of \( \text{MIC}_{\text{plate}} = 1.0 \) mM for this strain. Similar results were obtained from replicates. 

B, 100-fold lower cell density was used to reveal differential fraction of colony formation for the Cat1 strain. The number of colonies formed decreases steadily over sub-MIC Cm concentrations. The fraction of colonies formed per inoculum is plotted in Fig. 1B (green circles). 

C-D, The few Cat1 colonies that formed on plates with Cm concentration close to 80-90% of \( \text{MIC}_{\text{plate}} \) are not mutants. A colony chosen from the indicated plate (orange circle in panel B) was grown in the batch culture and replated on LB agar with the same range of Cm at low or high density (panels C and D respectively). The results obtained are similar to those from Cat1 cells that had never been exposed to Cm (compare panels A and B).
Fig. S3: The apparent MIC and viability of wild type (EQ4) cells in chloramphenicol from plate assays. Same as in fig. S2 panels A and B but for wild type cells. MIC\(_{\text{plate}}\) is determined to be 15 µM from high density plating (panel A). The number of colonies formed on each plate remains nearly constant as the Cm concentration approaches MIC\(_{\text{plate}}\) (panel B), in contrast to the decreasing CFU count in the Cm-resistant strain Cat1 at sub-MIC Cm concentrations. The fraction of colonies formed per inoculum is plotted in Fig. 1B (blue triangles).
Fig. S4: Design and characterization of a microfluidic culturing device for the maintenance of Cm concentration. We designed a microfluidic culturing device to monitor cell growth in a constant environment at the single cell level. The device is similar to a previous design (28), with a few modifications. Fresh medium flows actively through the outside reservoir (channels, in green) and diffuses into the cell growth chambers (center red in top-down view, 1mm×0.1mm×1.5µm). The small geometry of microfluidic chambers ensures the fast fluid exchange rate between the reservoir and the chamber. This is shown in panel B, where we abruptly introduced a fluorescent dye (Rhodamine 110, invitrogen) into a channel (black squares) and measured the changes in fluorescence at the edge (red circles) and in the middle (blue triangles) of the chamber. It took ~ 40 sec for fluorescence to reach 90% of its maximum level, suggesting that molecules of similar sizes as the dye are replenished approximately every 40 sec. The fast fluid exchange rate ensures that a constant chemical environment, particularly, the drug concentration, is maintained inside the chamber. This is important because the cells studied here express CAT, which modifies Cm. Thus, the Cm concentration in the growth
medium would otherwise decrease over the course of experiment, making it difficult for long-term observation.

In panel C, we compared the growth in a microfluidic chamber to that in batch culture. First of all, wild type (EQ4) cell and ΔmotA (EQ4m) cells grew at similar rates in the batch culture (solid black squares and circles, respectively) and the wild type (solid black square), with ~60 min doubling time (line) in minimal medium with glycerol and NH₄⁺. Next, the growth of the ΔmotA strain in the microfluidic chamber was characterized (open red circle) by counting the pixels occupied by growing colonies in phase contrast images; see Supplementary Methods). The growth rate is ~10% slower than that in batch culture.
Fig. S5: Ampicillin enrichment assay. This assay takes advantage of the fact that ampicillin (Amp) kills growing cells to detect the fraction of dormant cells in a batch culture (35, 105). A, Batch cultures grow in glass tubes containing media with various concentrations of Cm for 1-2 hours to allow cells to acclimate to Cm upshift (from Cm-free preculture). B, Cultures are diluted into identical medium with Cm, together with Amp (100 µg/ml). C, A small volume of culture from each tube is spread onto LB plates lacking antibiotics to determine CFU per culture volume prior to enrichment (the same is done at 100-fold dilution to ensure a countable number of cells). D, Cultures incubate in the amp-enriched media for 6-7 hours. During this incubation period, cultures become enriched for dormant cells as ampicillin kills growing cells rapidly but has little effect on non-growing cells (35). E, Identical volumes of cells from each post-enrichment culture are spread onto LB-agar plates lacking antibiotics. F, Plates incubate overnight at 37°C to
reveal colonies formed from cells that remained dormant during enrichment. Comparisons between post- and pre-enrichment CFU yields the fraction of dormant cells in each culture, plotted in Fig. 2F for strain Cat1. See also fig. S7.
Fig. S6: Ampicillin enrichment in microfluidic chambers. A. This assay was performed in the microfluidic chambers for GFP-expressing Cat1m cells (GCat1m) growing in minimal medium. We introduced 0.7 mM Cm into the medium at time zero, and added Amp (200 µg/ml) at t=9 hr. By t=24 hr, most cells appeared elongated, stopped dividing, and lost fluorescence (characteristic of the loss of cytoplasmic contents accompanying cell lysis and death (106)). After switching to drug-free medium at t=25hr, cells with fluorescence (those that were not killed by Amp) began growing again, becoming clearly visible after a 12-hr wait (t=37hr). B. Detail of (A) shows that only dormant cells survive ampicillin treatment. The history of microcolonies was tracked backwards in time, throughout and prior to enrichment. The red circle shows the history of a representative dormant cell after ampicillin was added at t=9 hrs. By visual inspection, we observed that this cell was not growing before and after ampicillin addition at t=9 hrs, but began growing after we removed Amp and Cm. Using this technique we found that cells...
surviving enrichment (and subsequently creating new microcolonies at t=25 hr) were always dormant prior to enrichment, with no colonies forming from cells growing during Cm or Amp+Cm treatment.
Fig. S7: The MCC of Cat1 and wild type cells. Cat1 and wild type (EQ4) cells were treated by the Ampicillin enrichment assay described in fig. S5. We measured viable cell densities of ~10^3 cells per µL culture with dilution plating prior to enrichment (giving ~5-8×10^4 colonies/plate, representative results on left). Comparisons between post- and pre-enrichment CFU yields the fraction of dormant cells in each culture. The concentrations of Cm used during enrichment are indicated beneath plates. A, The dormancy frequency is high at and above MIC (≥1.0 mM) as expected for Cm-inhibited cells (37). In the absence of Cm, we observed typical “background” dormancy frequencies of ~10^-3 in Amp-cultures as have been reported by others under similar conditions, attributed to the naturally occurring persisters (31, 32, 107, 108). Strikingly, colonies significantly (> 10-fold) above the background level were obtained from Amp-treated Cat1 cultures, when enriched with concentrations of Cm well below the MIC, e.g., at 0.4 mM Cm. The ‘minimal dormancy concentration’ (MCC), defined as the minimum Cm concentration above which at least 1% of the inoculant survives enrichment, is found to be between 0.3-0.4 mM for this strain (red arrow). A conservative cutoff of 1% is used here to avoid false positives due to variability (108) in the naturally appearing persisters (31, 32, 107) and possible variability in culture density. B, Wild-type cells could not survive ampicillin treatment even with Cm concentrations close to the MIC, e.g., for very slowly growing cultures with λ ≤ 0.15 hr^{-1} at [Cm]_{ext} > 12 µM (Fig. 3D). Less than 0.1% of wild-type cells formed colonies post-enrichment, for all sub-MIC Cm concentrations used during enrichment (indicated at the bottom), similar to the naturally occurring persisters in Amp enrichment without Cm (31, 32, 107, 108). The results suggest that most wild-type cells grew in sub-inhibitory Cm and were therefore killed by Amp. Only at very high Cm concentrations (≥ 20 µM) were EQ4 cells significantly protected from the bactericidal effects of Amp during incubation (plate to the far right).
**Fig. S8:** Tetracycline-resistant strain Ta1 showed bimodal behavior when exposed to Tc and wild type did not.  

