Environmental perturbations lift the degeneracy of the genetic code to regulate protein levels in bacteria

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The genetic code underlying protein synthesis is a canonical example of a degenerate biological system. Degeneracies in physical and biological systems can be lifted by external perturbations, thus allowing degenerate systems to exhibit a wide range of behaviors. Here we show that the degeneracy of the genetic code is lifted by environmental perturbations to regulate protein levels in living cells. By measuring protein synthesis rates from a synthetic reporter library in Escherichia coli, we find that environmental perturbations, such as reduction of cognate amino acid supply, lift the degeneracy of the genetic code by splitting codon families into a hierarchy of robust and sensitive synonymous codons. Rates of protein synthesis associated with robust codons are up to 100-fold higher than those associated with sensitive codons under these conditions. We find that the observed hierarchy between synonymous codons is not determined by usual rules associated with tRNA abundance and codon usage. Rather, competition among tRNA isoacceptors for aminoacylation underlies the robustness of protein synthesis. Remarkably, the hierarchy established using the synthetic library also explains the measured robustness of synthesis for endogenous proteins in E. coli. We further found that the same hierarchy is reflected in the fitness cost of synonymous mutations in amino acid biosynthesis genes and in the transcriptional control of σ -factor genes. Our study suggests that organisms can exploit degeneracy lifting as a general strategy to adapt protein synthesis to their environment.

codon bias \mid selective charging \mid translation efficiency \mid starvation \mid codon optimization

Degeneracy, the occurrence of distinct states that share a common function, is a ubiquitous property of physical and biological systems (1-3). Examples of degenerate systems include atomic spectra (4), condensed matter (5), the nervous system (2), and the genetic code (6, 7). Degeneracy in physical systems is often associated with underlying symmetries (1) and in biological systems with error minimization, evolvability, and robustness against perturbations (8). Degenerate states that are indistinguishable under normal conditions can exhibit distinct properties under the action of external perturbations (1). This effect, called degeneracy lifting, allows degenerate systems to exhibit a wide range of behaviors, depending on the environmental context (2). The genetic code governing protein synthesis is a highly degenerate system because 18 of the 20 amino acids have multiple synonymous codons and 10 of the 20 amino acids are aminoacylated (charged) onto multiple tRNA isoacceptors. Protein synthesis rates in living cells respond to diverse environmental perturbations, which raises the question of whether any of these perturbations modulates protein levels by lifting the degeneracy of the genetic code. Previous experiments found that both the concentration of charged tRNAs and the occupancy of ribosomes on synonymous codons undergo significant changes upon nutrient limitation (9-11). However, whether such environmental perturbations lift the degeneracy of the genetic code by modulating the expression level of proteins is unknown. Here, we propose to use amino acid limitation in the bacterium *Escherichia coli* as a model system to investigate whether the degeneracy of the genetic code can be lifted by environmental perturbations and how degeneracy lifting could provide a general strategy to adapt protein synthesis to environmental changes.

Results

Degeneracy Lifting upon Amino Acid Limitation. We considered synonymous codons for seven amino acids: Leu, Arg, Ser, Pro, Ile, Gln, and Phe. This set of seven amino acids is representative of the degeneracy of the genetic code, in that it includes six-, four-, three-, and twofold degenerate codon families. We constructed a library of 29 *yellow fluorescent protein (yfp)* gene variants, each of which had between six and eight synonymous mutations for one of the seven amino acids (Fig. 1A). In this library, we designed each *yfp* variant to characterize the effect of one specific codon on protein synthesis. We expressed the *yfp* variants constitutively at low gene dosage (two copies per chromosome, Fig. 1B) in *E. coli* strains that were auxotrophic for one or more amino acids. We monitored growth and YFP synthesis in these strains during amino acid-rich growth as well as during limitation for each of the seven amino acids (*Materials and Methods*).

During amino acid-rich growth, our measurements revealed that protein synthesis rates were highly similar across yfp variants, with less than 1.4-fold variation within all codon families (Fig. 1D, light gray bars). Thus, under rich conditions, the degeneracy of the genetic code remains intact with respect to protein synthesis. Strikingly, under amino acid-limited growth, codon families split into a hierarchy of YFP synthesis rates (Fig. 1 C and D). We found that some synonymous codons, such as CTA for leucine, were highly sensitive to environmental perturbation, causing YFP synthesis rates to be near zero in response to the limitation of these codons' cognate amino acids. Conversely, other synonymous codons, such as CTG for leucine, were more robust to the same perturbation with synthesis rates of YFP up to 100-fold higher than those of the sensitive ones. We define codons as robust when the synthesis rate from the corresponding yfp variant during cognate amino acid limitation is higher than the average synthesis rate within that codon family. Similarly, we define codons as sensitive when the synthesis rate from the corresponding *yfp* variant during cognate amino acid limitation is lower than the average synthesis rate within that codon family. In addition to fluorescence, the difference in robustness was reflected in protein levels measured with Western blotting (SI Appendix, Fig. S1). Notably, even a single substitution to a perturbation-sensitive codon in the yfp coding sequence resulted in more than a twofold difference in YFP synthesis rate during limitation for the cognate amino acid,

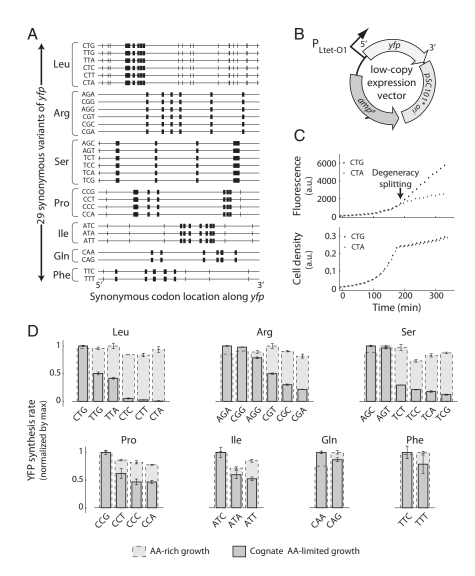
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without any effect on synthesis rate during amino acid-rich growth (*SI Appendix*, Fig. S2). Only those codons that were cognate to the limiting amino acid caused splitting of YFP synthesis rates (*SI Appendix*, Fig. S3). Interestingly, the splitting was more acute for codon families with sixfold degeneracy (Leu, Arg, and Ser), whereas splitting was weaker for codon families with four-, three-, and twofold degeneracies (Fig. 1D, *Upper* row vs. *Lower* row). These results support the idea that greater degeneracy typically allows systems to exhibit a wider range of responses to environmental perturbations (2). In subsequent experiments, we focused on the two codon families, leucine and arginine, that displayed the largest range of splitting. These two families constitute 16% of codons across the genome of *E. coli*.

Intracellular Determinants of the Hierarchy Among Synonymous Codons. We sought to identify the intracellular parameters that determine the observed hierarchy of degeneracy splitting during amino acid limitation. To this end, we quantified the robustness of synthesis rate to amino acid limitation as the ratio of YFP synthesis rates between amino acid-limited and amino acid-rich growth phases. Protein synthesis rate is known to be correlated with codon usage and tRNA abundance during artificial overexpression of proteins (12, 13). However, we found that robustness of YFP synthesis to amino acid limitation was not correlated with either codon usage or tRNA abundance (r^2 =

Fig. 1. Degeneracy lifting associated with amino acid limitation. (A) A library of 29 variants of the yellow fluorescent protein gene (yfp) was synthesized. In this library, each variant (represented as a horizontal line) was designed to measure the effect of one specific codon on protein synthesis rate. The identity of this codon and that of its cognate amino acid is indicated to the left of each yfp variant, and the locations of this codon along yfp are represented as thick vertical bars. Other codons for the same amino acid that were identical across all yfp variants in each codon family are represented as thin vertical bars. (B) Each yfp variant was constitutively expressed from a low-copy vector (SC101* ori, two copies per chromosome) in E. coli strains that were auxotrophic for one or more of seven amino acids. (C) To induce amino acid-limited growth, we adjusted the initial concentration of an amino acid in the growth medium to a level below that required for reaching saturating cell density. A methyl-ester analog of the amino acid supported steady growth in the amino acid-limited phase. Growth and fluorescence curves for two yfp variants, CTA, gray, and CTG, black, are shown as illustrative examples of degeneracy splitting upon limitation for the cognate amino acid, leucine. (D) Dark gray, YFP synthesis rates during limitation for cognate amino acid; light gray, YFP synthesis rates during amino acid-rich growth. YFP synthesis rate was defined as the rate of fluorescence change divided by the cell density. Synthesis rates were normalized by the maximum value within each synonymous codon family and separately in the amino acid-rich and amino acid-limited growth phases. Normalization factors (amino acidrich, limited): Leu, 94, 81; Arg, 89, 113; Ser, 217, 343; Pro, 306, 49; Ile, 295, 45; Gln, 185, 83; Phe, 311, 20 (arbitrary units). Error bars show standard error over three replicate cultures.

0.08 and 0.00, respectively, squared Spearman's rank correlation; SI Appendix, Fig. S4). We then considered determinants of protein synthesis that might be important specifically during amino acid limitation. tRNA isoacceptors are uniformly charged at about 80% under amino acid-rich conditions (14, 15). However, during perturbations such as amino acid limitation, some tRNA isoacceptors cognate to the amino acid are almost fully charged whereas other isoacceptors in the same family have charged fractions that are close to zero (10, 16). A theoretical model proposed that such selective charging arises from differences in the relative supply and demand for charged tRNA isoacceptors (9). Although it is unclear how this mechanism could solely control protein levels, charged tRNAs play an essential role as substrates for the elongation by ribosomes across individual codons (17). Consequently, we hypothesized that selective charging of tRNA isoacceptors also underlies the observed splitting in synthesis rates among yfp variants. Consistent with this hypothesis, charged fractions of leucine and arginine tRNA isoacceptors during limitation of cognate amino acid starvation measured in a previous work (10) were correlated with the robustness of synthesis rates from yfp variants after accounting for codon–tRNA assignments ($r^2 = 0.78$; SI Appendix, Fig. S5).

We experimentally tested whether varying the concentration of charged tRNA could change the hierarchy of protein synthesis

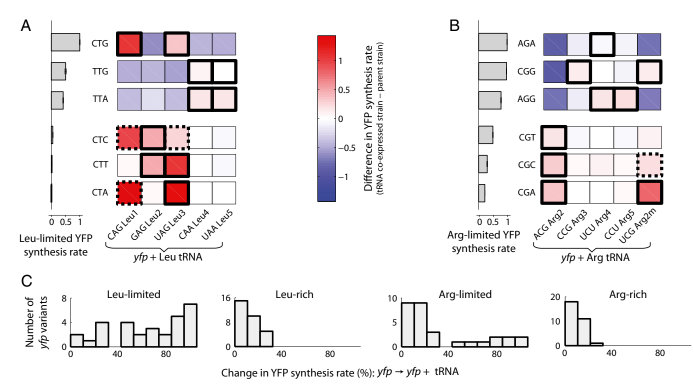


Fig. 2. Altering the hierarchy of degeneracy splitting among synonymous codons. The five leucine (arginine) tRNA isoacceptors were coexpressed together with each of the six leucine (arginine) yfp variants, resulting in 30 tRNA-yfp combinations for leucine (arginine). (*A* and *B*) Each square in the *Left (Right)* table corresponds to the difference in YFP synthesis rates of each yfp variant between the tRNA coexpressed strain and the parent strain without extra tRNA during leucine (arginine) limitation. YFP synthesis rates were defined in the same manner and normalized by the same factor as in Fig. 1*D*. (*Left*) YFP synthesis rate of the parent strain without extra tRNA during amino acid limitation is shown for each table (same data as in Fig. 1*D*). tRNA isoacceptor names are preceded by their unmodified anticodon sequences. Solid black-outlined squares correspond to codon–tRNA pairs that satisfy wobble-pairing rules after accounting for known posttranscriptional tRNA modifications (*SI Appendix*, Table S9). Dashed black-outlined squares correspond to codon–tRNA pairs that soacceptor. ^{UCG}Arg2_m is a nonnative arginine tRNA that was created by mutating the anticodon sequence of the ^{ACG}Arg2 gene. Standard error was less than 0.05 for all squares. (C) Histogram of differences in YFP synthesis rate of *yfp* variants upon tRNA coexpression. Amino acid-limited growth, 42% median difference; amino acid-rich growth, 9% median difference; (*n* = 60, aggregated for leucine and arginine). Change in YFP synthesis rate between each tRNA coexpressed strain and its parent strain of the largest value between the two YFP synthesis rates.

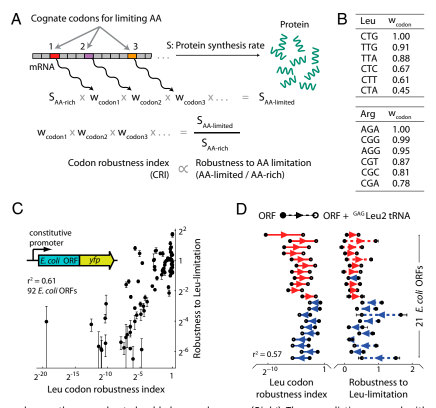
rates initially revealed by amino acid limitation. To this end, we coexpressed each one of the leucine or arginine tRNA isoacceptors together with each of the six leucine or arginine variants of yfp, respectively (Fig. 2). Previous work (16) showed that overexpression of a single tRNA isoacceptor cognate to a limiting amino acid enables it to compete better in the common charging reaction against other isoacceptors. As a result, charged tRNA concentration of the overexpressed isoacceptor increases, whereas charged tRNA concentrations of the remaining isoacceptors for that amino acid decrease or remain unchanged (16). We found that yfp variants constructed with perturbation-sensitive codons exhibited higher synthesis rates upon coexpression of tRNA isoacceptors cognate to those perturbation-sensitive codons (Fig. 2 A and B, Lower three rows, solid black-outlined squares). Conversely, yfp variants with perturbation-robust codons exhibited lower protein synthesis rates upon coexpression of noncognate tRNA isoacceptors (Fig. 2 A and B, Upper three rows, nonoutlined squares). These two patterns of changes in YFP synthesis rate mirror previously measured changes in charged tRNA concentration upon tRNA coexpression (16), thereby suggesting that the observed hierarchy in synthesis rates of yfp variants is tightly coupled with the concentrations of cognate charged tRNA isoacceptors during amino acid limitation. By contrast, tRNA coexpression had little effect on synthesis rates from yfp variants in the absence of perturbation, i.e., during amino acid-rich growth (Fig. 2C). We observed several codon-tRNA pairs with mismatches at the wobble position but that do not

satisfy known wobble-pairing rules (*SI Appendix*, Table S9) and that showed an increase in YFP synthesis rate upon coexpression of the tRNA isoacceptor during amino acid limitation (Fig. 2 *A* and *B*, dashed black-outlined squares).

Codon Robustness Index for Endogenous Proteins. We investigated whether the hierarchy of synthesis rates measured for the synthetic yfp variants also governs the synthesis of endogenous proteins of E. coli. We first devised a general parameter, hereafter called the codon robustness index (CRI), to characterize the robustness of any protein's synthesis rate to an environmental perturbation associated with limitation of a specific amino acid (Fig. 3A). We defined CRI as a product of codonspecific weights w_{codon} , and we inferred these weights from the synthesis robustness of yfp variants to limitation for their cognate amino acid (Fig. 3B). Our formulation of CRI is based on the simplifying assumption that each codon decreases protein synthesis rate by a factor w_{codon} that is independent of the codon's intragenic location, the presence of other codons in the coding sequence, or the specific cellular role of the encoded protein. By definition, $w_{\rm codon}$ is unity for codons that are not cognate to the limiting amino acid, and perturbation-robust codons have a higher w_{codon} value than perturbation-sensitive codons for the limiting amino acid.

