

Rapidly Characterizing the Fast Dynamics of RNA Genetic Circuitry with Cell-Free Transcription—Translation (TX-TL) Systems

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S Supporting Information

ABSTRACT: RNA regulators are emerging as powerful tools to engineer synthetic genetic networks or rewire existing ones. A potential strength of RNA networks is that they may be able to propagate signals on time scales that are set by the fast degradation rates of RNAs. However, a current bottleneck to verifying this potential is the slow design-build-test cycle of evaluating these networks *in vivo*. Here, we adapt an *Escherichia coli*-based cell-free transcription-translation (TX-TL) system for rapidly prototyping RNA networks. We used this system to measure the response time of an RNA transcription cascade to be approximately five minutes per step of the cascade. We also show that this response time can be



adjusted with temperature and regulator threshold tuning. Finally, we use TX-TL to prototype a new RNA network, an RNA single input module, and show that this network temporally stages the expression of two genes *in vivo*.

KEYWORDS: RNA genetic circuits, TX-TL, cell-free, response time, RNA synthetic biology

A central goal of synthetic biology is to control cellular behavior in a predictable manner.¹ Natural cellular behavior is governed by the expression of specific sets of genes needed for survival in different environments or developmental life stages. Genetic networks—webs of interactions between cellular regulatory molecules—are responsible for dynamically turning these genes on at the right time and place and, in effect, are the circuitry that implement behavioral programs in cells.² Because of this, a central focus of synthetic biology has been to control cellular behavior by engineering genetic networks from the bottom up.¹

Historically, work on engineering genetic networks has focused on combining sets of regulatory proteins to control their own expression in patterns that implement behaviors such as bistable memory storage,³ oscillations,^{4–6} layered logic gates,^{7,8} advanced signal processing,^{9,10} and spatial control of gene expression.¹¹ More recently, noncoding RNAs (ncRNAs) have emerged as powerful components for engineering genetic networks.¹² There are now examples of engineered ncRNAs that regulate nearly all aspects of gene expression,^{12–19} some as a function of intracellular molecular signals.^{15,16,20} In addition, new RNA structural characterization tools are enabling the engineering and optimization of these mechanisms.^{12,20–23} There are even large libraries of orthogonal RNA regulators,^{22–24} and there have been initial successes in engineering small genetic networks out of RNA regulators.^{17,19,25,26}

RNA genetic networks have several potential advantages over their protein counterparts.¹² First, networks constructed from RNA-based transcriptional regulators propagate signals directly as RNAs, thus eliminating intermediate proteins and making them potentially simpler to design and implement.¹⁹ Second, tools based on qPCR and next-generation sequencing have the potential to characterize the species, structural states, and interactions of RNAs across the cell at a level of depth and

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Figure 1. Schematic of the TX-TL design-build-test cycle for RNA circuits. Potential circuit designs are rapidly characterized in TX-TL by combining DNA-encoded RNA circuit components (colored circles) with the TX-TL reaction components. Overall circuit performance is monitored via the expression of fluorescent proteins enabling circuit designs to be rapidly benchmarked within a 2–3 h period. In addition, the openness of the TX-TL system allows characterization of circuit response via the addition of DNA encoded RNA regulators during the reactions. After multiple iterations of the design-build-test cycle, optimized circuit designs can be transformed into *E. coli* and tested for *in vivo* functionality.

detail not possible for proteins.¹² Finally, since the speed of signal propagation in a network is governed by the decay rate of the signal,²⁷ RNA networks have the potential to operate on time scales much faster than proteins. However, the design principles for engineering RNA circuitry are still in their infancy, and we have yet to fully test and verify these potential benefits. This is in part due to the slow nature of the current design-build-test cycle for engineering genetic networks *in vivo* that takes on the order of days even when testing circuits in bacteria with short doubling times.

Recently, cell-free protein synthesis systems have been developed into a platform to rapidly characterize the outputs of genetic networks.^{28–34} Cell-free reactions often consist of three components: cell extract or purified gene expression machinery, a buffer/energy mix optimized for gene expression, and DNA that encodes the genetic network^{28,35} (Figure 1). Fluorescent proteins are generally used as reporters, thus monitoring fluorescence over time allows the characterization of circuit dynamics. Because of their simplicity, cell-free reactions reduce the time for testing a constructed genetic circuit design from days to as little as an hour.^{33,36} Since these systems do not require selection markers or DNA replication to maintain circuitry constructs, there are no limitations on DNA circularization or on plasmid origin of replication and antibiotic compatibility.^{33,36} This flexibility allows for faster, multiplexed

generation of circuit constructs, further reducing design-buildtest cycle times. Since cell-free reactions lack a membrane, DNA encoding different regulators can be added at any time during the reactions, enabling the rapid characterization of network response as a function of perturbations that are extremely difficult or even impossible to do inside cells³⁴ (Figure 1). Additionally, there is increasing evidence that these *in vitro* characterizations correlate to *in vivo* results, including comparable rates of RNA degradation.^{29,33,36} Cell-free systems thus have intriguing potential to serve as an intermediate layer to rapidly prototype circuit design and response before porting the designs to the more complex environment of the cell.