A. Growth rates of tetracycline-resistant strain Ta1 growing in microfluidic chambers containing minimal medium with indicated concentrations of Tc. Growth rate drops abruptly in the device at 0.6 mM Tc.  

B. Green bars give the percentage of Ta1 cells to continue growing in microfluidic chambers after adding indicated concentration of Tc (as in Figure 1E).  

C. Growth rates of wild type (EQ4m) in microfluidic device. All wild type cells grew (none were dormant) for all sub-MIC Tc concentrations. Figure S12A gives the predicted MIC, MCC, and growth rates for this strain based on growth in batch culture.
**Fig. S9: Growth-mediated global effects on gene expression.** The data in Fig. 3C shows that the levels of “unregulated” or “constitutively expressed” genes decrease linearly with the growth rate upon translational inhibition. The rationale of this behavior is that under translational inhibition, the cell allocates growth-limiting resources, particularly the ribosomes, to increase the synthesis of the translational apparatus (signaled by ppGpp (109)). Since most of the ribosomes are engaged in translation according to the bacterial growth laws (16), this increase results in decreased synthesis of unregulated genes. Below we summarize the basic elements of a simple mathematical model that quantitatively predicts this growth-mediated global effect on gene expression, as established by Scott et al (16).

The model partitions the proteome into 3-sectors (pie chart above), each governed by a distinct growth-rate dependence: (i) The ribosomal sector R, whose mass fraction $\phi_R$ (blue) decreases as growth is slowed down by nutrient limitation but increases as growth is slowed down by translation (as illustrated by the figure above and to be described in more detail below); (ii) a growth rate independent sector Q, whose mass fraction $\phi_Q$ is growth-rate independent (green); and (iii) the remaining sector P whose mass fraction $\phi_P$ (red) is comprised of what remains after the ribosomal sector changes in response to growth-mediated changes, i.e.,

$$\phi_P = 1 - \phi_R - \phi_Q$$  \[[a]\]

Constitutively expressed genes are suggested and shown to belong to the P-sector (16).

It has been known for over 50 years that exponentially growing cultures of *E. coli* exhibit an obligatory linear relation (growth law) between the ribosomal content and the growth rate $\lambda$, when growth is varied by changing nutrient composition in the medium (44, 110). In terms of the ribosomal fraction $\phi_R$ introduced above, this growth law can be written mathematically as

$$\lambda = \gamma \cdot (\phi_R - \phi_0)$$  \[[b]\]

where the proportionality constant $\gamma$ is established to be the translation rate (16), and $\phi_0$ is an empirically determined offset.
Scott et al. (16) established another linear relation between $\phi_R$ and $\lambda$ if growth is varied by changing the translation rate $\gamma$, either by using various translational mutant or by applying sub-lethal doses of translation-inhibiting antibiotics, under a given saturating nutrient source,

$$\lambda = \nu \cdot (\phi_R^{\text{max}} - \phi_R). \quad [c]$$

This relation describes a trend of increase in the ribosomal fraction when growth is slowed by translational inhibition such as sub-lethal dose of Cm inhibition studied in this work. The proportionality constant $\nu$ is found to increase for nutrients of increasing quality, while $\phi_R^{\text{max}}$ is a constant that represents the maximum proteome fraction allocable to the ribosomal fraction.

The value of $\phi_R^{\text{max}}$, found empirically to be $\sim 50\%$ for a spectrum of nutrient sources, is interpreted by Scott et al (16) to arise from the fixed proteome fraction, $\phi_Q$. Thus,