To test the predictive power of the CRI, we selected 92 *E. coli* ORFs that span a broad range of leucine CRI values and functional categories (*SI Appendix*, Fig. S7 and Table S1). We expressed the corresponding proteins constitutively as N-terminal

Fig. 3. Degeneracy lifting for endogenous proteins. (A) The effect of each codon on the synthesis rate, S, of a protein during amino acid limitation was modeled by a codon-specific weight, w_{codon} . The codon robustness index (CRI) for any protein-coding sequence was defined as the product of w_{codon} values for all codons in that sequence that are cognate to the limiting amino acid. (B) w_{codon} values for leucine and arginine codons during limitation for their cognate amino acids were estimated from protein synthesis rates of the corresponding yfp variants (Materials and Methods). w_{codon} values for all codons not cognate to the limiting acid were set to 1. (C) Ninety-two ORFs from the E. coli genome were cloned as N-terminal fusions to YFP downstream of a constitutive promoter into a low-copy vector (Inset and Materials and Methods), Robustness to leucine limitation is quantified as the ratio of protein synthesis rates between leucine-limited and leucine-rich growth phases. This measured robustness was correlated with estimated Leu CRI values for the 92 ORF-yfp fusions ($r^2 = 0.61$, squared Spearman's rank correlation. $P = 10^{-20}$). Eleven ORFs had measured robustness below the lower limit of the vertical axis (SI Appendix, Table S1), but were included in the calculation of r^2 . Protein synthesis rates were normalized by the synthesis rate for the CTG variant of yfp. Error bars show standard error over three replicate cultures. (D) Two sets of ORF-yfp fusions (21 total ORFs) were coexpressed with GAGLeu2 tRNA. (Left) On the basis of the yfp data (Fig. 2A), we estimated a higher CRI for the first set (11 ORFs) and a lower CRI for the second set (10 ORFs) upon GAGLeu2 coexpression (Materials and Methods). Hence we predicted that the first set should show an increase in ro-



bustness of protein synthesis during leucine limitation whereas the second set should show a decrease. (*Right*) These predictions agreed with measured changes for 20 of the 21 ORFs ($r^2 = 0.57$, $P = 10^{-4}$). Error bars show standard error over three replicate cultures. Several error bars are smaller than data markers.

fusions with YFP in an E. coli strain auxotrophic for leucine (Fig. 3C, Inset). The YFP fusion partner was encoded by the CTG variant of *yfp* that has the highest, most robust synthesis rate during leucine limitation. Upon leucine limitation, we found a strong correlation between the robustness of protein synthesis rates from the 92 ORF-*yfp* fusions and their leucine CRI values (Fig. 3*C*, $r^2 = 0.61$, $P = 10^{-23}$, squared Spearman's rank correlation). Similarly, the arginine CRI was also strongly correlated with robustness of a library of 56 ORF-vfp fusions during arginine limitation ($r^2 = 0.59$, $P = 10^{-12}$; *SI Appendix*, Fig. S8 and Table S2). By contrast, standard measures of translation efficiency under amino acid-rich conditions such as codon adaptation index (18), tRNA adaptation index (19), or folding energy of the mRNA around the start codon (20) displayed only a weak correlation with protein synthesis rate from the ORF-yfp fusions during amino acid-rich growth ($r^2 = 0.10, 0.08$, and 0.02, resp.; *SI Appendix*, Fig. S9). We further found that changes in Leu CRI calculated from the yfp data could predict both the effect of tRNA coexpression and that of synonymous mutations on protein synthesis from E. coli ORFs during leucine limitation (Fig. 3D and SI Appendix, Fig. S10). Importantly, similar to our results using yfp reporters, neither tRNA coexpression nor synonymous mutations for E. coli ORF-yfp fusions had a significant effect on the synthesis rates from these ORFs during leucine-rich growth in the absence of environmental perturbations (SI Appendix, Fig. S11). Thus, the degeneracy of the genetic code underlies the levels of endogenous protein production only during response to environmental perturbations.

Consequences of Degeneracy Lifting for Fitness and Gene Regulation. Degeneracy splitting in physical systems can be exploited to encode information related to the environmental context (21, 22). We asked whether bacteria might similarly exploit the degeneracy splitting of genetic code during response to amino acid limitation. Hence we tested whether the expression of amino acid biosynthesis genes that enable bacteria to adapt to amino acid limitation is affected by the hierarchy between robust and sensitive codons. We found that mutating codons that are perturbation robust to those that are perturbation sensitive in the leucine-biosynthesis genes leuA, leuC, and leuD and the arginine-biosynthesis gene carA decreased their protein synthesis rate during cognate amino acid limitation, but not during amino acid-rich growth (SI Appendix, Fig. S12). Interestingly, in the case of *leuA* and *carA*, the same synonymous mutations also resulted in a fitness cost for prototrophic strains upon downshift from amino acid-rich to amino acidpoor conditions (Fig. 4A). Thus, synonymous mutations can have a significant fitness cost during an environmental perturbation, which is distinct from that measured under nutrient-rich conditions in the absence of any perturbation (20, 23). However, swapping codons that are perturbation robust with those that are perturbation sensitive in other biosynthesis genes (argA and leuC in Fig. 4A) did not significantly affect fitness, suggesting that the hierarchy of robust and sensitive codons might be selectively used by bacteria to regulate genes within a single metabolic pathway.

Perturbations associated with amino acid limitation in *E. coli* can result in two distinct outcomes, depending on the environmental conditions: On one hand, when substrates used in amino acid biosynthesis are still abundant in the environment, the cell up-regulates corresponding biosynthesis genes to mitigate the limitation of amino acids and resume growth. On the other hand, in the absence of substrates for amino acid biosynthesis, *E. coli* can survive a prolonged period in amino acid-poor environments through a cellular response mediated by σ -factors (24, 25). We found that genes encoding several stress-response σ -factors (*rpoS*, *rpoE*, and *rpoH*) are enriched in TTA and TTG, the leucine

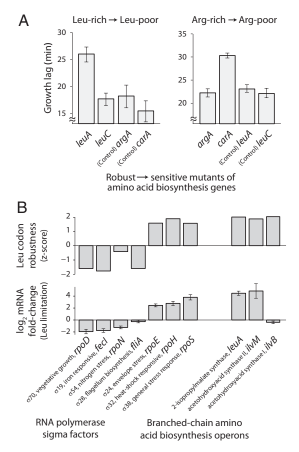


Fig. 4. Fitness cost and transcriptional control reflect degeneracy lifting. (A) Four different prototrophic E. coli strains were created. Each of these strains had one of the four amino acid biosynthesis genes argA (Arg), carA (Arg), leuA (Leu), and leuC (Leu) replaced at the native locus by a corresponding synonymous mutant ORF. These mutants were designed such that three to five perturbation-robust codons in a wild-type ORF were replaced by perturbationsensitive codons in the mutant ORF (SI Appendix, Fig. S12B). The strains were grown in medium supplemented with all 20 amino acids at 800 μ M and then diluted into a medium lacking either leucine (Left) or arginine (Right). Growth lag was calculated as the time taken by each strain to reach OD_{600} of 0.3 relative to a reference culture of the same strain grown in 800 μ M of all 20 amino acids. (Left) Difference in growth lag between the leuA mutant and the two controls during leucine downshift was 9.2 \pm 2.8 min, P = 10⁻³. (Right) Difference in growth lag between the carA mutant and the two controls during arginine downshift was 7.8 \pm 1.2 min, $P = 10^{-6}$. Standard errors were calculated over six biological replicates for each mutant. P values were calculated using a two-tailed t test between the leuA or the carA mutant and the corresponding controls. (B) (Upper) Genes encoding o-factors and leucine biosynthesis genes in E. coli are biased in their Leu CRI values, as quantified using a z-score that measures the normalized deviation from the expected CRI value based on genome-wide codon frequencies (SI Appendix). The most frequent leucine codon CTG was excluded in this analysis because its frequency varies significantly with expression level under nutrient-rich conditions (38). (Lower) Fold change in mRNA abundance in response to leucine limitation for o-factor genes and leucine biosynthesis operons was measured using RT-qPCR. Fold change of the gapA gene was used for internal normalization. Error bars show standard error over triplicate gPCR measurements.

codons that ensure robust protein synthesis during leucine limitation (Fig. 4B, Upper). By contrast, genes for the housekeeping σ -factor (*poD*) and a few minor σ -factors (*fecI*, *fliA*) are enriched for CTC and CTT, which are sensitive to leucine limitation. This contrasting pattern is observed for leucine (but not for arginine) and is further mirrored by the change in transcript abundance for σ -factor genes in response to leucine limitation (Fig. 4B, Lower). Hence degeneracy splitting in the genetic code might be exploited in concert with transcriptional control to regulate protein levels.

Discussion

In summary, we have found that the degeneracy of the genetic code does not have a role in regulating protein synthesis during amino acid-rich growth. By contrast, the splitting of this degeneracy upon reduction in amino acid supply has a potent effect on protein synthesis that results in up to 100-fold differences in protein synthesis rates between synonymous gene variants. Such a large role for synonymous codons in protein synthesis is surprising given that other posttranscriptional mechanisms such as protein degradation are known to play a significant role upon amino acid limitation (26). We identified competition between tRNA isoacceptors for aminoacylation as a key determinant of the hierarchy of protein synthesis rates during amino acid limitation. Low concentration of a charged tRNA isoacceptor can cause ribosomes to selectively pause at its cognate codon and trigger ribosome jamming (27), translation recoding (28), mRNA cleavage (29-31), or feedback-transcriptional control (32, 33). A recent genome-wide study found increased ribosome pausing at serine codons during serine-limited growth of E. coli. Interestingly, ribosomes paused significantly only at four of the six serine codons, and these four codons are precisely the same ones that caused YFP synthesis rate to be sensitive to serine limitation in our experiments (SI Appendix, Fig. S13). We measured the change in mRNA levels of different *yfp* variants in response to amino acid limitation. Changes in mRNA levels were correlated with corresponding changes in YFP synthesis rates upon amino acid limitation (SI Appendix, Fig. S14). However, changes in mRNA levels were smaller than expected, suggesting that changes in mRNA abundance induced by ribosome pausing might not be solely responsible for the observed changes in protein synthesis rate.

Here, we have investigated the effect of a specific environmental perturbation associated with amino acid limitation in the bacterium *E. coli*. However, this type of perturbation plays a crucial role in the life cycle of other bacteria such as *Myxococcus xanthus* and *Bacillus subtilis* that undergo differentiation cued by amino acid limitation (34, 35). Protein synthesis during such differentiation events might also be regulated by degeneracy lifting of the genetic code. Moreover, degeneracy lifting could be important during protein synthesis in eukaryotes, where clinically important conditions such as neoplastic transformation and drug treatment are often accompanied by a reduction in amino acid supply (36, 37). Therefore, lifting the degeneracy of the genetic code might emerge as a general strategy for biological systems to expand their repertoire of responses to environmental perturbations.

Materials and Methods

A summary of key methods is given below. Detailed methods for all experiments and analyses are included in *SI Appendix*.

Bacterial Strains. All strains used in this study were obtained from the *E. coli* Genetic Stock Center (CGSC), Yale University. Different auxotrophic strains were used, depending on the amino acid that was limiting in the growth medium (*SI Appendix*, Table S5).

Plasmids. The *pZ* series of plasmids (39) was used for expression of all genes constructed for this study. A low-copy plasmid, *pZS*11* [*SC101* ori* (3–4 copies per cell), Amp^R (*bla* gene), and a constitutive $P_{LtetO-1}$ promoter] was used for expression of all fluorescent reporter genes and their fusions. A medium-copy plasmid, *pZA32* [*p15A ori* (10–12 copies per cell), Chl^R (*cat* gene), and a $P_{LlaCO-1}$ promoter] was used for expression of all fluorescent.

Gene Synthesis and Cloning. A single *yfp* sequence was built de novo (synthesis by Genscript). All subsequent *yfp* variants were constructed using a sitedirected mutagenesis kit (Stratagene). tRNA genes and *E. coli* ORFs were amplified from the chromosome of wild-type *E. coli* MG1655 by PCR (Datasets S1 and S2). Amino Acid Limitation Experiments. Overnight cultures were inoculated from glycerol stocks or fresh colonies and grown in a MOPS-based rich-defined medium with 800 μ M of 19 amino acids and 10 mM serine at 30 °C with shaking. For experiments involving amino acid limitation, the amino acid whose limitation was to be induced was added at a reduced concentration and supplemented with its methyl-ester analog (40, 41) (*SI Appendix*, Table S6, Figs. S15 and S16). Growth and fluorescence were quantified using a standard 96-well plate reader integrated with a robotic system.

Analysis of Cell Density and Fluorescence Time Series. Matlab R2009 (Math-Works) was used for all analyses unless otherwise mentioned. All correlations and *P* values reported in this work were calculated using the Matlab command "corr". Growth and fluorescence time series were fitted with exponential and linear curves in the amino acid-rich and amino acid-limited growth regimes, respectively. Protein synthesis rate, *S* was calculated as

Protein synthesis rate, $S = \frac{1}{\text{Absorbance}} \times \frac{d(\text{Fluorescence})}{d(\text{time})}$.

Calculation of CRI. The CRI for a protein-coding sequence corresponding to a limiting amino acid was calculated by multiplying the w_i values for

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codons cognate to the limiting amino acid in that sequence. w_i values shown in Fig. 3*B* were calculated using the robustness of protein synthesis of the corresponding *yfp* variants during cognate amino acid limitation (Fig. 1*D*). On the basis of our noncognate limitation experiment (*SI Appendix*, Fig. 52), the w_i values for all codons other than those cognate to the limiting amino acid are set to be equal to 1. Revised w_i values based on *yfp* measurements in the presence of ^{GAG}Leu2 tRNA (Fig. 2) were used for calculation of the Leu CRI in the case of ^{GAG}Leu2 tRNA coexpression with *E. coli* ORFs (Fig. 3*D*).

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Appendix: Environmental perturbations lift the degeneracy of the genetic code to regulate protein levels in bacteria

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Bacterial strains

All strains used in this study were obtained from the *E.coli* Genetic Stock Center (CGSC), Yale University. For amino acid limitation experiments, standard auxotrophic strains (Table S5) were used depending on the amino acid that was limiting in the growth medium, unless mentioned otherwise. Strain CP78 was used for experiments involving leucine and arginine limitation. This strain has been used extensively in previous amino acid limitation studies [1, 2, 3, 4, 5, 6] and its multiple auxotrophy makes it a convenient choice for experiments involving limitation for several amino acids. The auxotrophic strains corresponding to the remaining amino acids are from the Keio-knockout collection [7], and are the commonly used auxotrophic strains for that amino acid (http://cgsc.biology.yale.edu/Auxotrophs.php).

For the growth lag measurements in Fig. 4A, the prototrophic strain MG1655 (Table S5) was used as the wild-type background. This background strain was tagged with yfp or rfp at the $attB\lambda$ locus (this tagging was a remnant from earlier experiments not related to this work, and has no relevance to any results presented here). Site-directed mutagenesis was used to create the synonymous mutant coding sequences for leuA, leuC, leuD, carA, argA and argF using the protocol described in the section on gene synthesis and cloning. Then to insert these mutant ORFs into their native locus without any additional markers, a two-step strategy based on λ Redmediated homologous recombination [8] was used: In the first step, the respective wild-type ORF was replaced by a kan resistance gene, and in the second step the kan gene was replaced by the mutant ORF without any additional markers by selecting on M9-glucose plates for prototrophy of the respective amino acid. Plasmid pSIM5 [9] was used as the helper plasmid and a previously published recombineering protocol [9] was used without any modifications.