In this work, we adapt an *E. coli* cell-free transcriptiontranslation (TX-TL) system^{28,37} for characterizing RNA genetic networks. Since this system was initially developed and optimized to test protein-based circuits,²⁹ we start by validating the functionality of RNA transcriptional attenuators¹⁹ in TX-TL and characterize the effect of different TX-TL experimental conditions including DNA concentration and batch-to-batch variation. We then show that a double-repressive RNA transcriptional cascade functions in TX-TL with characteristics that match its *in vivo* performance.¹⁹ The ability to spike in DNA encoding the top level of this cascade during the reaction allowed us to directly probe the response time of this RNA network. We found that the response time of this RNA cascade





Figure 2. Characterizing RNA transcriptional attenuators and circuits in TX-TL. (A) Fluorescence time courses of TX-TL reactions containing the pT181 attenuator reporter plasmid at 0.5 nM, with 8 nM antisense plasmid (+) or 8 nM no-antisense control plasmid (-). (B) SFGFP production rates were calculated from the data in (A) by calculating the slope between consecutive time points. Boxes represent regions of constant SFGFP production. Blue and red shaded regions in parts A and B represent standard deviations of at least seven independent reactions performed over multiple days calculated at each time point. (C) Average SFGFP production rates were calculated from the data in boxed regions in part B. Error bars represent standard deviations of those averages. The (+) antisense condition shows 72% attenuation compared to the (-) antisense condition in TX-TL. (D) Orthogonality of the pT181 attenuator (Att-1) to a pT181 mutant attenuator (Att-2). Average SFGFP production rates were calculated as in part C. Plots of SFGFP production rates can be found in Supporting Information Figure S2. Bars represent each attenuator at 0.5 nM with 8 nM of no-antisense control plasmid (blue), pT181 antisense plasmid (AS-1, red), or pT181 mutant antisense plasmid (AS-2, purple). (E) Schematic of an RNA transcriptional cascade. L1 is the same pT181 attenuator (Att-1) reporter plasmid used in parts A–D. In the plasmid transcribes the pT181-mut antisense (AS-2). (F) Average SFGFP production rates for the three combinations of the transcription cascade depicted in part E. L1 alone (blue bar) leads to high SFGFP production. L1+L2 (red bar) results in AS-1 repressing Att-1, thus lower SFGFP production. L1+L2+L3 (purple bar) results in a double inversion leading to high SFGFP production. Total DNA concentration in each reaction was held constant at 18.5 nM. In parts D and F, error bars represent standard deviations from at least seven independent reactions performed over multiple days.

is \sim 5 min per step of the cascade, matching our expectation of quick signal propagation due to the fast kinetics of RNA degradation. We then show that this response time can be tuned by either changing the temperature or effectively changing the threshold required for transcriptional repression by using tandem attenuators.¹⁹ To create a bridge to circuitry that we can implement and test in vivo, we show that we can use TX-TL to characterize the response time of similar cascades that use RNA regulators responsive to theophylline²⁰ to activate the cascade (Figure 1). The success of these experiments led to the forward design of a new RNA network motif, the single input module (SIM), which is responsible for staging the successive expression of multiple genes in natural pathways.³⁸ After characterizing the functionality of the individual SIM components in TX-TL, we transfer the final RNA-SIM circuit to E. coli, and show that this network dynamically stages the expression of two fluorescent reporter

proteins *in vivo*, solidifying the use of TX-TL for engineering RNA genetic circuits.

RESULTS AND DISCUSSION

RNA Transcriptional Cascades Function in TX-TL. In order to assess the feasibility of using TX-TL for characterizing RNA circuitry, we first tested the basic functionality of the central regulator in our RNA cascade, the pT181 transcriptional attenuator¹⁹ (Figure 2, Att-1). The pT181 attenuator lies in the 5'-untranslated region of a transcript, and functions like a transcriptional switch by either allowing (ON) or blocking (OFF) elongation of RNA polymerase.^{39,40} The OFF state is induced through an interaction with an antisense RNA (AS-1), expressed separately in our synthetic context¹⁹ (Supporting Information Figure S1). By transcriptionally fusing the pT181 attenuator to the super folder green fluorescent protein (SFGFP) coding sequence, we are able to assess functionality

of the attenuator by measuring SFGFP fluorescence with and without antisense RNA present (Supporting Information Figure S1).

To characterize antisense-mediated transcriptional repression in TX-TL, we first titrated concentrations of the pT181 attenuator (Att-1) reporter plasmid to determine an appropriate level of SFGFP output for our experimental setup. As expected, we observed a greater fluorescence output with increasing attenuator reporter plasmid concentration (Supporting Information Figure S2A). As noted in previous work with TX-TL, excessive amounts of input DNA can lead to resource competition and resource limitation within the reaction, which can confound circuit characterization.^{28,30,34,37} We found that an attenuator plasmid concentration of 0.5 nM (which corresponds to about one copy of plasmid into one *E. coli* cell) struck a balance between fluorescence signal and DNA concentration, and this concentration was used in subsequent experiments.

To test basic repression of the attenuator, we then characterized reactions that contained 0.5 nM of the attenuator reporter plasmid, and either 8 nM of the antisense-expressing plasmid (+), or 8 nM of a control plasmid that lacked the antisense coding sequence (-) (Supporting Information Figure S3 and Tables S3-4). As expected, we observed a substantial difference in the fluorescence trajectories between the (+) and (-) antisense conditions, with the (+) antisense condition resulting in an overall lower fluorescence output over time (Figure 2A). We note that in these experiments, we never observed a constant steady-state fluorescence signal due to the fact that SFGFP is not degraded (or is not diluted) in the TX-TL reaction during the time scale of the experiment (i.e., because SFGFP is not degraded, we always observed an increase in fluorescence over time even in the (+) antisense repressive condition).