$$\phi_R^{\text{max}} = 1 - \phi_Q. \quad [d]$$

Using Eqs. [c] and [d] in [a] immediately leads to

$$\phi_P = \frac{\lambda}{\nu} \quad [e]$$

for translation-inhibited growth. Eq. [e] is the basis of Eq. [3] in Fig. 3, i.e., the linear relation between constitutive gene expression and the growth rate. Equations [b] and [c] can be combined to obtain Eq. [S4] with $\lambda_0 = \lambda(\gamma_0, \nu)$ as the growth rate of drug-free cells, and $\lambda_0^{\text{max}} = \lambda(\gamma_0, \infty) \approx 2.85 / \text{h}$, which is the basis for the linear relation between doubling time and drug concentration.
Fig. S10: Inducible resistance gene expression also depends on growth rate.
Tetracyclines (Tc) induce gene expression from the $P_{tet}$ promoter that drives expression of the efflux pump, TetA. Data replotted from (111) (magenta) show how promoter activity is affected by growth rate (when inhibited by Tc) in a strain expressing a tetA-lacZ translational fusion gene from the native $P_{tet}$ promoter. At very low Tc concentrations (too low to inhibit growth) beta-galactosidase activity is induced to its maximum level. However, activity decreases in cultures containing higher concentrations of Tc, yielding an apparently linear dependence between growth rate and gene expression that is similar to the trend observed for constitutively expressed genes (background symbols rescaled and replotted from Fig. 3C).
Fig. S11: Growth curves of Cat1 cells in batch culture with chloramphenicol. Representative growth curves showing steady exponential growth in these cultures for up to seven generations, after which the experiments were usually halted. The lines are least squares fit of log-OD with time. Inverted triangles show unambiguously the non-growing cultures across the MIC.
Fig. S12: The MIC and growth rates of Ta1 cells in batch culture with tetracycline and minocycline. A. The relative decrease in the growth rate of Ta1 cells constitutively expressing TetA in minimal medium with various Tc concentrations (red disks) is well described by the growth feedback model (various lines). The solid black line is drawn with the \textit{in vitro} value of $I_{50}^{Tc}$ (table S5, see also panel D below), and $V_0/\kappa_{Tc}$ as fixed by equation \[S28\], with MIC=650±50 µM from batch culture growth data (red disks). Alternatively, we fit the model to only the empirical rates of growing cultures (i.e., when $\lambda/\lambda_0 > 0.15$), fitting either $V_0/\kappa_{Tc}$ alone, or $V_0/\kappa_{Tc}$ and $I_{50}^{Tc}$ simultaneously as free parameters (blue and dashed gray lines respectively). Fits using growth rates provide parameter values very close to those predicted by MIC, see table S5 for parameters and fitting procedures. All growth conditions are given in Supplementary Methods. B, Same as (A), except with minocycline instead of tetracycline. Each data point is growth rate from a single experiment, with experiments performed on three separate days. C, The
relative growth rates of tetracycline-sensitive tetA strain (Cat1) grown in minimal media with various tetracycline concentrations (red circles). In contrast to Tc-resistant strain Ta1, growth rate does not abruptly drop to near zero across some threshold tetracycline concentration. D. Data replotted from (C) shows that relative doubling time increases nearly linearly with drug concentration. Based on in vitro data we predict $I_{50}^{Tc} \approx 4 \, \mu M$ (table S5) such that doubling time should increase by a factor of two at $[Tc]_{ext} \approx 1 \, \mu M$ (because $[Tc]_{int} = 4 \cdot [Tc]_{ext}$, see discussion below Eq. [S55]). This prediction is consistent with doubling times presented here.
**Fig. S13: Dormancy of Ta1 cells in sub-MIC concentrations of Mn.** Results of batch culture Amp enrichment assay as described in fig. S5, but enriched with minocycline. Prior to enrichment incubation, cultures contained roughly 500 cells per µL as verified by control plates (50 µL culture produced ~2.5×10^4 colonies/plate, see example at left). After enrichment, the same volume of culture contained fewer CFUs in cultures containing 20 µM Mn or less (due to bactericidal activity of Amp). However, above 20 µM many cells remained viable, thus revealing dormant cells below the MIC (identified as 60 µM in batch cultures in fig. S11). The MCC for TetA-Mn interaction, defined as the minimum Mn concentration above which at least 1 % of the inoculant survives enrichment, is thus between 20 and 25 µM (between red and blue circles). This is comparable to our model prediction of MCC = 15-30 µM (end of lower branch in fig. S11, the value may be as high as 30 µM as described near equation [S55] of Supplementary Discussion).
Fig. S14: Determination of MIC\textsubscript{plate} for various CAT-expressing strains. These results identify MIC\textsubscript{plate} values of strains Cat2 through Cat6 (table S1); they are similar to those presented in fig. S2, but each strain harbors a unique level of constitutive CAT expression. The results are intended to complement the batch culture MIC determinations used in Fig. 4. For each strain shown, cells were diluted from log phase batch cultures lacking Cm and spread onto agar plates at a density of \~8\times10^4 cells per plate. Plates incubated overnight at 37°C. Chloramphenicol concentrations are indicated underneath each plate, and additionally as percentage of MIC\textsubscript{plate} for each strain (top row). MIC\textsubscript{plate} is taken as the Cm concentration above which colonies appear at a frequency of less than 42
~10^{-4}. At least two replicate experiments were performed for each strain; representative results are shown here. MIC_{plate} values were similar to those obtained with batch culture growth in minimal media with chloramphenicol (it is not uncommon to find MIC of an antibiotic in LB to correspond with MIC in minimal media (43). For the data plotted in Fig. 4B, we used only MIC data determined in batch cultures with minimal media because our model is formulated based on observations during batch culture growth (see Supplementary Discussion near equations [S5] and [S13]).
Fig. S15: Determination of MCC for various CAT-expressing strains. The Amp enrichment assay (fig. S5) was performed for CAT-expressing strains Cat2 through Cat6 to identify the MCC of each strain, representative plate results are shown here. Plates circled in red indicate that at least several percent of Amp-treated cells formed colonies, while those circled in blue indicate <1% of colony formation. The MCC value for each strain in the phase diagram of Figure 3B gives the average of at least two experiments strain (except for strain Cat4, for which we only performed the enrichment experiment once).
Fig. S16: Growth rates predicted by the measured MICs in the Cm-CAT system. To predict the growth rates of CAT-expressing strains in medium with various Cm concentrations (Fig. 4C-D), we used the value of $V_0/\kappa$ determined by MIC for strain Cat1 together with the relative CAT activities ($V_0$) measured for the different strains (bottom of Fig. 4B). Here, we instead used only the empirical MICs of the CAT-expressing strains (circles in Fig. 4B) to fix the quantity $V_0/\kappa$ for each strain with equation [S28]. This procedure is able to predict growth rates even if resistance enzyme activity is not easily assayable, e.g. in the Tc-TetA system for which the activity of drug resistance (provided by the efflux pump) is not readily assayable but for which MICs are available.
Fig. S17: Growth model without drug-induced feedback. Given the very nonlinear relation between the internal and external Cm concentrations for strongly resistant cells (Fig. 3B), we investigated whether a model without feedback, i.e., with a constant, growth-rate independent expression of CAT, might also explain the Cm-dependence of growth rate (data in Figs. 4C-3D, reproduced here as solid circles, together with Amp-enrichment data shown as open circles). The no-feedback model (Supp Discussion Eq. [S11] is used here to describe the data in this figure, using $V_{\text{max}} / \kappa$ as the only fitting parameter. The best-fit model for each strain is shown as the dashed gray line. While the no-feedback model can reasonably account for growth data of strains Cat6 and Cat5, which exhibit little or no growth bistability, the model fails to describe the growth data for the other strains that exhibit growth bistability and abrupt drops in growth rate. We fit the equation to the growth rates $\{[\text{Cm}]_{\text{ext}}, \lambda_\text{i} / \lambda_\text{r}\}$ using the Levenberg–Marquardt method with the “NonlinearFitModel” function in Mathematica; data from slow-growing or non-growing cultures ($\lambda_\text{i} / \lambda_\text{r} < 0.15$) were excluded from the fitting procedure to allow the no-feedback model the best possible chance to fit growth rates. Including the non-growing data into fits produced qualitatively similar results.