For RT-qPCR (Fig. 4B), a leucine auxotroph of MG1655 was created by deleting the leuB gene using the λ Redmediated homologous recombination protocol outlined above. For Western blots (Fig. S1), the auxotrophic strains in Table S5 were further modified by insertion of the *tet* repressor gene at the $attB\lambda$ site using a previous method based on λ integrase-mediated site-specific recombination [10]. The presence of Tet repressor enabled inducible control of YFP expression. The Western blots for leucine and arginine yfp variants were performed in an MG1655 auxotroph strain background instead of the CP78 strain. The CP78 strain has lower transformation efficiency which prevented integration of the *tet* repressor gene into the chromosome. Strains were stored as 20% glycerol stocks at -80° C either in 1ml cryo-vials or in 96-well plates (3799, Costar). In addition, for experiments involving over 25 strains, a temporary 20% glycerol stock was stored at -20° C in 96-well PCR plates.

Plasmids

The pZ series of plasmids [10] were used for extrachromosomal expression of genes. General features of the plasmid backbones are described here. Specific gene constructs that were cloned into these backbones is described in the section on gene synthesis and cloning. A low-copy plasmid, pZS^*11 [SC101* ori (3-4 copies/cell), AmpR (bla gene) and a constitutive P_L tetO1 promoter] was used for expression of all fluorescent reporter genes and their fusions. The synthetic ribosome binding site (RBS) in the original pZS^*11 backbone was replaced by a modified T7-based RBS that resulted in efficient protein expression from most coding sequences. A medium-copy plasmid, pZA32 [p15A ori (10-12 copies/cell), ChlR (cat gene) and P_L lacO1 promoter] was used for expression of all tRNA genes. Strains with pZA32 plasmids were grown with 1mM IPTG to ensure constitutive expression of all tRNA genes. Standard plasmids pUC18 and pUC19 (Invitrogen) were used as intermediate cloning vectors for sitedirected mutagenesis. Plasmid pSIM5 (13) was used as the helper plasmid expressing the λ -Red system for all chromosomal modifications in this project (except for Tet repressor insertion mentioned in the previous section).

Growth and fluorescence measurements

Overnight cultures were inoculated either from freshly grown single colonies or, in experiments involving more than 25 strains, from temporary glycerol stocks stored at -20C. Overnight cultures were grown in a modified MOPS richdefined medium [11] made with the following recipe: 10X MOPS rich buffer, 10X ACGU nucleobase stock and 100X 0.132M K2HPO4 (Teknova, Cat. No. M2105) were used at 1X final concentration as in the original recipe. In addition, the overnight growth medium contained 0.5% glucose as carbon source, 10^{-4} % thiamine and 800μ M of 19 amino acids and 10mM of serine. pH was adjusted to 7.4 using 1M NaOH and appropriate selective antibiotics $(50\mu g/m)$ ampicillin and/or $20\mu g/ml$ chloramphenicol) were added. Amino acids, glucose, thiamine and antibiotics were purchased from Sigma. 1ml overnight cultures were grown in 2ml deep 96-well plates (40002-014, VWR) at 30° C with shaking at 1350rpm (Titramax 100 shaker) for 12 to 16 hours.

For amino acid limitation experiments, overnight cultures were diluted 1:1000 into 1ml of the same MOPS rich-defined medium as the overnight cultures. However the amino acid whose limitation was to be induced was added at a reduced concentration and supplemented with its methyl ester analog (Table S6). Amino acid methyl esters are analogs of the corresponding amino acids and have been previously used for steady growth of E. coli under amino acid limiting conditions [12, 13] (see Figs. S15 and S16 for the effect of methyl ester on growth and robustness of YFP synthesis). Addition of the methyl esters results in a steady but limiting supply of the amino acid due to slow hydrolysis of the ester (see Note 1). Concentrations of the amino acid and its methyl ester were chosen such that the cultures consumed the limiting amino acid and entered amino acid-limited growth at an OD_{600} of 0.6-0.7 (corresponding to an OD_{600} value of 0.2-0.25 in our 96-well plate reader). Slight variations in the initial concentration of either the limiting amino acid or its methyl ester shift the transition to a higher or lower cell density without appreciable changes in growth rate (see Note 2). Except for a single limiting amino acid, the remaining 19 amino acids were present at the overnight culture concentrations during the amino acid limitation experiments. For proline limitation, no proline was necessary in the growth medium since proline methyl ester supported growth at the same rate as proline until the OD_{600} reached around 0.6.

Diluted overnight cultures were grown in 2ml deep 96-well plates for 3 hours at 30°C with shaking at 1350rpm (Titramax 100 shaker). After this time interval, 3 aliquots of 150μ l from each culture was pipetted into 3 wells of 3 different 96-well plates (3799, Costar). Wallac Victor2 plate reader (PerkinElmer) was used to monitor cell density (absorbance at 600nm) and YFP synthesis (fluorescence, excitation 504nm and emission 540nm). Each plate was read every 15 min using a robotic system (Caliper Life Sciences) and shaken in between readings (Variomag Teleshake shaker) for a total period of 6-10 hours. Temperature of 30°C and relative humidity of 60% was maintained throughout the experiment.

In the case of experiments without methyl ester (Figs. S15 and S16), the same protocol mentioned above was followed but the methyl esters were not added to the growth medium. For the RT-qPCR measurements shown in Fig. 4B, overnights cultures were diluted 1:1000 into the same medium. Then when the OD_{600} reached 0.5, the cells were spun down at 3000g for 5 min and then re-suspended in the same medium but either with or without leucine. Total RNA was extracted (see protocol below) after 30 min of shaking at 30°C, 200rpm.

For the growth lag measurements shown in Fig. 4A, overnight cultures of prototrophic strains were diluted 1:200 into medium either with or without one of leucine and arginine. Growth lag was measured as the difference in time taken to reach OD_{600} of 0.3 between two cultures of the same strain – one growing in the presence of either leucine or arginine and another growing in its absence.

Gene synthesis and cloning

All gene sequences constructed for this study are provided in the gene_sequences.fasta file. Plasmid backbone sequences are provided in the plasmid_sequences.genbank file. Primer sequences used for cloning will be provided upon request. For all primers, 18 to 22bp homologies without any special primer design criteria were sufficient for successful PCR amplification with Phusion High-Fidelity DNA polymerase (NEB). Initial yfp construct. All yfp variants used in this study were modified starting from a single yellow fluorescent protein gene sequence (called yfp0 in the sequence file and plasmid map). This yfp0 sequence encoded the fast-maturing 'Venus' variant of YFP [14]. All 238 codons of yfp0 were chosen such that they were decoded by abundant tRNA isoacceptors for each amino acid. Such a choice of codons ensured that the native level of demand for each tRNA isoacceptor inside the cell was minimally perturbed by the low-copy expression of fluorescent reporter genes. The yfp0 sequence was built de novo (synthesis by Genscript, USA). The synthesized yfp0 sequence was cloned between the KpnI and HindIII restriction sites of the pZS^*11 plasmid vector using standard molecularbiology techniques (19). The plasmid map of the resulting construct, pZS^*11 -yfp0 is shown in Fig. S17.

Synonymous variants of yfp. A subset of codons in $yfp\theta$ corresponding to 7 amino acids (Leu, Arg, Ser, Pro, Ile, Gln, Phe) were mutated to create the initial 29 synonymous variants of yfp (yfp1 - yfp29 in the gene_sequences.fasta file, sequences in the same order as shown in Fig. 1A). The 4 yfp variants corresponding to Pro (yfp19-yfp22) had all the Pro codons mutated to the most frequent CCG codon since the original yfp0 sequence had a few CCA and CCT codons that are more sensitive to Pro limitation. Similarly, all the Phe codons in yfp0 were mutated to the most abundant Phe codon TTT for the two Phe variants of yfp0 (yfp28-yfp29). Both these groups of variants (6 total) had higher overall fluorescence during amino-acid rich conditions than the rest of the 23 variants. This higher fluorescence is likely due to changes in secondary structure near the ribosome binding region on the mRNA as a consequence of mutations near ATG. However, this change is common across all variants within the Pro and Phe synonymous codon groups and hence is not responsible for the differential response to cognate amino acid limitation measured within these synonymous codon groups.

For constructing the 29 yfp variants, yfp0 from pZS*11yfp0 was first cloned into a *pUC19* cloning vector between the KpnI and HindIII restriction sites. A commercial site-directed mutagenesis kit (Quickchange Lightening Multi, Applied Biosystems) was used to introduce the mutations corresponding to each of the 29 variants and the manufacturer's protocol was followed. The resulting variants were verified by Sanger sequencing and then cloned into the pZS^*11 expression vector backbone between the KpnI and HindIII sites. The 22 single CTA variants of yfp (Fig. S2) were constructed using the same procedure as above. The 29 yfp variants for Western blotting (Fig. S1) were created using the same procedure as above, but with the addition of a 22 codon sequence at the 5' end that encoded a 3X-FLAG peptide recognized by a commercially available, anti-FLAG, antibody (Sigma). The 22-codon sequence is: GACTACAAAGACCATGACGGTGATTATAAA-GATCATGACATCGACTACAAGGATGACGATGACAAG.

tRNA expression vectors. The 5 distinct Leu tRNA isoacceptors encoded by the genes leuQ, leuU, leuW, leuX and leuZ, and the 4 distinct Arg tRNA isoacceptors encoded by the genes argV, argX, argU and argW were cloned between the EcoRI and HindIII sites of the pZA32 expression vector (Fig. S18). These genes were amplified by PCR from the chromosome of *E. coli* strain MG1655. In addition to these native tRNA genes, a synthetic tRNA gene $arg2_m$ cognate to the CGA Arg codon was also created. Normally, the ^{ACG}Arg2 tRNA with ICG anti-codon reads the CGA codon inefficiently through a purine-purine wobble pairing. Expressing a synthetic tRNA with an anticodon UCG restores efficient reading of this codon and is equivalent to increasing the supply

of the corresponding cognate tRNA isoacceptor. This synthetic tRNA isoacceptor was created from the pZA32-argV expression vector using overlap PCR to introduce the necessary single bp mutation in the anticodon of argV. The pZA32 vectors with the tRNA genes were electroporated into strains already containing the YFP expression vectors.

Library of *E. coli* ORF-*yfp* fusions. 92 *E. coli* Open Reading Frames (ORFs) were selected for experimental validation of the Leu Codon Robustness Index (Leu CRI). These ORFs were chosen to span a wide range of predicted Leu CRI values and functional categories (Fig. S7 and Table S1). First, a modified pZS^*11 -yfp0 vector backbone was created in which the start codon of yfp0 was replaced by a GGSGGS hexapeptide linker sequence: GGTGGATCCGGCGGTTCT containing a BamHI restriction site. Next, the 92 ORFs (without the stop codon) were amplified by PCR from the chromosome of E. coli strain MG1655 with 5'-KpnI and 3'-BamHI restriction site overhangs. These PCR fragments were cloned into the modified pZS^*11 -yfp0 vector backbone containing the BamHI restriction site. 13 of the 92 ORFs had either an internal KpnI or an internal BamHI site. In these cases, a larger fragment that included adjoining sections of the $pZS^*11-ufp0$ vector was constructed by overlap PCR and then cloned using other restriction sites (EcoRI or HindIII). Thus the final constructs had one of the 92 E. coli ORFs connected through a hexapeptide linker with yfp0. All the cloned sequences were verified by PCR for inserts of right length and around 40 ORF constructs were verified by Sanger sequencing. Two biological replicates of each ORF construct were compared for their synthesis robustness values as measured during the amino acid limitation assay and these values showed a high degree of correlation (Pearson $\rho = 0.93$, Fig. S19).

For validating the Arg codon robustness index (Arg CRI), 56 *E. coli* ORFs that included a subset of the above 92 ORFs were chosen (Table S2). The cloning procedure was exactly analogous to the above 92 ORFs but with one difference: the yfp0 part of the fusion construct was replaced by a synonymous variant of yfp0 (yfp7) that had the Arg codon AGA instead of the CGT and CGC codons in the yfp0 sequence. The codon AGA has the highest w_i value among the Arg codons (see Fig. 3B) and hence has a minimal effect on the measured robustness of the ORF fusions during Arg limitation.

Co-expression of ^{GAG}Leu2-tRNA with *E. coli* ORF-*yfp* **fusions.** Out of the 92 *E. coli* ORF-*yfp* fusions, 21 were chosen for co-expression with the ^{GAG}Leu2 tRNA that is cognate to the codons CTC and CTT. The 21 ORFs were chosen such that 11 of them had a lower Leu CRI prediction than their wild-type counterparts while the other 10 ORFs had a higher Leu CRI prediction than their wild-type counterparts (Table S3). This choice also corresponded respectively to either high frequency of the non-cognate TTA and TTG codons for ^{GAG}Leu2 or high frequency of the cognate codons CTC and CTT. The strains containing the 21 ORF fusions were each made electro-competent and then transformed with the *pZA32-leuU* plasmid that expresses ^{GAG}Leu2.

Synonymous variants of $E.\ coli\ ORF-yfp$ fusions. Out of the 92 $E.\ coli\ ORF-yfp$ fusions, 13 were selected for creating synonymous mutants (Table S2). These 13 ORFs had a high frequency of one or both of the Leu codons, TTA or TTG and these codons were mutated to the Leu codon, CTC. All these 3 codons, TTA, TTG and CTC occur at similar frequencies on average across the genome of $E.\ coli$. The 13 ORF-yfp fusions were amplified by PCR from the pZS^{*11} vectors between the EcoRI and XbaI restriction sites (see Fig. S17). These fragments were cloned between EcoRI and XbaI sites of the pUC19 cloning vector. A commercial site-directed mutagenesis kit (Quickchange Lightening Multi, Applied Biosystems) was used to introduce the TTA, TTG \rightarrow CTC mutations. A unique primer was designed for each of the TTG or TTA codons in the 13 ORFs, and these primers encoded the CTC mutation. All the primers corresponding to each ORF were mixed and then used in the mutagenesis reaction. This procedure resulted in mutant coding sequences with TTA, TTG \rightarrow CTC mutations at random locations. 10 colonies for each ORF were sequenced and each unique mutant sequence was then cloned into the pZS^*11 expression vector. At the end of the procedure, a total of 63 constructs were created that each had between one and seven TTA, TTG \rightarrow CTC mutations (see gene_sequences.fasta file for exact sequences).

Total RNA extraction

Total RNA was extracted for two different experiments (Figs. 4B, S14). Phenol-chloroform extraction method was used to obtain total RNA. Briefly, 3ml of cells were quickly mixed with 5ml of ice-cold water and harvested by centrifugation at 3000g for 10min. Cell pellets were re-suspended in 500μ l of 0.3M sodium acetate-10mM EDTA, pH 4.8 buffer. The resuspended cells were mixed with 500μ l of acetate-saturated phenol-chloroform at pH 4.8, 50μ l of 20% SDS and 500μ l of acid-washed glass beads (G1277, Sigma). The mixture was shaken in a vortexer for 20 min at 4C. The aqueous layer was extracted twice with acetate-saturated phenol-chloroform at pH 4.8 and once with chloroform. Total RNA was precipitated with an equal volume of isopropanol and washed with 70% ethanol-50mM sodium acetate pH 4.8 and finally re-suspended in 200μ l of RNase-free water. 20μ l of the total RNA was treated with DNase (EN0521, Fermentas) to remove residual DNA contamination (manufacturer's instructions were followed). The DNA-free RNA was re-suspended in 200μ l of RNase-free water. Intact RNA was confirmed by observation of sharp rRNA bands in native agarose gel electrophoresis.