Because of resource depletion effects that accumulate over time in TX-TL reactions, 2^{8} especially after 1–2 h of incubation, end points of these fluorescence trajectories can give misleading quantifications of repression. Since the action of the antisense RNA ultimately affects the rate of transcription of the attenuator, we sought a way to quantify antisense repression by comparing the rates of SFGFP expression.³³ Plotting the time derivative of the fluorescence trajectories allowed us to directly quantify the effective rate of SFGFP expression in the (+) and (-) antisense conditions as a function of time (Figure 2B). To further quantify antisense-mediated repression from these experiments, we computed a time average of the regions of constant maximum SFGFP production rate for each condition (Figure 2B boxed regions). We used maximum rate to reduce the confounding effects of resource depletion, which can cause production rates to go down over time (Figure 2B). Using these rates, we determined that with 8 nM of antisense RNA plasmid, we achieve 72% repression (Figure 2C), which is comparable to the 85% steady-state repression level previously observed in vivo.19,24

In order to move toward characterizing RNA circuitry in TX-TL, we needed to confirm the functionality of an orthogonal attenuator/antisense pair. We used a mutant pT181 attenuator/antisense pair (Att-2, AS-2) that had previously been shown to be perfectly orthogonal to the wild type attenuator *in vivo*.¹⁹ To test orthogonality in TX-TL, we performed reactions containing one of the two attenuator reporter plasmids at 0.5 nM, and one of the two antisense plasmids (AS-1 or AS-2) or the no-antisense control plasmid at 8 nM. For both attenuators, we found results consistent with those from *in vivo* experiments¹⁹—cognate antisense RNAs caused repression, while noncognate antisense yielded SFGFP expression rates that were within the error of the no-antisense conditions (Figure 2D, rate plots Supporting Information Figure S2B–D). We thus confirmed the orthogonality of the two attenuator/antisense pairs in TX-TL.

Interestingly, we observed that the region of maximum SFGFP production occurs at different times for different combinations of antisense-attenuators in the reaction. In particular, reactions with cognate (repressive) antisenseattenuator pairs have maxima that occur around 40 min, while reactions with noncognate (orthogonal), or just attenuator-reporter plasmids, have maxima that occur near the end of the reactions at ~100 min (Figure 2B, Supporting Information Figure S2B-D). Furthermore, cognate pairs show a decrease in SFGFP production rate for ~40 min after the maxima is reached (Figure 2B, Supporting Information Figure S2C). One reason for this decrease in production rate could be due to resource depletion caused by the cognate RNA-RNA interaction. Another reason could be from slow degradation of Att-1-SFGFP transcripts that escape attenuation at the start of the reaction. In fact, previous work has shown that the half-life in TX-TL of deGFP mRNA monitored by the malachite green aptamer⁴¹ is approximately 18 min.⁴² Independent of a specific cause of this effect, using maximum SFGFP production rate gives us a conservative estimate of attenuator repression that we use below.

The orthogonality of the two antisense-attenuator pairs allowed us to characterize a double-repression RNA transcriptional cascade in TX-TL (Figure 2E).¹⁹ The bottom level of the cascade (L1) consists of an SFGFP coding sequence controlled by the pT181 attenuator's (Att-1) interaction with its antisense (AS-1). AS-1 transcription is in turn controlled by the mutant antisense (AS-2), via a mutant attenuator sequence (Att-2) present upstream of AS-1 on the middle level of the cascade (L2). AS-2 is transcribed from the top level of the cascade (L3) (Figure 2E).

Previous work encoded L2 and L3 of the cascade on a highcopy plasmid (ColE1 origin, ~200 copies/cell), and L1 on a medium copy plasmid (p15A origin, ~15 copies/cell), and showed that the double repression cascade yielded a net activation of SFGFP expression at steady-state in vivo.¹⁹ Since there are no plasmid incompatibilities in TX-TL, we were able to use three separate plasmids for L1, L2, and L3 to characterize cascade function. Since three DNA elements are needed for the cascade, we first performed titrations of L2 versus 0.5 nM of L1 in order to find conditions that allowed us to observe repression without severe resource depletion (Supporting Information Figure S2E). We found that 4 nM of L2 caused a 72% repression, with no greater repression observed with higher concentrations of L2. To test the full cascade, we titrated different amounts of L3, from 10 nM to 18 nM, keeping the total amount of DNA in the TX-TL reaction constant with the addition of a control plasmid to approximately control for resource usage across conditions (Supporting Information Figure S2F). We found that L3 activates SFGFP expression with a rate that matches that of just L1 (Figure 2F).

This result proved that RNA circuitry functions in TX-TL reactions. Stated another way, these experiments demonstrate that the RNA circuitry tested only requires the machinery from the cytoplasmic extract contained in the TX-TL reaction. Furthermore, the flexibility of TX-TL allowed us to systemati-

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cally validate the functioning of each level of the cascade by adding successive levels one at a time. This is in contrast to the previous *in vivo* work where complex controls were needed to validate cascade performance since L2 and L3 were encoded on the same plasmid.¹⁹ Finally, we note that all of these experiments were performed at 29 °C, whereas the RNA transcriptional regulation and circuitry had only been previously tested at 37 °C, thus confirming its function over a range of temperatures.

Ideal TX-TL Batch Characteristics for Circuit Testing. There is known to be batch-to-batch variation in TX-TL preparations.²⁸ In order to assess the impact of batch-to-batch variation on RNA circuit characterization, we tested three different extract/buffer preparations by adding a range of concentrations of the no-antisense control plasmid to 0.5 nM of the L1 plasmid. Since extra control DNA causes resource competition, this experimental design allowed us to assess the maximum amount of DNA per reaction that each batch could support. As shown in Figure 3, we observed several important



Figure 3. Assessing batch-to-batch variation. (A) Fluorescence time courses of TX-TL reactions in three different extract and buffer preparations with 0.5 nM L1 and 15 nM no-antisense control DNA. Shaded regions represent standard deviations of at least 11 independent reactions over multiple days calculated at each time point. (B) Average maximum SFGFP production rates for the same three buffer and extract preparations from reactions with 0.5 nM L1 and 0, 5, 10, 15, and 20 nM no-antisense control DNA. Plots of maximum SFGFP production rates from which these were calculated can be found in Supporting Information Figure S4. Error bars represent standard deviations from at least 11 independent reactions.

features. First, for a fixed concentration of L1 and control DNA, we observed significant differences in the fluorescence time courses between the batches (Figure 3A). The end point fluorescence of batch 2 is more than twice that of batch 1 and 3. Second, batch 2 reaches constant SFGFP production faster than batch 1 and 3 for all conditions tested (Supporting Information Figure S4). Third, batch 1 had a lower fluorescence output than batch 3, but both reached constant SFGFP production at approximately the same time over all conditions.