For direct comparison between presence and absence of feedback, we include a similar fit for the growth feedback model using $V_0 / \kappa$ as the lone fitting parameter (solid
colored lines), again excluding the non-growing cultures from the fit procedure; the latter clearly account for the data as a whole much better. The function used to fit the data was the implicit solution to equations [1]-[3]: \( \frac{\lambda}{\lambda_0}([\text{Cm}]_{\text{ext}}; V_0/\kappa) \), with other parameters fixed (\( K_m \) and \( I_{50} \), table S2). (These lines differ from the solid lines in Fig. 4C-D because the latter are not fits; they are predictions of the model using the value of \( V_0/\kappa \) determined, respectively, from the measured MIC of Cat1 and the relative CAT activities of strains Cat2 through Cat6. Here we show the converse: that growth rates predict MIC reasonably well, despite the exclusion of non-growing cultures from the fitting procedure. This prediction is most striking for the strains with greatest CAT expression, in which growth rates accurately predict the location and approximate magnitude of the abrupt drop in growth. See also fig. S16).
Fig. S18: Growth bistability predicted by the growth-mediated feedback model for antibiotic resistance. Model predictions for the dependence of the growth rate ($\lambda$) and the intracellular antibiotic concentration ($[a]_{\text{int}}$) on the external antibiotic concentration ($[a]_{\text{ext}}$) at steady state. The concentrations are plotted in units of $I_{50}$, i.e., $a_i = [a]_{\text{int}} / I_{50}$ and $a_e = [a]_{\text{ext}} / I_{50}$. There are two dimensionless parameters: $\rho = V_{\text{max}} / (\kappa K_m)$ is the resistance efficacy and $\sigma = I_{50} / K_m$ is the saturability (see Eqs. [S17] and [S18]). Saturability is of the order 1 for both Cm and Tc studied in this work (tables S2, S5), i.e. $K_m \sim I_{50}$ such that the enzyme activity approaches $V_{\text{max}}$ prior to complete growth inhibition. For simplicity, we use $\sigma = 1$ for all plots shown here. A, The theoretical relation between $a_i$ and $a_e$ in our model is shown for a range of resistance efficacies ($0 \leq \rho \leq 200$) to give a sense of scale of this parameter. The family of curves show the region with multiple solutions (i.e., 3 values of $a_i$ for the same $a_e$) for $\rho > \rho_c$, where $\rho_c = 27$ for $\sigma = 1$ according to Eq. [S21]. B, The same relation is shown for a strain with high resistance efficacy ($\rho = 200$). The red dashed line shows that a single extracellular concentration may map to three intracellular concentrations in the region between $a_e^{\text{min}}$ and $a_e^{\text{max}}$. However, only two are predicted to
be stable solutions; intermediate internal antibiotic concentrations (red segments throughout the figure) are kinetically unstable (Supplementary Discussion near Eq. [S42]). The intracellular concentrations corresponding to $a_e^{\min}$ and $a_e^{\max}$ are $a_i^{\dagger}$ and $a_i^{\ast}$, respectively. C, Plots of the growth rate (relative to the drug-free case, $\lambda_0$) as a function of $a_e$ for various values of resistance efficacy, $\rho$. Note that solutions for the growth rate in the grey region are predicted to be kinetically unstable and, therefore, unobservable in experiments (red segments). D, Relative growth rate as a function of external antibiotic concentration for $\rho = 200$. The growth rate at $a_e^{\min}$ and $a_e^{\max}$ are denoted as $\lambda_{\dagger}$ and $\lambda^{\ast}$, respectively (empty and filled circles); see Eqs. [S28] and [S39]. $a_e^{\max}$ is identified as MIC and $a_e^{\min}$ as MCC.
Fig. S19: Key predictions from the model and approximations in the limit of “strong” resistance. Here we illustrate key predictions relating experimentally observable quantities to biochemical parameters in the model, revealing how observables depend on strength of resistance (see Supplementary Discussion surrounding Eqs. [S16]-[S36]).

A, The critical resistance efficacy \( \rho_c(\sigma) \) gives the minimal resistance efficacy required \( (\rho > \rho_c(\sigma)) \) in order for a strain to exhibit bistability over any range of drug concentrations. Quantities (B-E) are given for the Cm-CAT resistance system with saturability parameter \( \sigma = I_{50}/K_m \approx 0.46 \), see table S2 for details. B, The predicted internal antibiotic concentration \( a_i^* \) for cells growing in antibiotic medium just below the MIC, expressed as a function of resistance efficacy in absolute dimensionless units (top axis), or in units of critical resistance efficacy \( \rho_c(\sigma) \). It is obtained as the ArgMax of equation [S16] in the supplement. C, The plot shows how \( a_e^\text{max} \) varies with resistance efficacy. \( a_e^\text{max} \) is the highest external antibiotic concentration to allow growth in a strain with strong resistance, such that \( \text{MIC} = I_{50} \cdot a_e^\text{max} \) and provides the upper bound for the region of bistability. D, The growth rate \( \lambda^* \) at \( a_e^\text{max} \) gives the minimum growth rate achievable in sub-inhibitory concentrations (i.e. for \( [a]_{\text{ext}} \) just below MIC), and its value is independent of resistance efficacy for strains with strong resistance. E, For a strain of efficacy \( \rho \), the \( a_i^* \) gives the minimum internal antibiotic concentration possible at steady state along the “low-growth” branch of the bistable region. It is obtained as the ArgMin of equation [S16] and increases with the square root of resistance efficacy. F-G, The
external antibiotic concentration for a cell with $a_i = a^\dagger_i$ is given by $a^{\text{min}}_e$ and provides the lower boundary of the bistable region. This quantity also gives the MCC in the appropriate limit (Eq. [S36]), i.e. when the growth rate at this point $\lambda^{\dagger}_e$ is small (given in $G$).
Supplementary Movie Legends

**Supplementary Movie 1:** Example of bimodal growth in a microfluidic chamber
Genetically identical Cat1m cells express the CAT protein constitutively. Prior to imaging, cells grew exponentially for two generations in the microfluidic chambers under constant flow of minimal medium lacking Cm. At time t=0 hr (time indicated bottom right), media flowing into the device was switched to the same medium containing 0.9 mM Cm. Two cells in the chambers are visible in the frame at t=0 hr; one cell continues growing and the other becomes dormant.