RT-qPCR

Reverse transcription (RT) was performed using 4μ l of the DNA-free RNA (100-200ng) and Maxima reverse transcription kit (K1641, Fermentas), used according to the manufacturer's instructions. Random hexamer primers were used for priming the RT reaction. At the end of the RT reaction, the 20μ l reaction was diluted 100-fold and 10μ l of this diluted sample was used for qPCR in the next step. qPCR was performed using Maxima SYBR-Green qPCR kit (K0221, Fermentas) and manufacturer's instructions were followed. qPCR was performed in triplicates for each RT reaction and appropriate negative RT controls were used to confirm the absence of DNA contamination. gapA mRNA was used as internal reference to normalize all other mRNA levels. Standard curves with 6 serial dilutions were used to optimize reaction conditions and ensure amplification efficiency of between 90-100% for the *yfp* and *gapA* amplicons. $\Delta\Delta C_t$ method was used to obtain the change in mRNA levels due to amino acid limitation. The qPCR primer sequences are given in Table S8.

Western blotting

Fresh colonies were used to inoculate overnight cultures. These overnight cultures were then diluted 1:100 into 1ml of rich-defined medium with all 20 amino acids (see section on growth and fluorescence measurements for media composition). After approximately 3.5 hours of growth at 30° C when OD_{600} was ~0.4, cells were spun down at 9000g for 1 min, and then re-suspended in 1ml of rich-defined medium without the amino acid whose limitation was to be induced. This re-suspended culture was then split into two equal aliquots. The limiting amino acid was added to one aliquot (as a richmedium control) while the other aliquot did not have the limiting amino acid. The re-suspended medium also contained 200ng/ml of anhydro-tetracycline in order to induce the p_L tetO1 promoter that controls the 3XFLAG-yfp variants. After growth at 30° C for 60 min, cells were spun down at 12000g, 1 min and re-suspended in $40-400\mu$ l of CellLytic B buffer (Sigma, B7435). The buffer volume used was proportional to the OD_{600} measured at the time of harvesting the culture. The lysate was stored at -80° C. 10μ l of the lysate was mixed with 2X Laemmli Buffer (Biorad) and then loaded onto each lane of a pre-cast polyacrylamide gel (Biorad) and SDS-PAGE was carried out at 100V for 120 min. Proteins were transferred to a nitrocellulose membrane by semi-dry blotting at 180mA for 60 min. The membrane was blocked in 2% skim-milk-TBST overnight, and then incubated with a 1:2000 dilution of an anti-FLAG antibody (F3165, Sigma) in 10ml of 2% skim-milk-TBST with shaking at room temperature for 90 min. After washing 4 times for 5 min with TBST, the membrane was incubated with 1:2000 dilution of a secondary HRP-conjugated antibody (7076, Cell Signaling) in 15ml of 2% skim-milk-TBST with shaking at room temperature for 60 min. After washing 4 times for 5 min with TBST, the membrane was treated with an HRP substrate (L00354, Genscript) for 5 min and exposed for 30s to a luminescence imager.

Analyses

Matlab R2009b (Mathworks) was used for all analyses unless otherwise mentioned. All correlations and P-values reported in this work were calculated using the Matlab command 'corr' with the 'Type' option set to either 'Spearman' or 'Pearson' as appropriate.

Growth and fluorescence analysis. Background absorbance and fluorescence values (obtained from wells containing only growth media) were subtracted from the measured time series for each well. An exponential curve was fitted to the amino acid-rich growth regime for all data points located at least 50 min before the onset of amino acid limitation. A straight line was fitted to the amino acid-limited growth regime for all data points located at least 50 min after the onset time. These fits were performed using the Matlab command 'fit', and the in-built library options 'Exp1' and 'Poly1' respectively. To automatically identify the onset time, the intersection point between the two fitted curves was designated as the onset time of amino acid limitation. This inferred onset time coincided with the onset time identified through visual inspection of the growth curves.

To minimize noise in calculated protein synthesis rates, an exponential curve was fitted to the amino acid-rich regime of the fluorescence time-series and a straight line was fitted to the amino acid-limited regime of the fluorescence time-series. These fits were performed using the Matlab command *fit*, and the in-built library options 'Exp1' and 'Poly1' respectively. Protein synthesis rate, S was calculated as

Protein synthesis rate
$$S = \frac{1}{\text{Absorbance}} \times \frac{d(\text{Fluorescence})}{d(\text{time})}$$
[1]

First, the above formula was evaluated at the onset time of amino acid limitation using the exponential fits for absorbance and fluorescence data in the amino acid rich growth regime. Next, the same formula was evaluated at the onset time using the linear fits for absorbance and fluorescence data in the amino acid limited growth regime. These two values correspond to the protein synthesis rates reported for the amino acid rich and amino acid limited growth regimes (such as the data in Fig. 1D). The protein synthesis rates were normalized within each synonymous codon family and for each growth condition. Robustness of protein synthesis to amino acid limitation was calculated as the ratio of normalized protein synthesis rates between the amino acid rich and amino acid limited growth regimes.

In the case of the experiment without methyl ester (Fig. S15 and Fig. S16), the onset time of amino acid limited growth was determined exactly as above. Then starvation robustness was calculated as the normalized ratio of total fluorescence increase after the onset of amino acid limited growth to the fluorescence increase before this onset. Total fluorescence increase rather than protein synthesis rate was used for this analysis since protein synthesis rates decreased continuously to zero after the onset of amino acid limited growth in the absence of methyl ester analogs.

Calculation of CRI. CRI for a protein coding sequence corresponding to a limiting amino acid was calculated by multiplying the w_i values for codons cognate to the limiting amino acid in that sequence. w_i values in Fig. 3B were calculated using the robustness of protein synthesis of the corresponding yfp variants during cognate amino acid limitation (Fig. 1D). Based on the non-cognate amino acid limitation experiment (Fig. S2), the w_i values for all codons other than those cognate to the limiting amino acid are set to be equal to 1. For illustration, we demonstrate the calculation of w_i for the six Leu codons below. The exact same procedure was followed for other synonymous codon families. Taking $loq_2w_i \equiv W_i$ for each codon, and log_2 (robustness during amino acid limited growth) $\equiv SR$ for each yfp variant,

$$7 \times W_{CTA} + 15 \times W_{CTG} = SR_{yfp,CTA}$$
 [2]

$$7 \times W_{CTC} + 15 \times W_{CTG} = SR_{ufp,CTC}$$
^[3]

$$22 \times W_{CTG} = SR_{yfp,CTG} \qquad [4]$$

$$7 \times W_{CTT} + 15 \times W_{CTG} = SR_{yfp,CTT}$$
^[5]

$$7 \times W_{TTA} + 15 \times W_{CTG} = SR_{ufp,TTA}$$
 [6]

 $7 \times W_{TTG} + 15 \times W_{CTG} = SR_{yfp,TTA}$ $7 \times W_{TTG} + 15 \times W_{CTG} = SR_{yfp,TTG}$ [7]

The multiplicative factors on the LHS in front of W_i correspond to the frequency of the Leu codon i in the corresponding Leu variant of yfp (see Fig. 1A). The RHS is the measured (log2) robustness of protein synthesis from the corresponding yfp variant during Leu limitation (see Fig. 1D). These equations were solved simultaneously to determine the w_i value for each Leu codon. Revised w_i values (Table S7) based on yfp measurements in the presence of ^{GAG}Leu2 tRNA (Fig. 2) were used for calculation of Leu CRI in the case of ^{GAG}Leu2 tRNA co-expression with *E. coli* ORFs (Fig. 3D).

Leu and Arg CRI for *E. coli* ORFs. 4300 *E. coli* ORF sequences were parsed out from the MG1655 genome sequence (NCBI website, Accession number: NC_000913, downloaded on 14th Apr 2011). For each of these 4300 E. coli ORFs, Leu or Arg CRI was calculated by multiplying the w_i values for either all Leu or all Arg codons respectively in the ORF sequence. For the 63 synonymous variants of 13 ORFs (Fig.

S10), Leu CRI values were calculated using the same procedure as above after accounting for the synonymous mutations. For the 21 ORFs co-expressed with Leu2 tRNA (Fig. 3D), revised w_i values were first calculated using the method outlined in the previous section (Table S7), and using measurements on the 6 Leu variants of *yfp* complemented with ^{GAG}Leu2 tRNA (3rd column in Fig. 2A). These revised w_i values were then used to calculate Leu CRI under tRNA co-expression for the 21 tRNA co-expressed ORFs applying the same procedure as for the non co-expressed case.

Z-score for CRI. To quantify the deviation in CRI from its expected value for each of the 4300 ORFs in the *E. coli* genome, 1000 random coding sequences were generated for each ORF. Each random version preserved the original amino acid sequence, but the codons for a single amino acid were sampled randomly from a multinomial distribution based on the average frequency of codons for that amino acid in the genome. CRI values were calculated for each random version of the gene, and a distribution of CRI values was generated from the 1000 random trials. The average, μ_{CRI} and standard deviation, σ_{CRI} of this CRI distribution was used to calculate the Z-score for CRI as follows:

$$Z_{CRI} = \frac{CRI_{observed} - \mu_{CRI}}{\sigma_{CRI}}$$
[8]

In the case of the Z-score for leucine shown in Fig. 4B, the leucine codon CTG was not randomized in the above calculation and only the remaining 5 leucine codons: CTA, CTC, CTT, TTA, and TTG were randomized. This step is important since CTG, which is read by an abundant tRNA isoacceptor, is enriched in highly-expressed genes, and such genes will show up falsely as perturbation-robust genes because CTG is also the codon that is most robust to leucine limitation in our experiments (see Fig. 1D).

Codon-specific bioinformatic measures. Codon usage in Fig. S4 was calculated as the average frequency of each codon across the genome of *E. coli* MG1655 (4300 ORFs). tRNA concentrations in Fig. S4 were taken from previous work (see Table 2 in [15]). Concentrations of all cognate tRNAs for each codon were summed together. The codon-tRNA adaptation index in Fig. S4 is taken from literature (see Table S2 in [16]). The tAI value for the CGA codon was revised from the unrealistically low value of 0.00005 to 0.1333 as explained previously [17]. For inferring codon elongation rates from charged tRNA fractions (Fig. S5), we used the formula for codon elongation rate from [18]:

$$\frac{1}{v_k} = \tau_0 + \frac{1}{\sum_i t_i \alpha_i k_{ik}},$$
[9]

where v_k is the elongation rate of codon k, τ_0 is the codonindependent elongation time across any codon, t_i is the concentration of tRNA isoacceptor i that is cognate to codon k, k_{ik} is the second-order rate constant for binding of the ternary complex containing the charged isoacceptor i to the ribosome at codon k, and α_i is the charged fraction of isoacceptor i. We calculated the codon elongation rates during amino acid limitation using the measured charged fractions from [6]. For amino acid rich conditions, we set the charged fraction to be equal to unity. We used $\tau_0 = 0.05s^{-1}$, and $k_{ik} = 2 \times 10^7 M^{-1} s^{-1}$ similar to [18]. The ratio of codon elongation rates was then normalized within each codon family by the maximum value within that family. **ORF-specific bioinformatic measures.** Codon Adaptation Index was calculated for each *E. coli* ORF using the method in [19]. This calculation was implemented using the Codon-AdaptationIndex class in the CodonUsage module of BioPython (version 1.58). tRNA Adaptation Index was calculated for each *E. coli* ORF using the method in [20]. This calculation was implemented using the codonR package (http://people.cryst.bbk.ac.uk/~fdosr01/tAI/index.html, downloaded on 3rd Sep 2011). mRNA folding energy was calculated for the first 37nt of each *E. coli* ORF together with the 5 upstream nucleotides (GTACC) in the *pZS11* plasmid backbone. Calculation was implemented using the *hybrid-ss-min* command in UNAFold software v3.8 [21] with default parameter values for reaction conditions (NA = RNA, T = 37, [Na+] = 1, [Mg++] = 0, maxloop = 30).

Supplementary notes

Use of methyl esters in amino acid limitation experiments. Before we settled on the methyl ester analog-based experiments, we tested two other amino acid limitation assays that are commonly used in the literature. The first assay is a spin \rightarrow wash \rightarrow resuspend in amino acid+ / amino acid- medium [6]. We did not pursue this assay for most experiments since it is logistically difficult to perform this assay when working simultaneously with more than a dozen strains. However, we used this assay for the Western blotting and RT-qPCR measurements on a few strains (Figs. S1, S14 and 4B).

The second assay involves starting with a low initial concentration of an amino acid and letting the bacterial cultures exhaust the amino acid in the medium through exponential growth [22]. The bacteria then enter the amino acid limited regime in mid-log phase without any intervention from the experimenter. However in the absence of exogenous sources of amino acid in the amino acid limited regime, protein synthesis occurs only transiently for less than an hour under these conditions and YFP synthesis rates from all *ufp* variants drop below measurable levels at the end of this time period (Fig. S15). More importantly, there is no extended steady state during which differential protein synthesis rates can be measured accurately. Nevertheless, we have confirmed that the measurements with and without methyl esters give qualitatively similar results (Fig. S16). In addition, Western blotting done in the absence of methyl ester reproduced the heirarchy in protein levels between synonymous variants of YFP during amino acid limitation (Fig. S1).

In contrast to the assay without methyl ester, presence of methyl ester analogs in the growth medium results in a quasisteady state of amino acid limited growth due to hydrolysis of the ester, during which differential YFP expression can be measured easily (Fig. S15). Such partial amino acid limited growth is also likely to be the relevant scenario when prototrophic strains run out of amino acids in their growth media and have a limited supply of amino acids through protein degradation or partially up-regulated biosynthesis pathways.

Effect of varying the initial concentrations of amino acids and methyl esters. Increasing the initial concentration of the amino acid or its methyl ester results in a higher cell density for the onset of amino acid limitation, and when the corresponding concentrations are decreased, this onset happens at a lower cell density. Importantly, the observed differential robustness of protein synthesis (such as the data shown in Fig. 1D) is qualitatively the same upon 2-fold changes to the initial concentration of either the amino acid or its methyl ester. As an extreme example, see Figs. S15 and S16 for comparison between the cases with and without methyl ester analog in the growth medium.

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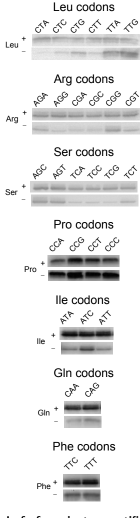
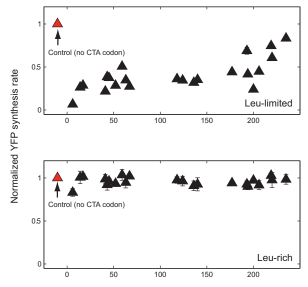


Fig. S1: Expression level of yfp variants quantified through Western blotting

Modified versions of 29 yfp variants (Fig. 1A) were created that had a 3X-FLAG tag at the 5' end. These yfp variants were transformed into the respective *E. coli* auxotrophs in which YFP synthesis was repressed by the TetR protein. Cells were harvested at an OD_{600} of 0.4 and re-suspended in medium with or without the corresponding amino acid. Expression of YFP was induced using 200ng/ml anhydrotetracyline, and cells were harvested after 60 min. For each set of yfp variants under a specific growth condition, the same amount of total protein (as measured by OD_{600} before cell lysis) was used for Western blotting.



Location of CTG to CTA synonymous mutation along yfp (codon)

Fig. S2: Effect of a single synonymous mutation on YFP synthesis rate

22 variants of yfp were synthesized, each of which had a single CTA codon at one of the 22 leucine codon locations along yfp. The remaining leucine codons in each variant were the perturbation-robust CTG codon. The 'control' yfp variant did not have any CTA codon. Vertical axis refers to the YFP synthesis rate from the 22 variants normalized by that of the control variant, either during leucine limitation (top panel) or during leucine-rich growth (bottom panel). Horizontal axis indicates the location of the CTA codon along each yfp variant (ATG start codon = 1). Error bars show standard error over three replicate cultures.