In terms of DNA loading effect, in all of the batches we see an increase in production rate when adding 5 nM of control plasmid (Figure 3B). For batches 1 and 3, production rates remained constant up to the maximum concentration of control DNA tested (20 nM). The production rate for batch 2 increased with the addition of 10 nM control DNA and then decreased for 15 and 20 nM control DNA (Figure 3B).

We hypothesize that the increase in production rate for all batches upon adding 5 nM control DNA is due to competition effects from the RNA degradation machinery (RNases). The control DNA has the same promoter as the attenuator and antisense plasmids from which an RNA terminator is made (Supporting Information Figure S3). While this RNA does not affect the attenuator in a mechanistic way, it does provide a decoy for RNases and could cause a decrease in degradation rate of the attenuator-SFGFP mRNA, and thus an overall increase in SFGFP production. The further increase in production rate from 5 to 10 nM control DNA in batch 2 suggests that different batches have different amounts of RNase machinery. To test this hypothesis, we added 0-250 ng of total yeast RNA to 0.5 nM of L1 plasmid in the three buffer/extract batches (Supporting Information Figure S5). The yeast RNA would provide a decoy for the RNases but should not sequester the bacterial translation machinery in TX-TL. As shown in Supporting Information Figure S5, we observed an increase in production rate with the addition of yeast RNA for all batches. This increase was more pronounced for batch 2, which responded to a lower concentration of yeast RNA, while it took a higher concentration of yeast RNA to see an effect for batches 1 and 3. These results support our conclusion from the control DNA experiments that there is batch-to-batch variation in RNase machinery concentration that contributes to the DNA loading effect. Because of this loading effect, we found it important to use an appropriately constructed control DNA when designing comparative experiments (Supporting Information Figure S3).

Additionally, these results show the importance of screening TX-TL extract and buffer batches to best match the needs for circuitry characterization. For our RNA attenuator circuitry, we chose a batch that struck a balance between high production rate for better signal/noise and invariance to DNA loading effects for testing circuitry with multiple components. We therefore used batch 3 in all further experiments.

Characterizing the Dynamics of RNA Circuitry with TX-TL. One of the potential advantages of RNA over protein circuitry is a faster response time due to the relatively fast degradation of RNA molecules.¹² The flexibility of TX-TL allowed us to directly measure the response time of the RNA transcription cascade (Figure 4). Since TX-TL is an open system, we designed an experiment that involved spiking in the DNA encoding L3 of the cascade into an ongoing reaction that was already expressing L1 and L2. We define the response time of this circuit, τ , as the time it takes to turn ON SFGFP



Figure 4. Determining cascade response time. (A) Schematic of spike experiment. L3 (or the no-antisense control plasmid) was spiked into an ongoing L1+L2 TX-TL reaction at time, t = 0 (represented by dashed box). Concentrations of DNA used are indicated beside the levels. (B) Normalized fluorescence curves combining three separate experiments performed at 29 °C with a total of 8 replicates over multiple days. An L1 (0.5 nM) + L2 (4 nM) reaction was setup for 25 min at which point L3 (14 nM, puple curve) or no-antisense control DNA (14 nM, red curve) was spiked into the reaction and time reset to 0. Inset shows the response time of the circuit to the addition of L3; defined as the time at which the L3 spike curve is statistically different from the L1+L2 curve ($\tau = 18.2 \pm 6.0$ min). (C) Normalized fluorescence curves combining three separate experiments performed at 37 °C with a total of 11 replicates over multiple days. The same experiment was setup as in part B except that the L1+L2 reaction ran for 20 min prior to the addition of L3. τ = 14.6 ± 4.8 min. Shaded regions represent standard deviations calculated at each time point.

production after spiking in L3. In order to determine τ , TX-TL reactions were setup with 0.5 nM L1 and 4 nM L2 following

our earlier results (Figure 2). This reaction was allowed to proceed for 25 min, at which time (t = 0) we spiked in 14 nM of either L3, or our no-antisense control plasmid. Fluorescence trajectories showed that the L3 spike caused a noticeable deviation from the control trajectory ~20 min after the spike (Figure 4B). By using Welch's *t* test to find the point at which the two trajectories differed significantly (Supporting Information Figure S6, Methods), we were able to quantify the response time over multiple experimental replicates to be 18.2 \pm 6.0 min (Figure 4B).

We can estimate τ by considering the three molecular events that need to occur to turn ON SFGFP production: (i) AS-2 needs to be transcribed from L3, (ii) the concentration of AS-2 needs to build up in order to repress the transcription of AS-1 via an interaction with Att-2, and (iii) any existing AS-1 must be degraded. Using a simple ordinary differential equation model of the expression of each level of the RNA cascade, we can derive an expression for the response time (Supporting Information Appendix 1):

$$\tau \sim \ln(2) \left(\frac{1}{d_2} + \frac{1}{d_1} \right) + \alpha \tag{1}$$

where d_2 and d_1 are the degradation rates of AS-2 and AS-1, respectively, and α is the maturation time of SFGFP. Since $\ln(2)/d$ is the half-life of each antisense species, we find that the response time is a sum of half-lives of the intermediate RNA signals, similar to a related analysis of response times of protein circuitry.^{2,27} The importance of antisense degradation rates for estimating circuit response times led us to determine approximate RNA degradation rates in our TX-TL batch. To do this, we used the fluorescent malachite green RNA aptamer as a representative small RNA (45 nucleotides), which allows a convenient fluorescence based assay to measure its abundance.⁴² By spiking in purified malachite green aptamer and ligand in TX-TL reactions and observing fluorescence decay, we measured its half-life in TX-TL batch 3 to be 2.7 \pm 0.44 min at 29 °C (Supporting Information Figure S7). While RNA degradation rates are structure and length dependent, this value is close to a previously determined half-life of the pT181 antisense RNA (91 nucleotides) of 5 min in vivo.⁴³ Using this value and a 5 min maturation time for SFGFP,44 we can estimate τ from eq 1 to be 15 min, in close agreement with our experimental observation.