**Supplementary Movie 2:** Non-growing cells can resume growth after Cm-downshift
In the same experiment depicted in Supplementary Movie 1 (with 0.9 mM Cm in the medium), some chambers contained only non-growing cells; this movie shows a typical non-growing cell. After 24 hours incubation the Cm concentration was reduced from 0.9 mM to 0.1 mM, a concentration well above MIC of wild type cells (see fig. S3). Growth of this cell resumed ~10 hours after Cm-downshift, indicating that the cell remained viable and retained resistance throughout the treatment despite remaining dormant.
**Table S1: Bacterial strains used in this study.** See Supplementary Materials and Methods for details of the construction.

<table>
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<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Background</th>
<th>Comments</th>
<th>AKA</th>
</tr>
</thead>
<tbody>
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<td>ΔlacIZY</td>
<td>MG1655</td>
<td><em>E. coli</em> K-12 MG1655 with the native lac operon removed</td>
<td>—</td>
</tr>
<tr>
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<td>ΔmotA</td>
<td>EQ4</td>
<td>immotile wild type strain</td>
<td>EQ79</td>
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<tr>
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<td>$P_{Lac-01}$-lacZ</td>
<td>EQ4</td>
<td>constitutive expression of lacZ at the lac locus, driven by the synthetic $P_{Lac-01}$ promoter</td>
<td>EQ5</td>
</tr>
<tr>
<td>Lac2</td>
<td>$P_{Lac-01}$-lacZ</td>
<td>MG1655 ΔlacI, ΔgalK, ΔrhyB</td>
<td>constitutive expression of lacZ at the lac locus, driven by the synthetic $P_{Lac-01}$ promoter</td>
<td>EQ37 from ref. (16)</td>
</tr>
<tr>
<td>Ta1</td>
<td>$Km^r;rrnBT$: $P_{Lac-01}$-tetA</td>
<td>EQ4</td>
<td>constitutive expression of tetA(B) from transposon Tn10 at the ycaD locus</td>
<td>EQ93</td>
</tr>
<tr>
<td>Cat1</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS1-cat</td>
<td>EQ4</td>
<td>constitutive expression of cat by the synthetic $P_{Lac-01}$ promoter at the ycaD locus; employs “RBS1” as the Shine-Dalgarno (SD) sequence*</td>
<td>EQ75</td>
</tr>
<tr>
<td>Cat2</td>
<td>$Km^r;rrnBT$: $P_{Lac-01}$-cat</td>
<td>EQ4</td>
<td>constitutive expression of cat by the synthetic $P_{Lac-01}$ promoter at the ycaD locus; employs RBS1 as SD sequence</td>
<td>EQ92</td>
</tr>
<tr>
<td>Cat3</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS3-cat</td>
<td>EQ4</td>
<td>as Cat1, but employs “RBS3” as the SD sequence</td>
<td>EQ94</td>
</tr>
<tr>
<td>Cat4</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS4-cat</td>
<td>EQ4</td>
<td>as Cat1, but employs “RBS4” as the SD sequence</td>
<td>EQ95</td>
</tr>
<tr>
<td>Cat5</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS5-cat</td>
<td>EQ4</td>
<td>as Cat1, but employs “RBS5” as the SD sequence</td>
<td>EQ96</td>
</tr>
<tr>
<td>Cat6</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS6-cat</td>
<td>EQ4</td>
<td>as Cat1, but employs “RBS6” as the SD sequence</td>
<td>EQ98</td>
</tr>
<tr>
<td>Cat1m</td>
<td>ΔmotA, $rrnBT$: $P_{Lac-01}$-RBS1-cat</td>
<td>EQ4m</td>
<td>as Cat1, but immotile</td>
<td>EQ107</td>
</tr>
<tr>
<td>GCat1m</td>
<td>$rrnBT$: $P_{Lac-01}$-RBS1-cat, ΔmotA, $Km^r;rrnBT$: $P_{Lac-01}$-gfp</td>
<td>EQ4m</td>
<td>as Cat1m, but also harboring constitutive expression of gfp at the intC locus</td>
<td>EQ108</td>
</tr>
</tbody>
</table>

* See table S3 for ribosomal binding site “RBS” sequences
Table S2: Biochemical parameters related to the Cm-CAT system for strain Cat1

<table>
<thead>
<tr>
<th>Parameter (units)</th>
<th>Description§</th>
<th>From literature, or measured</th>
<th>Fit from data</th>
<th>Predicted†</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\lambda_0$ (hr$^{-1}$)</td>
<td>growth rate in the absence of Cm</td>
<td>$0.62 \pm 0.03^a$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$K_D$ (µM)</td>
<td>Cm-ribosome dissociation constant</td>
<td>$0.6 – 2.7^b$</td>
<td>—</td>
<td>$1.3 \pm 0.12^k$</td>
</tr>
<tr>
<td>$K_m$ (µM)</td>
<td>Michaelis constant for CAT enzyme</td>
<td>$12^c$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$k_{cat}$ (s$^{-1}$)</td>
<td>turnover number for CAT enzyme</td>
<td>$600^c$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$\kappa$ (s$^{-1}$)</td>
<td>membrane permeability to Cm</td>
<td>$0.37 – 7.5^d$</td>
<td>—</td>
<td>$1.5^l$</td>
</tr>
<tr>
<td>$V_0$ (µM/OD$_{600}$/s)</td>
<td>CAT activity in the absence of Cm</td>
<td>$3.7 \pm 0.3^e$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$I_{50}$ (µM)</td>
<td>half-inhibition conc.</td>
<td>—</td>
<td>$5.5 \pm 0.5^h$</td>
<td>$2.8 – 12^k$</td>
</tr>
<tr>
<td>$V_0 / \kappa$ (mM)</td>
<td>CAT activity relative to permeability</td>
<td>—</td>
<td>$5.8 \pm 0.5^i$</td>
<td>$0.90 – 18^m$</td>
</tr>
<tr>
<td>MIC (µM)</td>
<td>min. inhibitory conc.</td>
<td>$950 \pm 50^{fg}$</td>
<td>—</td>
<td>$950 \pm 10^n$</td>
</tr>
</tbody>
</table>

§ All in vivo quantities refer to Cat1 cells growing exponentially in minimal glucose medium with Cm unless otherwise indicated.