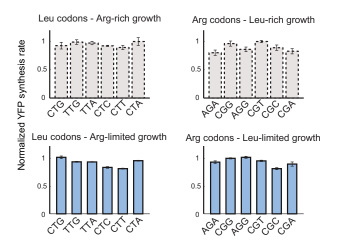


Fig. S3: YFP synthesis rate during limitation for a non-cognate amino acid

Leucine and arginine variants of yfp were expressed in an *E. coli* strain, CP78, that is auxotrophic for both leucine and arginine. Response of the 6 Leu variants to Arg limitation is determined by the Arg codons in yfp (CGT and CGC) that are common across all 6 Leu variants. Reciprocally, the response of the 6 Arg variants to Leu limitation is determined by the Leu codon that is common to the Arg variants of yfp (CTG). YFP synthesis rates are defined as in Fig. 1D. Error bars show standard error over three replicate cultures.

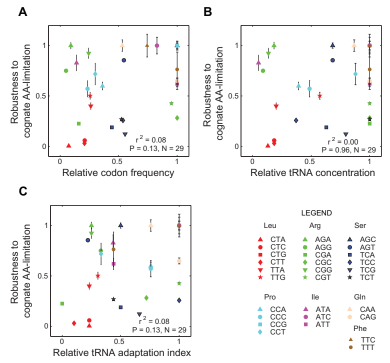


Fig. S4: Comparison of synthesis rate robustness with codon usage and tRNA concentration

(A) Codon usage was calculated as the average frequency of each codon across all protein coding sequences in *E. coli.* (*B*) tRNA concentration for each codon was calculated as the sum of tRNA concentrations for all cognate tRNAs [15]. (*C*) Since tRNAs can differ substantially in their affinity for their cognate codons, we also compared the measured robustness against the tRNA adaptation index for each codon [20]. This index accounts for different affinities of synonymous codons for the same tRNA isoacceptor. All three measures along the horizontal axes were normalized by the maximum value within each codon family. Robustness to amino acid limitation was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases. Error bars represent standard error over three replicate cultures. The data points that are not visible for a few codons overlap at the top right-hand corner of each plot.

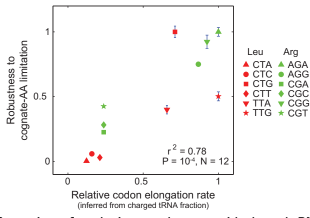


Fig. S5: Comparison of synthesis rate robustness with charged tRNA fraction

To compare YFP synthesis rates with charged tRNA fractions, the elongation rates for leucine and arginine codons were inferred from the measured charged fraction of leucine and arginine tRNA isoacceptors [6] (see section on codon specific bioinformatic measures). Previously assigned codon-tRNA assignments and kinetic parameters were used [18]. Note that charged tRNA fractions cannot be directly compared with synthesis rates of yfp variants due to overlapping and multiple codon assignments for several tRNA isoacceptors. Robustness to amino acid limitation was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases. Error bars represent standard error over three replicate cultures. Relative codon elongation rate is the ratio of codon elongation rates between amino acid starved and amino acid rich growth regimes, normalized to the maximum value within each synonymous codon family.

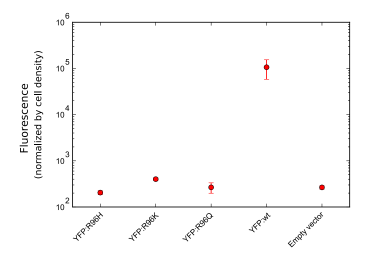


Fig. S6: Miscoding of a single arginine residue in YFP causes loss of fluorescence

To test whether mistranslation of arginine residues can underlie the high residual fluorescence of Arg yfp variant-Arg tRNA pairs (AGA: arg3, AGG: arg3, and CGG:arg4, arg5 in Fig. 2B), three YFP mutants were created that had one of three single point mutations at Arg96: R96H, R96K, and R96Q. The mutant and the 'wild-type' YFP proteins were expressed from a pUC18 high-copy vector. Each of the three mutations at Arg96 to a chemically similar amino acid (H, K or Q) decreased YFP fluorescence to background level (that of an empty pUC18 vector). Error bars denote standard deviation over five biological replicates.

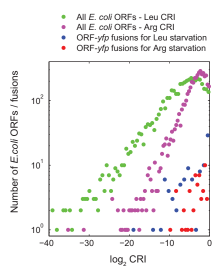


Fig. S7: Histogram of CRI

Green and purple data markers correspond to the Leu and Arg CRI values for 4300 ORFs in *E. coli*'s genome. Blue and red data markers correspond respectively to Leu and Arg CRI values for the *E. coli* ORF-yfp fusions that were used to experimentally validate CRI.

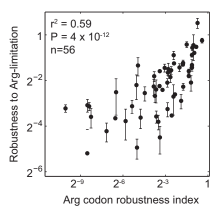


Fig. S8: Correlation of Arg CRI with measured robustness of 56 E. coli ORF-yfp fusions

The yfp sequence used for this experiment had the AGA codon at all Arg codon locations of yfp since AGA has the highest w_i value among arginine codons (see Fig. 3B). Correlation is reported as squared Spearman rank correlation. Error bars show standard error over three replicate cultures. Robustness to amino acid limitation was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases.

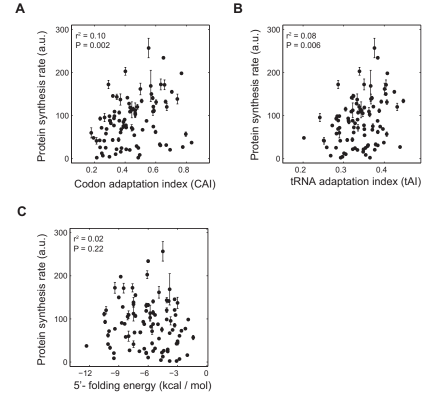
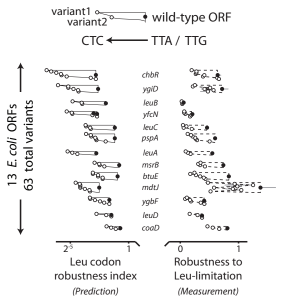


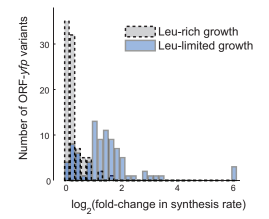
Fig. S9: Correlation of protein synthesis rate during amino acid rich growth with measures of translation efficiency

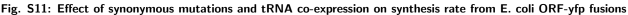
Protein synthesis rates from 92 *E. coli* ORF-*yfp* fusions during Leu-rich growth showed only a weak correlation with measures of codon adaptation, tRNA adaptation and 5' folding energy of mRNA. Folding energy was calculated from -5 to +37 nt of the ATG codon. Codon adaptation index (CAI) and tRNA adaptation index (tAI) were calculated using Biopython and codonR packages. Correlations are reported as squared Spearman rank-correlation coefficient.





Sixty three synonymous variants of 13 ORF-yfp fusions were constructed by mutating wild-type TTG or TTA codons in the ORF sequence to the codon CTC that causes sensitive protein synthesis rate under leucine limitation. The number of mutations was between 1 and 6 and the location of these mutations was random. 59 of the 63 variants displayed a decrease in their robustness during Leu limitation (dashed lines) that was predicted by Leu CRI (solid lines). In addition, magnitude of the changes in robustness during Leu limitation were positively correlated with magnitude of the changes in Leu CRI (r^2 = 0.19, P = 10⁻⁴). Filled circles indicate values for ORF_{wild-type} and open circles indicate values for ORF_{variant}. Different open circles within a single polygon correspond to distinct ORF variants for the same wild-type ORF. Robustness to amino acid limitation was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases. Error bars show standard error over three replicate cultures. Most error bars are smaller than data markers. DNA sequences for variants are provided in gene_sequences.fasta supplementary file.





We analyzed the change in protein synthesis rates from the 21 ORF-yfp fusions co-expressed with ^{GAG}Leu2 (Fig. 3D) and the 63 different ORF-yfp variants with synonymous mutations (Fig. S10). Several of the ^{GAG}Leu2-coexpressed as well as the synonymously-mutated ORF-yfp variants (84 total variants) had significantly altered protein synthesis rates compared to their non-tRNA co-expressed or non-mutated counterparts (referred as wild type) during leucine limited growth (green histogram, median fold-change in protein synthesis rates = 2.37). By comparison, most of the 84 variants had similar protein synthesis rates to their wild-type counterparts during leucine rich growth (grey histogram, median fold-change in protein synthesis rates = 1.12). Protein synthesis rates were defined as in Fig. 1D.

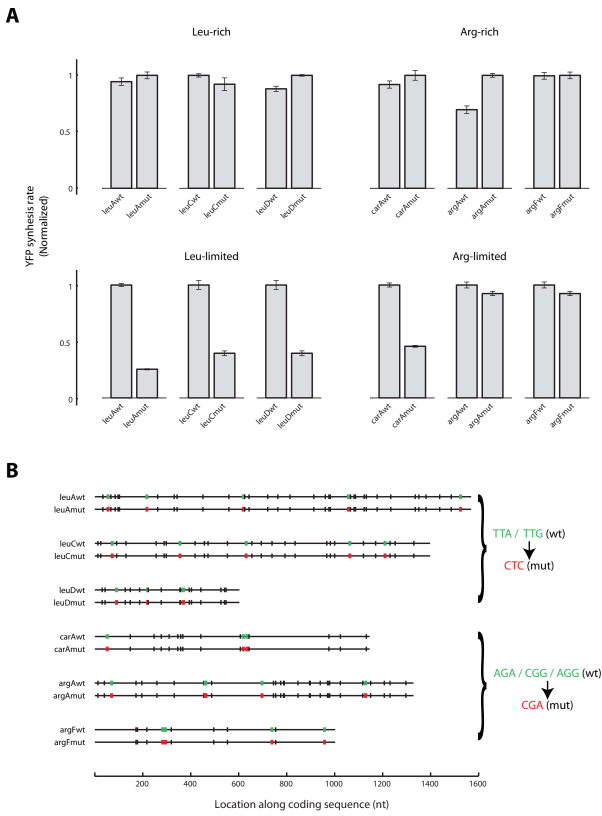


Fig. S12: Effect of synonymous mutations on synthesis rate from amino acid biosynthesis genes

(A) Synthesis rates from leuA, leuC, leuD and carA, argA, argF -yfp fusions encoding either wild-type or mutant ORF sequences during amino acid rich and amino acid limited growth. The synthesis rates were normalized for each pair of wild-type and mutant ORF-yfp fusions, and also for each growth condition. Error bars show standard error over six replicate cultures. (B) Position and identity of synonymous mutations in wild-type and mutant sequences used for the experiment in (A). The black vertical bars correspond to the non-mutated leucine codons in the case of leuA, leuC and leuD, and to the non-mutated arginine codons in the case of carA, argA and argF.

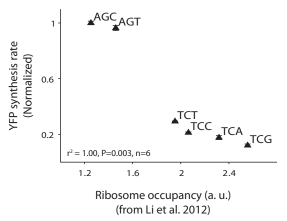


Fig. S13: Correlation of protein synthesis rate with ribosome occupancy at serine codons during serine-limited growth

Protein synthesis rate of serine synonymous variants of yfp during serine limitation (same data as in Fig. 1D, third panel) is negatively correlated with genome-wide ribosome occupancy at serine codons during serine-limited growth of *E. coli*. The increased occupancy at perturbation-sensitive serine codons is consistent with selective ribosome pausing at these codons. Ribosome occupancy data was taken from a recent ribosome profiling experiment in *E. coli* [23].

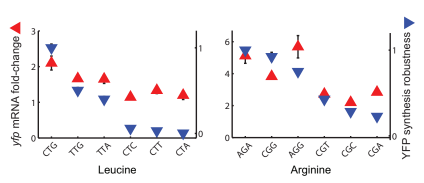


Fig. S14: Change in mRNA level of yfp variants in response to cognate amino acid limitation

We measured the change in mRNA levels of different yfp variants in response to amino acid limitation. Total RNA was extracted either during exponential amino acid rich growth or 60 min after amino acid limited growth in the presence of the amino acid methyl ester. mRNA levels were quantified by RT-qPCR relative to gapA mRNA. Error bars show standard error of triplicate qPCR measurements. Synthesis rate robustness to amino acid limitation was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases.

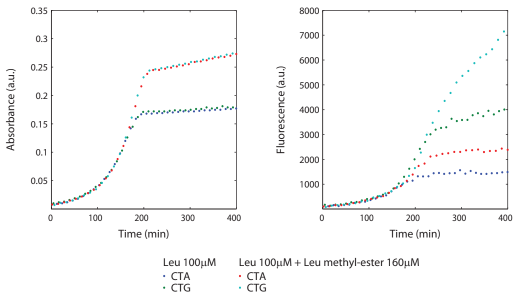
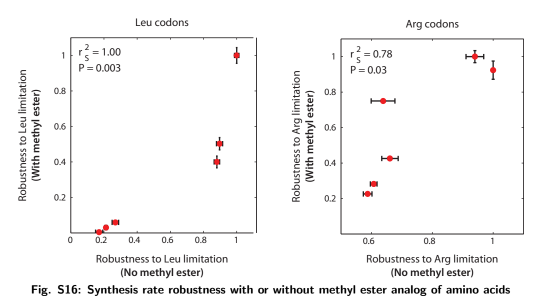


Fig. S15: Raw absorbance and fluorescence curves with or without methyl ester analog of amino acids

Growth and fluorescence curves for two yfp variants corresponding to CTA and CTG codons are shown here as representative examples for amino acid limited growth in the presence or absence of methyl ester analogs in the growth medium. Absorbance as measured using spectrometry is proportional to cell density. Presence of methyl ester analogs caused an increase in the time and cell density at which amino acid limited growth began. More importantly, inefficient metabolism of methyl ester analogs resulted in a slow but steady growth in amino acid limited regime. This residual growth ensured that YFP synthesis continued robustly from the CTG yfp variant under these conditions. By contrast, in the absence of methyl esters in the growth medium, YFP synthesis from all yfp variants eventually dropped to zero.



Robustness to amino acid limitation in the absence of methyl ester analogs was calculated as the ratio of fluorescence change between the amino acid limited growth phase and amino acid rich growth phase. This ratio was further normalized by the maximum value within each codon family. Robustness to amino acid limitation in the presence of methyl ester analogs was quantified as the ratio of normalized YFP synthesis rates between amino acid limited and amino acid rich growth phases. Error bars show standard error over three replicate cultures.

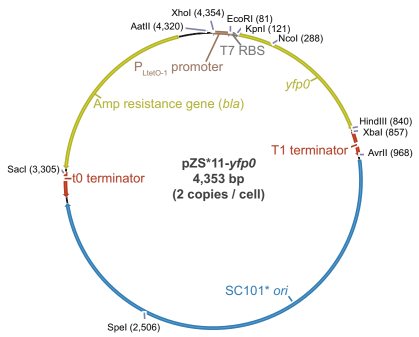


Fig. S17: Plasmid map of expression vector for yfp and E. coli ORF-yfp fusions

A specific plasmid construct with $yfp\theta$ is shown here. In the case of ORF-yfp fusions, yfp was fused in-frame to the 3'-end of the ORF with a GGSGGS hexa-peptide linker sequence that encoded a BamHI restriction site and the resulting coding sequence of the fusion protein was cloned between the KpnI and HindIII restriction sites in the above vector.

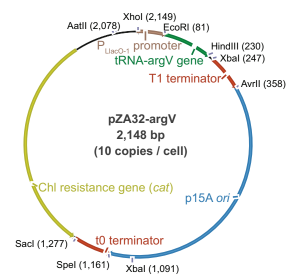


Fig. S18: Plasmid map of expression vector for tRNA genes

A specific construct encoding an Arg tRNA is shown here.