In E. coli, RNA degradation is primarily controlled by the RNA degradasome, a multiprotein complex that degrades RNA species.⁴⁵ Since RNA degradation is enzymatic, we expect its rate to increase with increasing temperature, and thus expect increasing temperature to decrease τ . To test this, we repeated both the malachite green degradation and DNA spike experiments with reactions running at 37 °C and determined the malachite green aptamer half-life to be 1.4 \pm 0.08 min (Supporting Information Figure S7) and τ for the RNA cascade to be 14.6 ± 4.8 min (Figure 4C). This response time is in remarkable agreement to our 15 min estimate of τ from eq 1, which was derived from antisense degradation rates measured at 37 °C in vivo.43 While the average response time at 37 °C was lower than that at 29 °C, the difference between the two averages was not statistically different due to the large error bars on the measurements (p = 0.1694, Supporting Information Table S2).

These results represent the first measurement of RNA circuitry response times. Furthermore, they confirm our

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Figure 5. Determining the response time to tandem attenuators. (A) Schematic of a single input module (SIM). Colored circles represent nodes of the circuit, a repression cascade in which two genes (purple circles) are temporally controlled by a single species (blue circle). Temporal control is accomplished by varying thresholds of action (β_1 , β_1') for the intermediate cascade species (green circle). Plots are schematic response curves for the spike experiment described in Figure 4. The regulatory species (blue) is added in at t = 0. This species shuts off production of the intermediate species (green) once threshold β_2 is reached at time δ_2 (see Supporting Information Appendix 1). The final genes are activated after time τ and τ' once the intermediate species falls below the thresholds β_1 and β_1' , respectively.² (B) Schematic of spike experiment. Experimental setup was analogous to Figure 4A except that L1 contains tandem pT181 attenuators (Att-1-Att-1). (C) Normalized fluorescence curves combining three separate experiments performed at 37 °C with a total of 12 replicates over multiple days. L1 (0.5 nM) + L2 (4 nM) reaction was setup for 20 min at which point L3 (14 nM, purple curve) or no-antisense control DNA (14 nM, red curve) was spiked into the reaction and time reset to 0. Inset shows the response time of the circuit to the addition of L3 ($\tau = 19.4 \pm 5.0$ min). Shaded regions represent standard deviations calculated at each time point.

expectation of a quick response time that is dependent on the degradation rates of the intermediate RNA species in the network. Our simple estimate shows that for circuits constructed from the attenuation system, we can expect the response time to be ~5 min for each step in the circuit. This is in stark contrast to an analogous protein-mediated cascade, which has been shown to have a response time of ~140 min as measured *in vivo.*⁴⁶ In fact, because protein degradation is typically very slow, the response times of protein-mediated circuitry is often set by the rate of cell division, which is the dominant source of protein decay.²⁷

Cascade Response Time Can Be Tuned by Using Tandem Attenuators. The success of the cascade response time measurements led us to begin the forward design of an RNA version of the single input module (SIM).³⁸ The SIM is a network motif in which a single regulatory molecule controls the expression of multiple outputs (Figure 5A). In nature, SIMs are used to regulate the genes of biosynthetic pathways and stress response systems so that they are expressed in the order at which they are needed.² The SIM provides this temporal regulation by encoding different regulatory thresholds of each gene. As the master regulator increases in concentration, it successively traverses these thresholds and thus activates genes at different times³⁸ (Figure 5A).

The ability to configure tandem attenuators upstream of a coding sequence¹⁹ provides a mechanism to adjust the threshold at which the RNA cascade responds to antisense concentrations (Figure SB). Since tandem attenuators are more sensitive to antisense RNA concentration,¹⁹ we hypothesized that the response time of a cascade with tandem attenuators in L1 (Att-1-Att-1) would be slower than that of the single attenuator cascade—that is, it would take a longer time for AS-1 to decay below the threshold required for repressing the Att-1-Att-1 attenuator, thus causing a longer response time (Figure SA). We repeated the L3 spike experiment with the Att-1-Att-1 cascade at 37 °C and determined the response time to be 19.4 \pm 5.0 min (Figure SC, Supporting Information Table S1). This

response time was statistically significant from the single Att-1 cascade at 37 °C (p = 0.0303, Figure 4C) and again matched the estimation of 20 min based on a modified version of eq 1 that takes into account the different threshold of the tandem Att-1-Att-1 (Supporting Information Appendix 1). We thus showed that the Att-1-Att-1 tandem attenuator could be effectively used to tune cascade response time, making it suitable for its use as a component in designing an RNA SIM.