†These values are predicted using other parameter values listed on the two left columns; they are provided to illustrate consistency between model results and literature.

a. Growth rate of strain Cat1 in glucose minimal media in absence of antibiotics, fit from slope of log-OD versus time with standard deviation from four replicates.

b. The range of dissociation constant values reported for Cm-ribosome binding (22, 84).

c. Michaelis constant and the turnover number for CAT enzyme (23). Error for the $K_m$ value is described as “less than 10%” in the reference.

d. Membrane permeability for Cm were estimated using the methods of Nikaido (88). Briefly, permeability of Cm through cytoplasmic membrane is calculated based on Cm hydrophobicity and molecular weight (97, 98). Permeability through E. coli outer membrane is calculated using available data for permeabilities of antibiotics of similar hydrophobicity, molecular weight, size and charge (85, 102, 103). Based on simple flux balance across each membrane, an effective permeability $\kappa$ is defined in terms of cytoplasmic and outer membrane permeabilities using reported dimensions of cell volume and membrane surface area (88) (see expression [S46] and equation [S49] in supplement).
c. \( V_0 \), the CAT activity of exponentially growing Cat1 cells in Cm-free medium, is quantified using a spectrophotometric assay (77) (see Supplementary Methods). We provide the estimated in vivo activity here as acetylation rate per cytoplasmic volume by the following conversion. We divide the molar rate of Cm acetylated/OD\(_{600} \) by a factor (112) of 1.6 to convert our measurements from per-OD\(_{600} \) to per-OD\(_{460} \), and then divide by cell volume per mass (113) of 0.34 ± 0.04 µL cells/(mL culture-OD\(_{460} \)) to obtain the rate relative to equivalent cell volume in the reaction. Cell volume per mass changes little across strains or growth conditions in E. coli (113–115). We finally divide this quantity by 0.8 to take into account that cytoplasm comprises approximately 80% of total cell volume in E. coli K-12 (116, 117) to obtain the result in desired units: \( V_0 \approx 8.5 \pm 0.8 \) mM/s.

f. MIC from the abrupt growth rate drop in batch cultures. See, for example, growth curves in fig. S11.

g. MIC\(_\text{plate} \) from LB plating, see fig. S2.

h. The value of half-inhibitory concentration obtained from the fit of the growth data of wild type cells in sub-inhibitory concentrations of Cm (Fig. 3D) to equation [3], assuming \([\text{Cm}]_{\text{int}} \sim [\text{Cm}]_{\text{ext}}\). We performed a simple weighted least squares fit using weights \( w = (\text{s.e.})^2 \) for each data point, with s.e. given by the standard error of doubling time for each determination (error bars in Fig. 3D). Uncertainty in the fit parameter \( I_{50} \) is a function of the estimated variance and is reported as standard error of the regressed parameter, with estimated variance \( \sigma^2 = \sum \text{wt} \cdot (\text{fit residual})^2 / (\text{D.F.}) \).

i. Parameter \( V_0 / \kappa \) is obtained from solving equation [S28] for with the empirical (batch culture) MIC obtained for strain Cat1, with the value MIC=950 µM taken from \((f,g)\), the value of \( I_{50} = 5.5 \) µM taken from \((h)\), and \( K_m \) from \((c)\). Error represents uncertainties propagated from MIC, \( I_{50} \), and \( K_m \).

j. This is an alternative estimate obtained by fitting the growth rate data of Fig. 4A with an equation for \( \dot{\lambda}(\text{Cm}_{\text{ext}}) \) obtained from equations 1-3, with \( V_0 / \kappa \) as the only free parameter (as described in fig. S17). To fit the data independently of MIC, data from non-growing cultures were excluded from fit and we used only the data for which growth rates were non-zero. This value obtained by fitting growth rates is similar to the value predicted by MIC in \((f)\).

k. The half-inhibitory concentration is expected to be proportional to the Cm-ribosome dissociation constant as discussed in the supplementary text \((I_{50}=K_D \lambda_{\text{max}/}\lambda_{0}, \text{equation } [S5])\), independent of a strain’s CAT activity. To provide the value of \( K_D \) predicted by our model and growth data, we solve this equation for \( K_D \) using the fit value of \( I_{50} = 5.5 \) µM from \((h)\), \( \lambda_{0} = 0.67 \) hr\(^{-1} \) for WT strain EQ4 from \((a)\), and \( \lambda_{\text{max}} = 2.85 \) hr\(^{-1} \) from Scott et al\(^9\). Conversely, we also calculate an estimate for \( I_{50} \) using equation [S5] and the \( K_D \) range from the literature given in \((b)\).

l. We obtain this estimate \( \kappa = 1.5 \) s\(^{-1} \) from the \( V_0 / \kappa = 5.8 \) mM obtained by MIC \((f)\) and the in vitro value for \( V_0 \) reported in \((e)\). In Fig. 4, we obtain the \( V_0 / \kappa \) values for strains Cat2-Cat6 by assuming all strains have this same permeability to Cm. With this assumption, \( V_0 / \kappa \) for all strains are determined simply by the relative CAT activities displayed in Figure 3B (grey bars).
This estimate of the key parameter of the model uses the parameter values $V_0$ and $\kappa$ given in (e) and (d) respectively.

MIC calculated using the value of $V_0/\kappa$ obtained from fitting our model to growth rates (j) and equation [S28] from the Main Text, with the value of $I_{50} = 5.5 \mu M$ taken from (h), and $K_m$ from (c). Error propagated from standard error of fit in (j) and uncertainties of $I_{50}$ and $K_m$. 
### Table S3: RBS substitutions

<table>
<thead>
<tr>
<th>RBS ID</th>
<th>Strain</th>
<th>PLlac-O1 and RBS</th>
</tr>
</thead>
<tbody>
<tr>
<td>RBS1</td>
<td>Cat1</td>
<td>CTCGAGAATTGTAAGCGGATAACAATTTGACATTGTGAGCGGA&lt;br&gt;TAACAAGATACCTGA&lt;br&gt;GCACATCGACGGACGACTGAGCGAATTCTCATTAAGAGAGGAG&lt;br&gt;AAAGGTACCATG</td>
</tr>
<tr>
<td>RBS3</td>
<td>Cat3</td>
<td>CTCGAGAATTGTAAGCGGATAACAATTTGACATTGTGAGCGGA&lt;br&gt;TAACAAGATACCTGA&lt;br&gt;GCACATCGACGGACGACTGAGCGAATTCTCATTAAGAGAGGAG&lt;br&gt;AAAGGTACCATG</td>
</tr>
<tr>
<td>RBS4</td>
<td>Cat4</td>
<td>CTCGAGAATTGTAAGCGGATAACAATTTGACATTGTGAGCGGA&lt;br&gt;TAACAAGATACCTGA&lt;br&gt;GCACATCGACGGACGACTGAGCGAATTCTCATTAAGAGAGGAG&lt;br&gt;AAAGGTACCATG</td>
</tr>
<tr>
<td>RBS5</td>
<td>Cat5</td>
<td>CTCGAGAATTGTAAGCGGATAACAATTTGACATTGTGAGCGGA&lt;br&gt;TAACAAGATACCTGA&lt;br&gt;GCACATCGACGGACGACTGAGCGAATTCTCATTAAGAGAGGAG&lt;br&gt;AAAGGTACCATG</td>
</tr>
<tr>
<td>RBS6</td>
<td>Cat6</td>
<td>CTCGAGAATTGTAAGCGGATAACAATTTGACATTGTGAGCGGA&lt;br&gt;TAACAAGATACCTGA&lt;br&gt;GCACATCGACGGACGACTGAGCGAATTCTCATTAAGAGAGGAG&lt;br&gt;AAAGGTACCATG</td>
</tr>
</tbody>
</table>