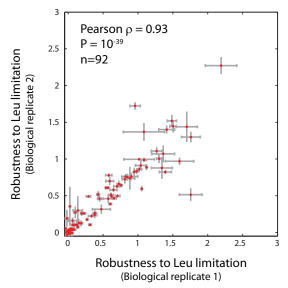


Fig. S19: Reproducibility of measurements between biological replicates of 92 E. coli ORF-yfp fusions

Two different colonies were picked after cloning the 92 *E. coli* ORF-yfp fusions and the same leucine limitation assay that was used for the data in Fig. 3C was performed on these two biological replicates on two different days. None of the clones for replicate 2 were sequence-verified and hence the few outliers seen above could be the result of errors in the cloned sequences. The data reported in Fig. 3C is from replicate 1 for which about 40 constructs were sequence-verified. Robustness to Leu limitation was calculated as the ratio of normalized YFP synthesis rates between Leu limited and Leu rich growth phases.

Table S1: 92 E. coli ORF-yfp fusions used for Leu CRI validation

Genes are arranged by increasing values of Leu CRI. $S_{Leu-rich}$ and $S_{Leu-limited}$ refer to respective protein synthesis rates (a.u. per sec per cell). Robustness refers to the ratio between the two protein synthesis rates after normalization by the corresponding value for the CTG variant of yfp (which is the yfp tag in these ORF-yfp fusions). \pm refers to standard error of measurement.

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$	Gene product
1	polB	10.4 ± 0.3	0.6 ± 0.6	0.063 ± 0.062	-19.45	DNA polymerase II
2	$_{\rm thiH}$	69.7 ± 1.1	-0.7 ± 0.8	-0.012 ± 0.013	-14.52	tyrosine lyase, involved in
						thiamin-thiazolemoiety synthe-
						sis
3	aat	26.7 ± 0.8	1.3 ± 0.3	0.054 ± 0.015	-12.52	leucyl/phenylalanyl-tRNA-
						protein transferase
4	gdhA	20.4 ± 0.3	-0.2 ± 0.2	-0.013 ± 0.009	-11.62	glutamate dehydrogenase,
						NADP-specific
5	serC	91.3 ± 0.5	-1.7 ± 0.6	-0.020 ± 0.008	-11.6	3-
						phosphoserine/phosphohydroxy
						threenine aminotransferase
6	gpsA	111.8 ± 3.3	1.9 ± 1.9	0.020 ± 0.018	-11.38	glycerol-3-phosphate dehydro-
						genase $(NAD+)$
7	mlrA	44.1 ± 1.5	-0.7 ± 0.8	-0.017 ± 0.019	-11.09	DNA-binding transcriptional
						regulator
8	ybeU	41.5 ± 7.6	0.5 ± 0.3	0.017 ± 0.013	-11	conserved protein, DUF1266
						family
9	argS	67.6 ± 1.9	1.0 ± 0.6	0.017 ± 0.010	-10.93	arginyl-tRNA synthetase
10	ilvD	59.2 ± 5.6	3.4 ± 0.9	0.064 ± 0.017	-10.39	dihydroxyacid dehydratase
11	ugpC	88.5 ± 2.7	1.9 ± 0.9	0.023 ± 0.011	-10.33	glycerol-3-phosphate trans-
						porter subunit
12	argI	44.5 ± 4.5	5.3 ± 2.6	0.138 ± 0.067	-10.24	ornithine carbamoyltransferase
						1
13	ttdA	86.1 ± 0.2	-0.0 ± 0.5	-0.000 ± 0.006	-10.06	L-tartrate dehydratase, alpha
						subunit
14	leuS	143.4 ± 3.7	-2.5 ± 0.5	-0.019 ± 0.003	-9.46	leucyl-tRNA synthetase
15	hisB	119.8 ± 3.7	-0.3 ± 3.0	-0.002 ± 0.027	-9.31	fusedhistidinol-
						phosphatase/imidazoleglycerol-
						phosphatedehydratase

Table S1 (contd.): 92 E. coli ORF-yfp fusions used for Leu CRI validation

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$	Gene product
		- Dea Tren	- Dea timitea			
16	rpoA	71.8 ± 7.5	0.0 ± 0.6	-0.001 ± 0.009	-9.05	RNA polymerase, alpha subunit
17	uvrY	173.2 ± 8.8	-5.2 ± 9.3	-0.027 ± 0.061	-8.36	DNA-binding response regula-
						tor in two-component regula-
18	rob	24.9 ± 0.7	0.5 ± 0.2	0.024 ± 0.007	-7.94	tory system with BarA right oriC-binding transcrip-
10	100	24.9 ± 0.7	0.0 ± 0.2	0.024 ± 0.007	-1.94	tional activator, AraC family
19	agaS	73.0 ± 3.3	1.3 ± 0.4	0.019 ± 0.005	-7.48	tagatose-6-phosphate ke-
						tose/aldose isomerase
20	lysS	111.8 ± 4.4	3.7 ± 2.1	0.035 ± 0.020	-7.17	lysine tRNA synthetase, consti-
21	sdaB	121.0 ± 5.7	36.4 ± 4.1	0.333 ± 0.044	-7.14	tutive L-serine deaminase II
$\frac{21}{22}$	purA	35.5 ± 2.2	-1.2 ± 0.4	-0.039 ± 0.015	-7.12	adenylosuccinate synthetase
23	rpoD	77.5 ± 2.2	2.3 ± 1.1	0.032 ± 0.014	-6.99	RNA polymerase, sigma 70
						(sigma D) factor
24	melR	44.5 ± 3.3	15.1 ± 0.4	0.378 ± 0.031	-6.76	DNA-binding transcriptional
25	kefF	99.2 ± 3.0	3.3 ± 0.5	0.038 ± 0.007	-6.52	dual regulator flavoprotein subunit for the
20	KCII	55.2 ± 5.0	0.0 ± 0.0	0.050 ± 0.001	-0.02	KefC potassium efflux system
26	uhpA	22.4 ± 1.5	0.4 ± 0.7	0.019 ± 0.034	-6.5	DNA-binding response regula-
						tor in two-component regula-
97	ndoN	41.6 ± 9.7	1.9 ± 0.7	0.050 + 0.015	C E	tory system with UhpB
27	ydcN	41.6 ± 2.7	1.9 ± 0.7	0.050 ± 0.015	-6.5	predicted DNA-binding tran- scriptional regulator
28	aspS	62.3 ± 4.2	10.9 ± 1.5	0.191 ± 0.014	-6.42	aspartyl-tRNA synthetase
29	relB	87.9 ± 2.3	0.8 ± 0.9	0.009 ± 0.010	-6.09	Qin prophage; bifunctional
						antitoxin of theRelE-RelB
						toxin-antitoxin system/ tran-
30	guaA	122.6 ± 2.3	6.9 ± 2.2	0.061 ± 0.019	-5.81	scriptional repressor GMP synthetase (glutamine
	84441	12210 ± 210	0.0 ± =.=	01001 ± 01010	0.01	aminotransferase)
31	phnM	203.7 ± 9.5	20.9 ± 1.2	0.113 ± 0.007	-5.81	carbon-phosphorus lyase com-
20	4.1.D	45 104	71 ± 0.9		F 70	plex subunit
32	tdcD	4.5 ± 0.4	7.1 ± 0.2	1.755 ± 0.163	-5.76	propionate kinase/acetate ki- nase C, anaerobic
33	ompR	114.2 ± 18.7	-1.4 ± 2.2	-0.019 ± 0.027	-5.74	DNA-binding response regula-
	_					tor in two-component regula-
2.4	1 T			0.000 + 0.010	F F 1	tory system with EnvZ
34	phnL	75.7 ± 5.3	5.7 ± 1.1	0.082 ± 0.013	-5.51	carbon-phosphorus lyase com- plex subunit
35	purH	29.6 ± 1.4	10.2 ± 0.4	0.383 ± 0.025	-5.27	fused IMPcyclohydrolase /
	1					phosphoribosyl aminoimidazole
	~					carboxamide formyltransferase
36	argG	2.5 ± 0.2	3.0 ± 0.2	1.343 ± 0.150	-5.07	argininosuccinate synthetase
37	rimM ubiC	107.5 ± 3.7	13.3 ± 0.9	0.137 ± 0.014	-4.95	16S rRNA processing protein
38 20		93.6 ± 3.9	7.5 ± 2.2	0.087 ± 0.022	-4.87	chorismate–pyruvate lyase leu operon leader peptide
39	leuL	48.9 ± 2.3	0.7 ± 0.5	0.017 ± 0.011	-4.75	
40	asnS ribB	9.0 ± 0.4	7.2 ± 0.5	0.884 ± 0.071	-4.65	asparaginyl tRNA synthetase
41	ridd	107.6 ± 7.8	12.2 ± 1.0	0.127 ± 0.020	-4.49	3,4-dihydroxy-2-butanone-4- phosphate synthase
42	smpB	15.8 ± 0.8	1.5 ± 0.4	0.101 ± 0.023	-4.39	trans-translation protein
43	guaB	31.0 ± 0.3	5.2 ± 0.6	0.184 ± 0.022	-4.21	IMP dehydrogenase
44	proC	69.2 ± 3.1	9.4 ± 1.6	0.151 ± 0.030	-3.92	pyrroline-5-carboxylate
						reductase, NAD(P)-binding
45	$_{\rm pth}$	89.0 ± 2.6	25.2 ± 0.9	0.313 ± 0.018	-3.1	peptidyl-tRNA hydrolase
46	ivbL	1.8 ± 0.3	1.6 ± 0.1	1.085 ± 0.295	-2.89	ilvB operon leader peptide
47	chbR	25.6 ± 1.0	24.4 ± 0.8	1.054 ± 0.008	-2.81	rRepressor, chb operon forN,N'-
48	leuA	78.1 ± 2.2	53.6 ± 0.8	0.756 ± 0.019	-2.77	diacetylchitobiose utilization 2-isopropylmalate synthase
40	argD	85.7 ± 4.4	53.0 ± 0.8 51.0 ± 1.8	0.657 ± 0.019	-2.72	bifunctional acetylor-
10		-	··			nithine aminotrans-
						ferase/succinyldiaminopimelate
20	C 3.	00.0 1.01	040 1 1 5		0.51	aminotransferase
50	yfcN	98.8 ± 2.1	24.2 ± 1.5	0.270 ± 0.011	-2.71	conserved protein

Table S1 (contd.): 92 E. coli ORF-yfp fusions used for Leu CRI validation

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	${\bf Robustness}$	$\log 2($ Leu CRI $)$	Gene product
51	ygiD	7.3 ± 0.3	9.9 ± 0.4	1.503 ± 0.046	-2.62	predicted dioxygenase, LigB
52	mdtJ	12.5 ± 1.6	17.5 ± 0.2	1.505 ± 0.204	-2.52	family multidrug efflux system trans-
53	agaR	12.5 ± 1.0 80.6 ± 2.5	31.4 ± 2.0	0.428 ± 0.018	-2.39	porter DNA-binding transcriptional
54	hisA	22.0 ± 0.9	27.6 ± 0.9	1.392 ± 0.087	-1.86	repressor of the aga regulon N-(5'-phospho-L-ribosyl-
01	111071	22.0 ± 0.0	2110 ± 010	1.002 1 0.001	1.00	formimino)-5-amino-1- (5'-phosphoribosyl)-4- imidazolecarb oxamide iso- merase
55	ygbF	72.7 ± 21.5	26.4 ± 1.3	0.475 ± 0.130	-1.65	probable ssRNA endonuclease, CRISP-associatedprotein
56	rpoH	89.2 ± 1.8	50.5 ± 0.9	0.623 ± 0.005	-1.53	RNA polymerase, sigma 32 (sigma H) factor
57	leuD	128.4 ± 2.7	70.0 ± 0.6	0.601 ± 0.018	-1.48	3-isopropylmalate dehydratase small subunit
58	dinJ	58.9 ± 2.8	34.4 ± 1.5	0.648 ± 0.050	-1.48	antitoxin of YafQ-DinJ toxin- antitoxin system
59	nuoI	25.2 ± 1.8	30.6 ± 1.4	1.361 ± 0.164	-1.48	NADH:ubiquinone oxidoreduc- tase, chain I
60	luxS	109.5 ± 0.7	70.3 ± 3.9	0.706 ± 0.035	-1.34	S-ribosylhomocysteine lyase
61	leuC	104.5 ± 6.0	54.0 ± 3.4	0.573 ± 0.069	-1.18	3-isopropylmalate dehydratase large subunit
62	pspA	69.8 ± 2.6	51.4 ± 0.6	0.813 ± 0.039	-1.18	regulatory protein for phage- shock-proteinoperon
63	pyrI	138.9 ± 13.7	112.5 ± 6.2	0.906 ± 0.091	-1.15	aspartate carbamoyltransferase, regulatorysubunit
$\begin{array}{c} 64 \\ 65 \end{array}$	btuEmsrB	149.1 ± 4.6 153.6 ± 14.7	$\begin{array}{c} 132.9 \pm 3.6 \\ 131.7 \pm 2.8 \end{array}$	$\begin{array}{c} 0.982 \pm 0.020 \\ 0.958 \pm 0.074 \end{array}$	-0.95 -0.8	glutathione peroxidase methionine sulfoxide reductase
						В
66	coaD	94.4 ± 0.3	96.2 ± 0.8	1.122 ± 0.012	-0.71	pantetheine-phosphate adeny- lyltransferase
67	sfsB	72.2 ± 2.4	18.9 ± 1.5	0.291 ± 0.032	-0.61	DNA-binding transcriptional activator of maltosemetabolism
68	nirD	113.2 ± 17.8	22.1 ± 1.5	0.223 ± 0.027	-0.52	nitrite reductase, NAD(P)H- binding, smallsubunit
69	fdnI	62.4 ± 1.6	57.8 ± 3.4	1.019 ± 0.034	-0.43	formate dehydrogenase- N, cytochrome B556 (gamma)subunit, nitrate- inducible
$70 \\ 71$	greA hupB	172.5 ± 12.4 287.5 ± 19.6	90.0 ± 1.2 116.5 ± 4.0	$\begin{array}{c} 0.581 \pm 0.045 \\ 0.450 \pm 0.030 \end{array}$	-0.38 -0.33	transcript cleavage factor HU, DNA-binding transcrip-
72	glpE	144.2 ± 2.0	110.0 ± 1.0 142.0 ± 2.6	1.086 ± 0.035	-0.33	tional regulator, betasubunit thiosulfate:cyanide sulfurtrans-
73	ogrK	95.8 ± 9.5	145.2 ± 3.3	1.699 ± 0.153	-0.33	ferase(rhodanese) positive regulator of P2 growth
10	0511	00.0 ± 0.0	110.2 ± 0.0	1.000 ± 0.100	0.00	(insertion of P2ogr gene into the chromosome)
74	rplD	20.9 ± 1.5	41.0 ± 1.6	2.191 ± 0.226	-0.33	50S ribosomal subunit protein L4
75	dmsB	116.2 ± 5.3	156.3 ± 5.1	1.486 ± 0.067	-0.28	dimethyl sulfoxide reductase, anaerobic, subunitB
76	rpsP	241.7 ± 3.7	118.5 ± 0.3	0.540 ± 0.007	-0.19	30S ribosomal subunit protein S16
77	rplX	150.6 ± 3.2	83.7 ± 3.5	0.613 ± 0.031	-0.19	50S ribosomal subunit protein L24
78	tpiA	138.5 ± 11.4	87.1 ± 2.2	0.700 ± 0.047	-0.19	triosephosphate isomerase
79	gapA	39.5 ± 2.0	62.6 ± 1.6	1.757 ± 0.135	-0.19	glyceraldehyde-3-phosphate de- hydrogenase A
80	rpsT	174.7 ± 12.0	133.7 ± 4.5	0.849 ± 0.050	-0.14	30S ribosomal subunit protein