Theophylline Responsive Antisense Provides a Bridge to Move RNA Circuitry In Vivo. While the tandem attenuator cascade provided a necessary component of the RNA SIM, we have thus far been probing circuit response time by spiking in antisense RNA via a DNA plasmid-a perturbation not possible in an in vivo experiment. We therefore changed L3 to encode the theophylline-responsive antisense RNA developed by Qi et al.,²⁰ which has previously been shown to only attenuate transcription in the presence of theophylline in vivo. The response time probing experiment was adjusted so that either theophylline (2 mM in the reaction) or water (as a control) was spiked into each TX-TL reaction containing L1, L2, and aptamer-L3 (Figure 6A). The response time was then calculated by comparing with (+) and without (-) theophylline fluorescence trajectories in the same way as described previously. This was done for both the single (Att-1) and tandem (Att-1-Att-1) versions of the cascade resulting in response times of 59.3 \pm 7.3 min and 45.2 \pm 11.7 min respectively (Figure 6B-C, Supporting Information Table S1).

Notably, both response times are greater than what we observed in the DNA spike experiments (Figures 4, 5). To verify this was not an issue due to theophylline toxicity, we first tested if theophylline was toxic to the basic TX-TL reactions (Supporting Information Figure S9). While there was a 20% decrease in SFGFP production rate from our L1 (Att-1) construct with 2 mM theophylline (Supporting Information Figure S9A), it did not increase the response time of the core RNA cascade, as determined by spiking in theophylline with L3



Figure 6. Determining the response time to a theophylline regulated cascade. (A) Schematic of experiment. L3 of the cascade in Figures 4A and 5A has been replaced with AS-2 fused to a theophylline aptamer from Qi et al.²⁰ AS-2-theo is only active in the presence of theophylline. Theophylline was spiked into an ongoing L1 + L2 + aptamer-L3 TX-TL reaction at t = 0 (represented by dashed box). (B) Single attenuator (Att-1) cascade. Normalized fluorescence curves combining three separate experiments performed at 37 °C with a total of 12 replicates over multiple days. L1 (Att-1, 0.5 nM) + L2 (4 nM) + aptamer-L3 (14 nM) reaction was setup for 20 min at which point theophylline (final concentration 2 mM, puple curve) or ddH₂O (red curve) was spiked into the reaction and time reset to 0. Inset shows the response time of the circuit to the addition of theophylline ($\tau = 59.3 \pm 7.3$ min). (C) Tandem attenuator (Att-1-Att-1) cascade. Normalized fluorescence curves combining three separate experiments at 37 °C with a total of 9 replicates over multiple days. L1 (Att-1-Att-1, 0.5 nM) + L2 (4 nM) + aptamer L3 (14 nM) reaction was setup for 20 min at which point theophylline (final concentration of theophylline ($\tau = 45.2 \pm 11.7$ min). Shaded regions represent standard deviations calculated at each time point.

DNA to mimic the experiments in Figure 4 (Supporting Information Figure S9B).

Interestingly, we also observed that the response time for the Att-1 cascade was longer than that of the Att-1-Att-1 cascade in the theophylline spike experiments. We attribute this to the dip in the Att-1 fluorescence trajectory with (+) theophylline seen between 0 and 45 min (Figure 6B, Supporting Information Figure S8), which is not present in the Att-1-Att-1 (+) trajectory (Figure 6C). In the theophylline spike experiments, we hypothesize that the presence of unbound aptamer-AS-2 that builds up in concentration during the prespike reaction competes with AS-1 for RNA degradation machinery, or causes general resource depletion effects, which slows overall reaction rates. Because of the high concentration of aptamer-AS-2 used in these experiments (14 nM), the aptamer-AS-2 RNAs would be present in a higher concentration than AS-1. This could cause a bottleneck in RNA degradation that would lead to a dip in the fluorescence trajectory. The SFGFP production rate could transition from a slow to a fast phase once the excess unbound aptamer-AS-2 is cleared from the reaction allowing AS-1 to be degraded and the cascade to be fully activated. In addition, resource depletion caused by the extra 14 nM of aptamer-AS-2 DNA in the prespike incubation period could lower overall production rates leading to a longer than expected response time. We note that we do not observe the dip in the Att-1-Att-1 cascade, which could be due to the overall lower SFGFP expression from the Att-1-Att-1 construct and the noise of the experiments.

To test these ideas, we performed an experiment in which the aptamer-L3 DNA and theophylline (or water as a control) were cospiked into ongoing TX-TL reactions at the same time (Supporting Information Figure S10). While this experiment does not provide a bridge to cells as we are spiking in a DNA construct, it does remove the dip from the with (+) theophylline curve, bringing the Att-1 response time down to 41.9 ± 16.9 min. This result supports our hypothesis of either a bottleneck in RNA degradation or depletion of TX-TL resources leading to the dip in Figure 6B.

Even without the confounding trajectory dips, the response times of the theophylline-mediated (Att-1 and Att-1-Att-1)



Figure 7. RNA single input module (SIM) functions *in vivo*. (A) Schematic of the network motif. L1 contains Att-1-RFP and Att-1-Att-1-SFGFP. Expression of both proteins is controlled by AS-1 in L2, which in turn is controlled by the interaction of Att-2 with its antisense AS-2-theo (aptamer-L3). AS-2-theo is a fusion with the theophylline aptamer, which is only active when the aptamer is in the bound state.²⁰ All three plasmids were cotransformed into *E. coli* TG1 cells. Plasmid origins are noted by the cascade levels. Theophylline is added to one of the split cultures once in logarithmic growth at which point time was set to zero (represented by dashed box). (B) Normalized fluorescence time courses for cultures with (+) and without (-) theophylline at 2 mM. Response time, τ , was calculated by determining the time at which the (+) and (-) curves were statistically different. τ (RFP) = 41.7 ± 13.4 min. τ (SFGFP) = 40.0 ± 9.5 min. (C) Schematic of the network motif in (A) with fluorescent reporters switched in L1. L1 for this network contains Att-1-SFGFP and Att-1-Att-1-RFP. L2 and aptamer-L3 remain the same. (D) Normalized fluorescence time courses for cultures with (+) and without (-) theophylline at 2 mM. τ (SFGFP) = 42.5 ± 10.6 min. τ (RFP) = 72.7 ± 20.5 min. Shaded regions represent standard deviations calculated from 12 independent transformants at each time point.

cascades are still much slower than the nontheophyllinemediated cascades. The DNA/theophylline cospike experiment (Supporting Information Figure S10) eliminated any experimental setup differences leading to the slower than expected response times. From eq 1, we see that the response time is governed by the degradation rates of the two intermediate species. It could be that structural differences caused by the aptamer sequence or bound ligand alters the stability of the aptamer-AS-2 RNA enough to account for these observed differences. However, the ability to observe cascade activation upon addition of theophylline provided the necessary bridge to move forward with the RNA SIM construction and characterization *in vivo*.