**Modifications of the RBS region in constitutive promoters.** The -35 (TTGACA) and -10 (GATACT) elements are underlined and the transcriptional start site (+1; indicated a bolded capital “A”) and the first codon (ATG) are bolded. RBS is highlighted in cyan, nucleotide substitutions in yellow, and nucleotide insertion in magenta. RBS1=original RBS; RBS3=4 base insertion; RBS4=1-base substitution; RBS5=1-bp substitution plus 4-base insertion; and RBS6=2-bp substitution plus 4-base insertion.
Table S4: Primers used in this study

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’ to 3’)</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cat-Kpn-F</td>
<td>ATAGGTACCATGGAGAAAAAATCAGGGATATAC</td>
<td>Cloning cat into pKDT_Llac-01 or pKDT_PLet</td>
</tr>
<tr>
<td>Cat-Bam-R</td>
<td>ATTGGATCCCTTACGCCCGCCCTGCCCACCT</td>
<td>Cloning cat into pKDT_Llac-01 or pKDT_PLet</td>
</tr>
<tr>
<td>tetA-Kpn-F</td>
<td>AATGGTACCATGAATATGTTGACAAAAAGATCCGCTTG</td>
<td>Cloning tetA into pKDT_Llac-01</td>
</tr>
<tr>
<td>tetA-Bam-R</td>
<td>TATGGATCCTAAGCCTTTGCTCTCCTGTCT</td>
<td>Cloning tetA into pKDT_Llac-01</td>
</tr>
<tr>
<td>Plac-RBS1</td>
<td>CATTGGGATATATCAACGGTGGTTATATCCAGTGATTTTTTTTCCATGGTACCTTTAGGAGCTTCCTTTAATGAATTC</td>
<td>RBS region mutation in P_Llac-01</td>
</tr>
<tr>
<td>Plac-RBS3</td>
<td>CATTGGGATATATCAACGGTGGTTATATCCAGTGATTTTTTTTCCATGGTACCTTTAGGAGCTTCCTTTAATGAATTC</td>
<td>RBS region mutation in P_Llac-01</td>
</tr>
<tr>
<td>Plac-RBS4</td>
<td>CATTGGGATATATCAACGGTGGTTATATCCAGTGATTTTTTTTCCATGGTACCTTTAGGAGCTTCCTTTAATGAATTC</td>
<td>RBS region mutation in P_Llac-01</td>
</tr>
<tr>
<td>Plac-RBS5</td>
<td>CATTGGGATATATCAACGGTGGTTATATCCAGTGATTTTTTTTCCATGGTACCTTTAGGAGCTTCCTTTAATGAATTC</td>
<td>RBS region mutation in P_Llac-01</td>
</tr>
<tr>
<td>Plac-RBS6</td>
<td>CATTGGGATATATCAACGGTGGTTATATCCAGTGATTTTTTTTCCATGGTACCTTTAGGAGCTTCCTTTAATGAATTC</td>
<td>RBS region mutation in P_Llac-01</td>
</tr>
<tr>
<td>PlaclacZ-P1</td>
<td>GCATTTTAGTTTACACCCATGCACTCGATGCAGTGTATGAGTTCGAGCTTCATGACCGACTCCGAGCTTC</td>
<td>Chromosomal substitution of P_Llac-01 to lacI and PlacZYA</td>
</tr>
<tr>
<td>PlaclacZ-P2</td>
<td>CGTGTAAAAACGACCGGCAGTCACTAACCGGAAGCTTAATTAGGAGCAGCTTCAGCTTC</td>
<td>Chromosomal substitution of P_Llac-01 to lacI and PlacZYA</td>
</tr>
<tr>
<td>Lac-ver-R</td>
<td>CAACTGTGGAGGCCGCGTGAGCTGGTGGAGCTTC</td>
<td>Verification of chromosomal P_Llac-01</td>
</tr>
<tr>
<td>Cat-P1</td>
<td>AGACGCAGTGGCATATCTTCTGAAGAGCATAGCGGAAAATAGAGTTGCTTCTGTTAGCTGCGCTTC</td>
<td>Chromosomal integration of P_Llac-01-cat or P_Ltet-cat to ycaD locus</td>
</tr>
<tr>
<td>Cat-P2</td>
<td>GGTGAAAATACGCGATATCCACGGCACTCGATGAACATGCTGGCTGTGGAGCTTCAGCTGGCTTC</td>
<td>Chromosomal integration of P_Llac-01-cat or P_Ltet-cat to ycaD locus</td>
</tr>
<tr>
<td>ycaD-ver-R</td>
<td>GCCAGAGTCAACAAAGCAGGCA</td>
<td>Verification of chromosomal P_Llac-01-cat or P_Ltet-cat</td>
</tr>
</tbody>
</table>
Table S5: Biochemical parameters related to the TetA system for Tc (top) and Mn (bottom)