Table S1 (contd.): 92 E. coli ORF-yfp fusions used for Leu CRI validation

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$	Gene product
81	rpsJ	170.2 ± 37.2	142.3 ± 1.6	1.015 ± 0.219	-0.14	30S ribosomal subunit protein S10
82	rplT	120.5 ± 8.5	112.6 ± 3.7	1.039 ± 0.077	-0.14	510 50S ribosomal subunit protein L20
83	ahpC	57.0 ± 5.0	64.9 ± 1.7	1.267 ± 0.078	-0.14	alkyl hydroperoxide reductase, C22 subunit
84	rpsK	109.9 ± 9.2	139.0 ± 3.5	1.411 ± 0.113	-0.14	30S ribosomal subunit protein S11
85	tsf	200.4 ± 2.8	103.8 ± 2.1	0.570 ± 0.008	0	protein chain elongation factor EF-Ts
86	rplU	173.3 ± 12.4	92.5 ± 4.9	0.595 ± 0.056	0	50S ribosomal subunit protein L21
87	rpmI	158.7 ± 4.0	104.2 ± 1.4	0.725 ± 0.025	0	50S ribosomal subunit protein L35
88	yjgF	131.9 ± 5.3	98.1 ± 3.0	0.824 ± 0.057	0	conserved protein, UPF0131 family
89	ppiB	133.6 ± 4.5	114.6 ± 2.7	0.945 ± 0.011	0	peptidyl-prolyl cis-trans iso- merase B (rotamaseB)
90	yjbJ	134.6 ± 4.4	158.7 ± 1.1	1.300 ± 0.035	0	conserved protein, UPF0337 family
91	rpsI	27.5 ± 2.8	78.6 ± 6.0	3.198 ± 0.324	0	30S ribosomal subunit protein S9
92	rpsF	31.7 ± 1.6	97.1 ± 0.8	3.392 ± 0.183	0	30S ribosomal subunit protein S6

Table S2: 56 E. coli ORF-yfp fusions used for Arg CRI validation

Genes are arranged by increasing values of Arg CRI. $S_{Arg-rich}$ and $S_{Arg-limited}$ refer to respective protein synthesis rates (a.u. per sec per cell). Robustness refers to the ratio between the two protein synthesis rates after normalization by the corresponding value for the AGA variant of yfp (this AGA variant was also used as the yfp tag in these ORF-yfp fusions). \pm refers to standard error of measurement.

Number	Gene	$S_{Arg-rich}$	$S_{Arg-limited}$	Robustness	$\log 2(\mathbf{Arg}\ \mathbf{CRI})$	Gene product
1	leuS	100.5 ± 2.6	16.1 ± 1.7	0.107 ± 0.012	-10.05	leucyl-tRNA synthetase
2	phoR	16.4 ± 3.2	0.7 ± 0.6	0.027 ± 0.029	-8.52	sensory histidine kinase in two- componentregulatory system with PhoB
3	glnG	61.0 ± 3.4	10.8 ± 2.3	0.118 ± 0.022	-8.51	fused DNA-binding response regulator intwo-component reg- ulatory system with GlnL: re- sponseregulator/sigma54 inter- action protein
4	asnS	15.5 ± 1.8	2.6 ± 0.1	0.114 ± 0.014	-8.38	asparaginyl tRNA synthetase
5	guaA	91.5 ± 4.2	11.7 ± 3.7	0.083 ± 0.024	-8.25	GMP synthetase (glutamine aminotransferase)
6	$_{\rm thiH}$	53.9 ± 1.1	4.4 ± 1.3	0.054 ± 0.015	-7.16	tyrosine lyase, involved in thiamin-thiazolemoiety synthe- sis
7	fruR	44.7 ± 1.5	5.3 ± 0.1	0.079 ± 0.004	-6.54	DNA-binding transcriptional dual regulator
8	argG	10.0 ± 0.9	2.5 ± 1.1	0.175 ± 0.088	-6.49	argininosuccinate synthetase
9	gpsA	81.9 ± 3.9	9.1 ± 4.2	0.073 ± 0.032	-5.81	glycerol-3-phosphate dehydro- genase (NAD+)
10	rpoA	60.9 ± 1.6	10.5 ± 2.1	0.114 ± 0.022	-5.36	RNA polymerase, alpha subunit
11	yiiD	31.2 ± 1.8	9.8 ± 2.9	0.218 ± 0.078	-5.05	predicted acetyltransferase
12	agaR	53.2 ± 1.8	2.6 ± 0.8	0.032 ± 0.010	-5.01	DNA-binding transcriptional repressor of the agaregulon
13	rimK	17.9 ± 2.6	9.8 ± 1.5	0.393 ± 0.106	-4.94	ribosomal protein S6 modifica- tion protein

Table S2 (contd.): 56 E. coli ORF-yfp fusions used for Arg CRI validation

Number	Gene	$S_{Arg-rich}$	$S_{Arg-limited}$	Robustness	$\log 2(Arg \ CRI)$	Gene product
14	rpoH	75.2 ± 2.8	9.3 ± 2.9	0.081 ± 0.023	-4.76	RNA polymerase, sigma 32
	-					(sigma H) factor
15	melR	43.7 ± 0.4	11.1 ± 2.3	0.170 ± 0.036	-4.46	DNA-binding transcriptional dual regulator
16	tdcB	32.1 ± 2.0	10.2 ± 2.9	0.208 ± 0.051	-3.83	catabolic threonine dehy- dratase, PLP-dependent
17	serC	68.2 ± 2.3	28.9 ± 4.2	0.284 ± 0.043	-3.79	3-
						phosphoserine/phosphohydroxy threonine aminotransferase
18	rnc	65.7 ± 2.5	15.4 ± 2.6	0.157 ± 0.028	-3.74	RNase III
19	chbR	22.3 ± 1.2	6.2 ± 2.1	0.190 ± 0.064	-3.66	rRepressor, chb operon forN,N'- diacetylchitobiose utilization
20	rsuA	100.3 ± 5.9	12.8 ± 2.8	0.084 ± 0.015	-3.65	16S rRNA U516 pseudouridine synthase
21	tauC	70.1 ± 3.2	7.4 ± 1.3	0.071 ± 0.016	-3.5	taurine transporter subunit
22	smpB	55.3 ± 1.3	17.5 ± 0.8	0.211 ± 0.010	-3.42	trans-translation protein
23	carA	28.2 ± 1.1	14.0 ± 0.8	0.331 ± 0.005	-3.4	carbamoyl phosphate syn- thetase small subunit, glutamine amidotransferase
24	ubiC	70.6 ± 1.8	4.7 ± 1.9	0.044 ± 0.018	-3.39	chorismate–pyruvate lyase
25	$_{\rm pth}$	71.2 ± 2.4	20.0 ± 0.8	0.187 ± 0.001	-3.21	peptidyl-tRNA hydrolase
26	rpsF	26.6 ± 1.9	14.7 ± 2.3	0.366 ± 0.043	-3.06	30S ribosomal subunit protein S6
27	gapA	34.3 ± 0.9	12.1 ± 1.9	0.234 ± 0.030	-2.86	glyceraldehyde-3-phosphate de- hydrogenase A
28	yihL	43.2 ± 3.4	6.7 ± 1.1	0.105 ± 0.022	-2.85	predicted DNA-binding tran-
29	allR	36.0 ± 1.5	17.1 ± 0.8	0.318 ± 0.026	-2.82	scriptional regulator DNA-binding transcriptional
						repressor for all(allantoin) and gcl (glyoxylate) operons;glyoxylate-induced
30	yfcN	74.2 ± 4.8	38.9 ± 7.3	0.353 ± 0.068	-2.8	conserved protein
31	fdnI	42.7 ± 1.8	17.8 ± 3.2	0.277 ± 0.042	-2.65	formate dehydrogenase- N, cytochrome B556
						(gamma)subunit, nitrate- inducible
32	ruvA	53.2 ± 4.1	26.7 ± 1.1	0.341 ± 0.040	-2.59	component of RuvABC resolva-
33	adiY	47.5 ± 8.2	6.1 ± 0.9	0.086 ± 0.003	-2.55	some, regulatorysubunit DNA-binding transcriptional
34	holD	15.1 ± 3.6	2.9 ± 1.3	0.153 ± 0.066	-2.43	activator DNA polymerase III, psi sub-
35	dmsB	68.1 ± 2.2	54.7 ± 6.5	0.537 ± 0.067	-2.42	unit dimethyl sulfoxide reductase,
36	bglJ	10.5 ± 1.3	6.6 ± 0.7	0.420 ± 0.010	-2.39	anaerobic, subunitB DNA-binding transcriptional
50	Dgi	10.0 ± 1.0	0.0 ± 0.7	0.420 ± 0.010	-2.03	activator for silentbgl operon,
						requires the bglJ4 allele to function; LuxRfamily
37	yfdT	71.4 ± 1.4	17.6 ± 0.7	0.164 ± 0.005	-2.32	CPS-53 (KpLE1) prophage;
38	argF	64.9 ± 2.8	35.7 ± 2.5	0.371 ± 0.044	-2.01	predicted protein ornithine carbamoyltransferase
39	rplU	102.0 ± 7.4	20.5 ± 4.2	0.134 ± 0.029	-1.89	2, chain F; CP4-6prophage 50S ribosomal subunit protein
40	luxS	90.3 ± 1.4	54.9 ± 2.4	0.406 ± 0.018	-1.77	L21 S-ribosylhomocysteine lyase
40 41	coaD	50.3 ± 1.4 70.9 ± 1.5		0.400 ± 0.018 0.426 ± 0.036	-1.75	pantetheine-phosphate adeny-
42	glnB	199.0 ± 6.0	61.9 ± 7.2	0.207 ± 0.021	-1.75	lyltransferase regulatory protein P-II for glu-
43	uidR	32.2 ± 1.7	35.7 ± 1.5	0.745 ± 0.064	-1.5	taminesynthetase DNA-binding transcriptional
'						repressor

Table S2 (contd.): 56 E. coli ORF-yfp fusions used for Arg CRI validation

Number	Gene	$S_{Arg-rich}$	$S_{Arg-limited}$	Robustness	$\log 2(\mathbf{Arg} \ \mathbf{CRI})$	Gene product
44	relB	69.6 ± 4.3	29.8 ± 0.5	0.287 ± 0.016	-1.35	Qin prophage; bifunctional antitoxin of theRelE-RelB toxin-antitoxin system/ tran- scriptionalrepressor
$\begin{array}{c} 45\\ 46\end{array}$	btuE argR	$\begin{array}{c} 106.1 \pm 1.4 \\ 37.3 \pm 3.4 \end{array}$	52.7 ± 4.0 13.7 ± 4.4	$\begin{array}{c} 0.331 \pm 0.021 \\ 0.235 \pm 0.054 \end{array}$	-1.35 -1.19	glutathione peroxidase DNA-binding transcriptional dual regulator,L-arginine-
47	$\mathrm{ogr}\mathrm{K}$	53.3 ± 1.5	29.3 ± 0.9	0.367 ± 0.013	-1.14	binding positive regulator of P2 growth (insertion of P2ogr gene into the shromescame)
48	hupB	127.5 ± 3.6	71.2 ± 1.0	0.373 ± 0.008	-1.13	chromosome) HU, DNA-binding transcrip- tional completant betagehumit
49	ppiB	95.0 ± 2.7	57.2 ± 10.5	0.401 ± 0.069	-1.13	tional regulator, betasubunit peptidyl-prolyl cis-trans iso- merase B (rotamaseB)
50	yjgF	92.2 ± 4.7	74.4 ± 3.6	0.540 ± 0.032	-1.13	conserved protein, UPF0131 family
51	kefF	68.6 ± 2.5	69.7 ± 7.6	0.677 ± 0.058	-1.11	flavoprotein subunit for the KefC potassiumefflux system
52	yjbJ	92.0 ± 5.1	96.6 ± 30.0	0.730 ± 0.272	-1.02	conserved protein, UPF0337 family
$53 \\ 54 \\ 55 \\ 56$	leuL rimM ydjO mdtJ	$\begin{array}{c} 36.7 \pm 1.4 \\ 86.4 \pm 1.7 \\ 9.9 \pm 0.9 \\ 21.0 \pm 0.4 \end{array}$	$\begin{array}{c} 31.8 \pm 2.2 \\ 89.9 \pm 3.3 \\ 20.7 \pm 1.6 \\ 26.4 \pm 1.5 \end{array}$	$\begin{array}{c} 0.577 \pm 0.023 \\ 0.694 \pm 0.029 \\ 1.432 \pm 0.219 \\ 0.841 \pm 0.060 \end{array}$	-0.97 -0.82 -0.74 -0.41	leu operon leader peptide 16S rRNA processing protein predicted protein multidrug efflux system trans- porter

Table S3: 21 E. coli ORF-yfp fusions co-expressed with GAG Leu2 tRNA

Genes are arranged by increasing values of Leu CRI as calculated for GAG Leu2 tRNA co-expression. $S_{Leu-rich}$ and $S_{Leu-limited}$ refer to respective protein synthesis rates (a.u. per sec per cell) under GAG Leu2 tRNA co-expression. Robustness refers to the ratio between the two protein synthesis rates after normalization by the corresponding value for the CTG variant of yfp (see Fig. 1D). \pm refers to standard error of measurement. Refer to Table S1 for corresponding values without GAG Leu2 tRNA co-expression.

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$	Gene product
1	ygiD	8.2 ± 0.6	3.8 ± 0.8	0.515 ± 0.103	-5.69	predicted dioxygenase, LigB family
2	chbR	26.5 ± 2.5	7.3 ± 1.5	0.299 ± 0.043	-5.68	rRepressor, chb operon forN,N'- diacetylchitobiose utilization
3	vfcN	92.7 ± 2.6	12.1 ± 2.9	0.142 ± 0.030	-5.47	conserved protein
4	mdtJ	12.7 ± 0.8	5.7 ± 1.7	0.513 ± 0.165	-4.46	multidrug efflux system trans- porter
5	ilvD	62.5 ± 1.5	23.6 ± 0.8	0.417 ± 0.022	-4.29	dihydroxyacid dehydratase
6	aspS	54.1 ± 2.0	18.5 ± 1.8	0.379 ± 0.046	-4.24	aspartyl-tRNA synthetase
7	lysS	106.3 ± 4.7	35.2 ± 0.8	0.367 ± 0.025	-4.22	lysine tRNA synthetase, consti- tutive
8	leuC	100.7 ± 0.5	19.0 ± 1.7	0.208 ± 0.019	-4.04	3-isopropylmalate dehydratase large subunit
9	ygbF	69.4 ± 3.4	8.1 ± 0.4	0.129 ± 0.009	-3.99	probable ssRNA endonuclease, CRISP-associated protein
10	pspA	65.1 ± 1.2	25.9 ± 2.0	0.437 ± 0.025	-3.91	regulatory protein for phage- shock-protein operon
11	guaA	122.0 ± 2.8	51.1 ± 1.9	0.462 ± 0.018	-3.27	GMP synthetase (glutamine aminotransferase)
12	btuE	137.7 ± 4.6	57.5 ± 3.9	0.462 ± 0.044	-3.23	glutathione peroxidase
13	purA	73.3 ± 1.8	34.8 ± 2.7	0.522 ± 0.035	-3.16	adenylosuccinate synthetase
14	ompR	116.4 ± 1.0	50.7 ± 3.2	0.479 ± 0.027	-2.86	DNA-binding response regula- tor in two-component regula- tory system with EnvZ

Table S3 (contd.): 21 E. coli ORF-yfp fusions co-expressed with GAG Leu2 tRNA

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$	Gene product
15	phnM	68.2 ± 4.4	48.4 ± 1.4	0.786 ± 0.044	-2.72	carbon-phosphorus lyase com- plex subunit
16	msrB	113.4 ± 3.1	37.9 ± 2.1	0.370 ± 0.028	-2.68	methionine sulfoxide reductase B
17	purH	71.4 ± 1.3	7.9 ± 2.0	0.120 ± 0.028	-2.61	fused IMPcyclohydro- lase/phosphoribosyl aminoim- idazole carboxamide formyl- transferase
18	coaD	98.3 ± 7.4	46.7 ± 1.9	0.526 ± 0.018	-2.58	pantetheine-phosphate adeny- lyltransferase
19	$\operatorname{rim}M$	114.3 ± 5.9	51.3 ± 1.1	0.498 ± 0.030	-1.99	16S rRNA processing protein
20	relB	79.9 ± 3.0	66.6 ± 3.3	0.923 ± 0.077	-1.73	Qin prophage; bifunctional antitoxin of theRelE-RelB toxin-antitoxin system/ tran- scriptional repressor
21	proC	78.1 ± 2.0	36.7 ± 4.4	0.519 ± 0.070	-1.72	pyrroline-5-carboxylate reductase,NAD(P)-binding

Table S4: 63 synonymous mutants of 13 E. coli ORF-yfp fusions

 $S_{Leu-rich}$ and $S_{Leu-limited}$ refer to respective protein synthesis rates (a.u. per sec per cell) of mutant ORFs. Robustness refers to the ratio between the two protein synthesis rates after normalization by the corresponding value for the CTG variant of yfp (see Fig. 1D). \pm refers to standard error of measurement. Refer to Table S1 for corresponding values of wild-type ORFs. The DNA sequence of the mutant variants below is provided in the gene_sequences.fasta file. Three of the sequences below did not have any mutations compared to the wild-type ORF and were included as internal controls.