An RNA SIM Functions *In Vivo*. Our success in demonstrating different response times using single and tandem attenuators in TX-TL led us to design an RNA SIM and

characterize its function *in vivo*. To construct the SIM, we combined the single and tandem attenuator cascades in a single circuit controlling the expression of two different fluorescent proteins, red fluorescent protein (RFP) and SFGFP, respectively (Figure 7A).

The first step in implementing the proposed RNA SIM was to develop a three-plasmid version of the cascade. To do this, we placed L1 on a low copy pSC101 backbone with kanamycin resistance, L2 on a medium copy p15A backbone with chloramphenicol resistance, and L3 on a high copy ColE1 backbone with ampicillin resistance (Supporting Information Figure S11A). A steady state test of this circuit in E. coli TG1 cells showed 68% attenuation with L1+L2 present, and recovery of 68% of the L1 only signal with the full cascade present (Supporting Information Figure S11B). Differences between these attenuation and recovery levels, and those observed in TX-TL (Figure 2F) could be due to plasmid concentrations not obeying the same DNA concentration ratios that we used in TX-TL. However, our observation of recovery upon adding L3 in vivo was sufficient to measure the response time of the circuit.

To construct the SIM in the three-plasmid architecture, we placed a single pT181 attenuator in front of the RFP coding sequence followed by tandem pT181 attenuators in front of the SFGFP coding sequence on the same pSC101 backbone, each under the control of its own promoter (Supporting Information Figure S12A). This plasmid was cotransformed into E. coli TG1 cells along with the L2 and aptamer-L3 plasmids to complete the SIM motif. After overnight growth of replicate colonies, cultures were split in pairs and subcultured in minimal media for 4 h until logarithmic growth was reached. At this point, theophylline was added to one of the subcultures from each colony to a final concentration of 2 mM. RFP and SFGFP fluorescence was then monitored every 10 min for a total of 90 min (see Materials and Methods). RFP and SFGFP fluorescence trajectories for the with and without theophylline conditions are shown in Figure 7B. From these curves, we calculate a response time of 41.7 ± 13.4 min for Att-1-RFP and 40.0 ± 9.5 min for Att-1-Att-1-SFGFP.

On the surface, the Att-1 and Att-1-Att-1 response times are similar. However, after correcting for the slower maturation time of RFP compared to SFGFP (42 min⁴⁷ and 5 min,⁴⁴ respectively), we find the Att-1-Att-1 element is activated 35 min later than the Att-1 element according to the SIM design. We verified this finding by switching the protein reporters between the Att-1 and Att-1-Att-1 regulators. In this orientation, L1 contained Att-1-SFGFP and Att-1-Att-1-RFP constructs (Figure 7C, Supporting Figure S12B). Using this L1 we would expect to see a distinctive difference in the response times since the slower maturing protein monitors the slower responding Att-1-Att-1 element. The experiment was repeated in the same manner as above and we calculated a response time of 42.5 \pm 10.6 min for Att-1-SFGFP and 72.7 \pm 20.5 min for Att-1-Att-1-RFP (Figure 7D), again providing evidence of a functioning SIM.

Comparing the two SIM versions allows a further check on SIM function. In fact, the Att-1-Att-1-RFP response time (Figure 7D) is approximately 30 min longer than Att-1-Att-1-SFGFP response time (Figure 7B) as we would expect from the longer maturation time of RFP compared to SFGFP. However, the Att-1-SFGFP response time (Figure 7D) was slower than expected, and was within error of the Att-1-RFP response time (Figure 7B). This could be due to the increase in the Att-1SFGFP (-) theophylline fluorescence trajectory (Figure 7D) causing a longer calculated response time. This increasing background trajectory could be due to incomplete repression of L1 by L2 under the (-) theophylline condition, which we observed in the incomplete repression of Att-1-SFGFP by L2 at steady state (Supporting Information Figure S11B). On the other hand, the Att-1-Att-1 constructs would not be affected in the same way since Att-1-Att-1 offers tighter control.¹⁹ In future applications, the OFF level of the RNA SIM could be improved by using Att-1-Att-1 vs Att-1-Att-1 constructs or ribosome binding site (RBS) optimization of the protein output to reduce leak. The constructs used in this work were developed for high protein expression, thus a strong RBS was used, however, RBS tuning has been shown to improve the antisense-attenuator OFF state.¹⁹

Despite the Att-1-SFGFP inaccuracy, the two orientations of the SIM confirm that we observe a clear difference in response time between the Att-1 and Att-1-Att-1 components of the RNA SIM in *E. coli*. This network motif allows for temporal control of two genes in response to a theophylline signal, and could have applications in a variety of contexts where this level of control could be useful such as optimizing metabolic pathways.⁴⁸

CONCLUSIONS

In conclusion, we have demonstrated the utility of TX-TL reactions for rapidly prototyping and characterizing RNA circuitry. Each of the TX-TL experiments performed took a matter of hours to complete, and if DNA constructs are already available, several experiments can be completed with TX-TL in a single day. This is a significant speed up of the biological design-build-test cycle and demonstrates the power of TX-TL as a bridge to engineering fully functioning genetic circuitry that operates *in vivo*.