<table>
<thead>
<tr>
<th>Parameter (units)</th>
<th>Description</th>
<th>From literature, or measured</th>
<th>Fixed by MIC</th>
<th>Fit in model*</th>
<th>Predicted</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\lambda_0$ (hr$^{-1}$)</td>
<td>growth rate in absence of antibiotic</td>
<td>0.68 $^a$</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$K_d^{Tc}$ (µM)</td>
<td>Tc-ribosome binding dissociation constant</td>
<td>1 $^b$</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$K_m^{Tc}$ (µM)</td>
<td>Michaelis constant for TetA pump and Tc</td>
<td>10 $^c$</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>$I_{50}^{Tc}$ (µM)</td>
<td>growth half-inhibition constant for Tc</td>
<td>—</td>
<td>—</td>
<td>3.6 ± 0.4 $^d$</td>
<td>4.2 $^e$</td>
</tr>
<tr>
<td>$V_0 / \kappa^{Tc}$ (mM)</td>
<td>TetA-Tc activity relative to Tc permeability in strain Ta1</td>
<td>—</td>
<td>4.20 ± 0.30 $^f$</td>
<td>4.41 ± 0.12 $^d$</td>
<td>4.43 ± 0.08 $^g$</td>
</tr>
</tbody>
</table>

| $K_d^{Mn}$ (µM) | Mn-ribosome binding dissociation constant | 0.2 $^h$ | — | — | — |
| $K_m^{Mn}$ (µM) | Michaelis constant for TetA pump and Mn | 2 $^i$ | — | — | — |
| $I_{50}^{Mn}$ (µM) | Growth half-inhibition constant for Mn | — | — | 0.89 ± 0.1 $^j$ | 0.84 $^k$ |
| $V_0 / \kappa^{Mn}$ (µM) | TetA-Mn activity relative to Mn permeability in strain Ta1 | — | 368 ± 16 $^l$ | 334 ± 22 $^j$ | 324 ± 14 $^m$ |

$^§$ All in vivo quantities refer to Cat1 cells growing exponentially in minimal glucose medium with indicated antibiotic unless otherwise noted.

$^*$ “Fit in model” parameters show that fitting the model to empirical growth rates gives similar results to those predicted by MIC alone. The fit values for $I_{50}$ also illustrate consistency with values calculated from in vitro data.

$^†$ These values are predicted using other parameter values listed in the far left column; they are provided to illustrate consistency between model results and predictions based on literature.

$^a$ Growth rate of strain Ta1 determined in glucose minimal media batch culture in absence of antibiotics

$^b$ The dissociation constant for Tc-ribosome binding to its single high-affinity site (83, 118). Some reports indicate a lower binding affinity, however this is likely due to an
inability to control for non-saturable binding in some experiments; see (83, 119) and references therein.

c. Michaelis constant for TetA efflux of tetracycline (88).

d. Here, the half-inhibitory concentration \( I_{50} \) and TetA activity \( V_0 / \kappa \) were obtained by simultaneously fitting our model (equations [1]-[3]) to growth rate data of strain Ta1 growing in Tc, with \( K_m \) taken from (c), and \( \lambda_0 \) from (b). Errors are standard error of fit, and fitting procedures are identical to those used for Cm-CAT system described in fig. S15, except that we fit both \( V_0 / \kappa \) and \( I_{50} \) simultaneously. See also equation [S60].

The half-inhibitory concentration is expected to be proportional to the Tc-ribosome dissociation constant as discussed in the supplement (\( I_{50} = K_D \cdot \lambda_{\text{max}} / \lambda_0 \), equation [S5]), independent of a strain’s efflux activity, and is calculated here with \( K_D \) given in (b), \( \lambda_0=0.68 \text{ hr}^{-1} \) of strain Ta1 given in (a), and \( \lambda_{\text{max}}=2.85 \text{ hr}^{-1} \).

e. This parameter is obtained by solving equation [S28] for \( V_0 / \kappa \) with the empirical MIC obtained for strain Ta1 in Tc (MIC = 650 ± 50 µM, fig. S11A), the value of \( I_{50}^{\text{Tc}} \) taken from (e), and \( K_m \) from (c). Error represents uncertainties propagated from MIC.

f. Here we fit \( V_0 / \kappa \) as the only free parameter in our model to growth rate data of strain Ta1 growing in Tc with the value of \( I_{50}^{\text{Tc}} \) taken from (e), and \( K_m \) taken from (c). Errors are standard error of fit, and fitting procedures are identical to those used for Cm-CAT system described in fig. S16.

The dissociation constant for Mn-ribosome binding is reported in table 2 of Olson et al. (119). We obtain a similar value from the relative binding-affinities of minocycline and tetracycline for 30S-ribosome, given by competitive binding assays between \(^3\text{H}\)-tetracycline and each compound (table 3 in Olson et al. (119)). EC\(_{50}\) for each compound gives the concentration at which \(^3\text{H}\)-tetracycline binding is reduced by 50% due to competitive binding. Then \( K_{D}^{\text{Mn}} = \frac{\text{EC}_{50}^{\text{Mn}}}{\text{EC}_{50}^{\text{Tc}}} \cdot K_D^{\text{Tc}} = 0.2 K_D^{\text{Tc}} \), with \( K_D^{\text{Tc}} \) taken from (b).

g. Michaelis constant for TetA efflux of minocycline is taken as the concentration of minocycline required to inhibit TetA-mediated influx of \(^3\text{H}\)-tetracycline into everted membrane vesicles by 50%, reported elsewhere (120, 121).

h. The half-inhibitory concentration \( I_{50} \) and TetA activity \( V_0 / \kappa \) for Mn are obtained as in note (e), except using growth rate data from growth in Mn, with \( K_m \) taken from (i).

i. The half-inhibitory concentration \( I_{50} \) and TetA activity \( V_0 / \kappa \) for Mn are obtained as in note (e), except using growth rate data from growth in Mn, with \( K_m \) taken from (i).

j. The half-inhibitory concentration \( I_{50} \) and TetA activity \( V_0 / \kappa \) for Mn are obtained as in note (e), except using growth rate data from growth in Mn, with \( K_m \) taken from (i).

k. The half-inhibitory concentration is expected to be proportional to the Mn-ribosome dissociation constant as discussed in the supplement (\( I_{50} = K_D \cdot \lambda_{\text{max}} / \lambda_0 \), equation [S5]), independent of a strain’s efflux activity, and is calculated here with \( K_D \) given in (h).

l. This parameter is obtained by solving equation [S28] for \( V_0 / \kappa \) with the empirical MIC obtained for strain Ta1 in Mn (MIC ≈ 58 ± 3 µM, see Fig. 10B), the value of \( I_{50}^{\text{Mn}} \) taken from (k), and \( K_m \) from (i). Error represents uncertainties propagated from MIC.

m. This \( V_0 / \kappa \) was obtained by fitting our model as in note (g), except using growth rate data from growth in Mn with \( I_{50}^{\text{Mn}} \) taken from (k), and \( K_m \) from (i).
Supplementary References


76. M. Kim et al., Need-based activation of ammonium uptake in Escherichia coli, Molecular Systems Biology 8, 616 (2012).