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$
1	btuE-yfp mutant 1	137.0 ± 5.3	50.1 ± 1.1	0.403 ± 0.007	-2.22
2	btuE-yfp mutant 4	131.5 ± 3.5	20.5 ± 0.5	0.172 ± 0.009	-3.06
3	btuE-yfp mutant 5	138.9 ± 1.8	34.1 ± 0.1	0.270 ± 0.003	-2.62
4	btuE-yfp mutant 7	123.4 ± 4.9	48.2 ± 1.8	0.433 ± 0.032	-2.22
5	btuE-yfp mutant 8	130.7 ± 5.2	28.8 ± 0.9	0.243 ± 0.015	-3.06
6	btuE-yfp mutant 9	128.9 ± 0.9	40.0 ± 0.6	0.342 ± 0.003	-2.62
7	chbR-yfp mutant 1	29.5 ± 1.7	17.7 ± 0.3	0.663 ± 0.028	-2.81
8	chbR-yfp mutant 2	26.7 ± 2.2	15.1 ± 0.4	0.634 ± 0.072	-3.25
9	chbR-yfp mutant 3	31.7 ± 0.9	11.3 ± 0.6	0.393 ± 0.029	-5.31
10	chbR-yfp mutant 4	36.0 ± 2.5	9.0 ± 0.3	0.279 ± 0.027	-6.2
11	chbR-yfp mutant 5	28.9 ± 2.4	10.7 ± 0.2	0.414 ± 0.033	-5.76
12	chbR-yfp mutant 6	32.6 ± 2.7	5.7 ± 0.5	0.197 ± 0.021	-7.03
13	chbR-yfp mutant 7	32.1 ± 1.2	7.7 ± 0.5	0.267 ± 0.023	-5.8
14	chbR-yfp mutant 9	31.3 ± 0.9	7.9 ± 0.5	0.278 ± 0.026	-6.64
15	coaD-yfp mutant 1	92.3 ± 1.3	39.1 ± 1.3	0.467 ± 0.021	-1.89
16	coaD-yfp mutant 2	91.7 ± 4.1	55.5 ± 0.8	0.669 ± 0.034	-1.1
17	coaD-yfp mutant 3	94.0 ± 2.4	47.7 ± 2.3	0.558 ± 0.014	-1.5
18	leuA-yfp mutant 6	70.5 ± 2.1	11.3 ± 0.3	0.177 ± 0.010	-4.4
19	leuA-yfp mutant 8	77.9 ± 2.2	6.6 ± 1.4	0.093 ± 0.020	-5.23
20	leuA-yfp mutant 9	70.0 ± 2.1	8.4 ± 0.2	0.132 ± 0.007	-4.84
21	leuB-yfp mutant 7	273.5 ± 8.7	6.1 ± 0.9	0.025 ± 0.003	-4.81
22	leuB-yfp mutant 8	273.3 ± 10.6	9.6 ± 1.0	0.039 ± 0.005	-4.02
23	leuB-yfp mutant 9	274.9 ± 8.1	8.4 ± 1.7	0.034 ± 0.007	-3.97
24	leuC-yfp mutant 5	86.8 ± 0.6	8.5 ± 1.7	0.108 ± 0.021	-3.64
25	leuC-yfp mutant 7	95.6 ± 1.9	5.1 ± 1.7	0.059 ± 0.019	-3.3
26	leuC-yfp mutant 8	100.5 ± 1.7	4.3 ± 0.9	0.047 ± 0.009	-3.69
27	leuC-yfp mutant 9	80.0 ± 4.9	14.3 ± 0.8	0.198 ± 0.013	-3.25
28	leuD-yfp mutant 2	123.6 ± 0.9	18.3 ± 1.2	0.163 ± 0.011	-2.76
29	leuD-yfp mutant 3	123.1 ± 3.7	39.5 ± 1.0	0.355 ± 0.020	-1.48
30	leuD-yfp mutant 4	122.0 ± 1.0	32.9 ± 0.3	0.297 ± 0.002	-1.92
31	mdtJ-yfp mutant 1	17.3 ± 0.6	9.1 ± 0.4	0.580 ± 0.020	-4.15
32	mdtJ-yfp mutant 2	12.2 ± 0.5	5.8 ± 0.3	0.520 ± 0.008	-3.31
33	mdtJ-yfp mutant 4	10.0 ± 0.9	4.3 ± 0.1	0.475 ± 0.047	-4.54

Number	Gene	$S_{Leu-rich}$	$S_{Leu-limited}$	Robustness	$\log 2($ Leu CRI $)$
24		10.0 1 0.1			
34	mdtJ-yfp mutant 5	13.6 ± 2.1	11.4 ± 0.5	0.979 ± 0.190	-4.15
35	mdtJ-yfp mutant 6	6.4 ± 0.9	5.1 ± 0.1	0.903 ± 0.125	-3.75
36	mdtJ-yfp mutant 7	5.9 ± 3.1	4.3 ± 0.3	1.251 ± 0.488	-4.15
37	mdtJ-yfp mutant 8	10.2 ± 1.5	8.3 ± 0.1	0.932 ± 0.117	-4.19
38	mdtJ-yfp mutant 9	10.3 ± 0.8	9.6 ± 0.2	1.049 ± 0.087	-3.36
39	msrB-yfp mutant 1	76.6 ± 1.9	23.7 ± 0.8	0.342 ± 0.014	-2.48
40	msrB-yfp mutant 2	65.3 ± 1.6	20.7 ± 0.8	0.348 ± 0.005	-2.92
41	msrB-yfp mutant 3	99.1 ± 2.7	27.8 ± 2.5	0.310 ± 0.031	-2.52
42	msrB-yfp mutant 4	94.0 ± 1.5	36.2 ± 0.4	0.424 ± 0.005	-2.08
43	pspA-yfp mutant 1	68.2 ± 1.9	16.3 ± 0.6	0.264 ± 0.006	-2.81
44	pspA-yfp mutant 2	64.2 ± 0.5	15.0 ± 0.7	0.257 ± 0.011	-2.85
45	pspA-yfp mutant 3	62.3 ± 1.5	28.5 ± 0.6	0.505 ± 0.016	-1.97
46	pspA-yfp mutant 5	73.3 ± 1.3	14.0 ± 0.9	0.211 ± 0.018	-3.2
47	pspA-yfp mutant 6	70.1 ± 3.0	18.3 ± 0.9	0.288 ± 0.013	-2.76
48	pspA-yfp mutant 7	73.5 ± 1.4	8.5 ± 0.2	0.127 ± 0.002	-3.64
49	pspA-yfp mutant 8	73.0 ± 1.2	12.3 ± 0.8	0.185 ± 0.012	-3.25
50	yfcN-yfp mutant 1	103.0 ± 3.5	8.3 ± 1.3	0.088 ± 0.012	-4.82
51	yfcN-yfp mutant 5	86.0 ± 3.6	14.9 ± 1.7	0.192 ± 0.022	-3.1
52	yfcN-yfp mutant 6	90.5 ± 1.6	5.6 ± 0.7	0.069 ± 0.009	-5.31
53	yfcN-yfp mutant 8	91.5 ± 4.1	14.1 ± 1.7	0.170 ± 0.017	-2.71
54	yfcN-yfp mutant 9	85.5 ± 1.1	5.2 ± 0.4	0.067 ± 0.005	-4.87
55	ygbF-yfp mutant 2	80.6 ± 2.6	6.2 ± 0.3	0.085 ± 0.006	-3.28
56	ygbF-yfp mutant 3	68.5 ± 3.9	25.4 ± 1.4	0.413 ± 0.044	-1.65
57	ygbF-yfp mutant 5	79.0 ± 4.1	25.1 ± 0.2	0.352 ± 0.017	-2.1
58	ygbF-yfp mutant 6	65.3 ± 2.9	10.8 ± 0.6	0.183 ± 0.019	-2.44
59	ygbF-yfp mutant 7	80.5 ± 4.0	8.3 ± 0.2	0.113 ± 0.004	-3.33
60	ygbF-yfp mutant 8	70.4 ± 3.1	10.8 ± 0.9	0.170 ± 0.019	-2.89
61	ygiD-yfp mutant 1	11.0 ± 1.0	2.1 ± 0.2	0.216 ± 0.009	-5.17
62	ygiD-yfp mutant 3	11.3 ± 0.9	4.2 ± 0.3	0.417 ± 0.056	-4.73
63	ygiD-yfp mutant 4	10.6 ± 0.5	5.4 ± 0.6	0.572 ± 0.090	-4.34
64	ygiD-yfp mutant 5	7.2 ± 1.1	3.3 ± 0.5	0.514 ± 0.041	-5.17
65	ygiD-yfp mutant 6	6.8 ± 1.2	2.7 ± 0.6	0.476 ± 0.131	-4.29
66	ygiD-yfp mutant 7	12.3 ± 2.5	2.7 ± 0.4	0.251 ± 0.016	-6.06

Table S5: List of strains

Limiting AA	Strain designation	CGSC number	Genotype
Leu, Arg	CP78	4695	W3110, argH- leuB- thr- his- thi-
Ser	JW2880-1	10234	BW25113, $\Delta serA$
Pro	JW0233-2	8468	BW25113, $\Delta proA$
Ile	JW3745-2	10733	BW25113, $\Delta i lvA$
Gln	JW3841-1	10775	BW25113, $\Delta g lnA$
Phe	JW2580-1	10048	BW25113, $\Delta pheA$
-	MG1655	6300	wild-type strain with known genome sequence

Table S6: Concentration of amino acids and methyl esters used for amino acid limitation experiments

Amino Acid	Amino acid concen- tration in overnight cultures (μM)		Amino acid methyl es- ter concentration for amino acid limitation experiments (μM)	8
Leu	800	100	160	L1002 (Sigma)
Arg	800	150	160	11030 (Sigma)
Ser	10000	5000	800	412201 (Sigma)
Pro	800	0	160	
Ile	800	100	160	I0522 (VWR)
		0 100		287067 (Sigma)

Amino Acid		Amino acid concen- tration for amino acid limitation exper- iments (μM)		amino acid methyl es-
Gln	800	400	400	68604 (Astatech)
Phe	800	50	50	P17202 (Sigma)

Table S6 (contd.): Concentration of amino acids and methyl esters used for amino acid limitation experiments

Table S7: w_i values for Leu codons under ^{GAG}Leu2 co-expression

Codon	- log w_i (with ^{GAG} Leu2 co-expression)
CTA CTC CTG	0.48 0.05 0.04
$\begin{array}{c} \text{CTT} \\ \text{TTA} \\ \text{TTG} \end{array}$	$0.02 \\ 0.26 \\ 0.33$

Table S8: qPCR primer sequences

Gene	Forward primer	Reverse Primer
yfp	TCATGCTGTTTCATGTGATC	AGGGTGATGCTACTTATGGC
gapA	GCTGAAGGCGAAATGAAAGG	GTACCAGGATACCAGTTTCACG
rpoD	TGAAGCGAACTTACGTCTGG	AGAACTTGTAACCACGGCG
rpoS	TCTCAACATACGCAACCTGG	AGCTTATGGGACAACTCACG
rpoH	TCGTAATTATGCGGGCTATGG	CAGTGAACGGCGAAGGAG
rpoE	CCAGAAGGGAGATCAGAAAGC	TACCACATCGGGAACATCAC
rpoN	TGATCCAACTCTCCCAATTCG	TCGTGATTGGCTAACAGATCG
$\overline{f}ecI$	ACTACCACAGCTTCCTTAACG	TTTCGCTGACCATTACCCG
fliA	CGCTATGCTGGATGAACTTCG	CTAAACGTTCCGCTACCTCAG
leuA	GTCGCTAACTACAACGGTCG	GCACGCCAGATATTGTTCAG
ilvM	GTTTCCACGTCTGCTCAATG	CTGACTAAACAGTAAGTCGACCG
ilvB	TGAGTTTCCGTGTCCAATCC	ATCTGATGCTGACCAACGTC

Table S9: Codon-tRNA assignments

tRNA	Unmodified Anticodon	Modified Anticodon (if known)	Cognate Codons	Reference
Leu1	CAG	-	CUG	[15]
Leu2	GAG	-	CUC, CUU	15
Leu3	UAG	$\mathrm{cmo}^{5}\mathrm{UAG}$	CUA, CUG, CUU	[24]
Leu4	CAA	CmAA	UUA, UUG	[15, 25]
Leu5	UAA	$\mathrm{cmnm}^{5}\mathrm{UmAA}$	UUA, UUG	[15, 25]
Arg2	ACG	ICG	CGU, CGC, CGA	[15, 25]
Arg3	CCG	-	CGG	[15]
Arg4	UCU	$\mathrm{mnm}^{5}\mathrm{UCU}$	AGA, AGG	[18, 25]
Arg5	CCU	-	AGG	[15]
Ser1	UGA	$\mathrm{cmo}^{5}\mathrm{UGA}$	UCA, UCU, UCG	[15, 25]
Ser2	CGA	-	UCG	[15]
Ser3	GCU	-	AGC, AGU	[15]
Ser5	GGA	-	UCC, UCU	[15]
Pro1	CGG	-	CCG	[15]
Pro2	GGG	-	CCC, CCU	[15]
Pro3	\mathbf{UGG}	$ m cmo^5 UGG$	CCA, CCU, CCG	[15, 25]
Ile1	GAU	-	AUC, AUU	[15]

tRNA	Unmodified Anticodon	Modified Anticodon (if known)	Cognate Codons	Reference
Ile2	CAU	$k^2 CAU$	AUA	[15, 25]
Gln1	UUG	$\mathrm{mnm}^{5}\mathrm{s}^{2}\mathrm{UUG}$	CAA, CAG	[15, 25]
Gln2	CUG	-	CAG	[15]
Phe	GAA	-	UUC, UUU	[15]

Table **S9** (contd.): Codon-tRNA assignments