In addition to establishing TX-TL reactions as a way to characterize RNA circuitry, we have also used it to directly measure circuit response times with DNA spike experiments. With these experiments, we provide the first evidence that RNA circuits can propagate signals at time scales set by their degradation rates and showed that this leads to circuit response times on the order of 15–20 min. This is nearly 6 times faster than circuits constructed from stable proteins, which have response times set by the cell doubling time.^{27,46} Thus, RNA circuitry shows enormous speed up compared to protein circuitry, which could become even more important in designing circuitry that needs to operate in slowly dividing cells.

We also showed how TX-TL reactions could be used to systematically prototype components for larger circuit designs. Part of our success in constructing a SIM and verifying its function *in vivo* was the result of the ability to use TX-TL to characterize its individual subparts and confirm that tandem attenuators could be used to tune circuit response time. This led to the construction and characterization of the first RNA SIM in *vivo*, demonstrating that RNA circuitry can be used to create temporal programs of gene expression.

One goal of the TX-TL system is to serve as a molecular breadboard that would help guide circuit design *in vivo*.²⁸ While more and more studies are showing correlations between TX-TL and *in vivo* characterization^{29,30,33,34} it is important to investigate the differences relevant to each circuit component. Here, we have uncovered some important guidelines for using TX-TL to prototype RNA circuitry: (i) A key component that varies batch-to-batch are RNase activities, which affect

degradation times and RNA circuit performance. Therefore, it is important to design appropriate controls when adding multiple constructs to a reaction and screening of extract batches might be necessary. (ii) A single TX-TL reaction is resource limited; therefore, it is ideal to minimize total DNA input. This resource limitation may lead to differences in performance when compared to an *in vivo* environment, and one might consider supplementing or replenishing necessary buffer components. (iii) TX-TL serves as a great platform for optimizing individual circuit components and DNA concentration ratios. This characterization can be transferred to copy numbers and promoter design for *in vivo* constructs, though more work needs to be done to make this transfer process precise and predictable.

In addition, while we clearly observed functioning RNA cascades in TX-TL and in vivo, there are differences in the response times measured in these two contexts. In particular, the in vivo Att-1-RFP response (Figure 7B) was ~35 min faster than the TX-TL Att-1-SFGFP response (Supporting Information Figure S10), after correcting for protein maturation rate. Similarly, but to a lesser extent, the in vivo Att-1-Att-1-SFGFP response time (Figure 7B) was faster than the TX-TL response time (Figure 6C) by \sim 5 min. Since the response times are related to RNA degradation rates (eq 1), this suggests that the cellular degradation machinery is more abundant or faster, or processes such as cell doubling are contributing to additional degradation rate increases that allow in vivo circuitry to operate faster. More work needs to be done to quantitatively translate TX-TL circuit performance into predictions for in vivo circuit function.

Finally, we note that the initial portions of this work establishing TX-TL for characterizing RNA circuitry were performed at the inaugural Cold Spring Harbor summer course in Synthetic Biology in July–August 2013. During this intensive two week course, four of us (V.S., K.J.S., S.A.K., and C.P.F.), who had little to no prior experience in performing TX-TL reactions or engineering RNA circuitry, were able to confirm the functioning of the RNA cascade and prototype a version of the RNA SIM. The rapid time scale of TX-TL experiments for characterizing genetic circuitry and its simple experimental design²⁸ thus provide an ideal tool to teach the next generation of synthetic biologists in a cutting-edge research setting.

METHODS

Plasmid Construction and Purification. A table of all the plasmids used in this study can be found in Supporting Table S4, with key sequences found in Supporting Information Table S3. The pT181 attenuator and antisense plasmids, pT181 mutant attenuator and antisense plasmids, and the no-antisense control plasmid were constructs pAPA1272, pAPA1256, pAPA1273, pAPA1257, and pAPA1260, respectively, from Lucks et al.¹⁹ The theophylline pT181 mutant antisense plasmid was construct pAPA1306 from Qi et al.²⁰ The second level of the cascade was modified from construct pAPA1347 from Lucks et al.¹⁹ The double attenuator and SIM constructs were created using Golden Gate assembly.⁴⁹ Plasmids were purified using a Qiagen QIAfilter Plasmid Midi Kit (Catalog number: 12243) followed by isopropanol precipitation and eluted with double distilled water.

TX-TL Extract and Buffer Preparation. *Extract Preparation.* Cell extract and reaction buffer was prepared according to Shin and Noireaux³⁷ and Sun et al.²⁸ In brief, *E. coli* BL21 Rosetta cells were grown to an OD600 of 1.5, pelleted via centrifugation, and washed with a buffer at pH 7.7 containing Mg-glutamate, K-glutamate, Tris, and DTT. Lysis was performed via bead-beating, followed by centrifugation to remove beads and cell debris. The resulting supernatant was incubated at 37 °C for 80 min and then centrifuged, to remove endogenous nucleic acids. The supernatant was dialyzed against a buffer at pH 8.2, containing Mg-glutamate, K-glutamate, Tris, and DTT. The extract was then centrifuged, and the supernatant flash-frozen in liquid nitrogen and stored at -80 °C. The cell extract for Batch 1 had a protein concentration of 37 mg/mL, and its expression was optimized via the addition of 4 mM Mg and 20 mM K. For Batch 2: 29 mg/mL protein, 4 mM Mg, and 80 mM K.

Buffer Preparation. The reaction buffer was prepared according to Sun et al.,²⁸ and consists of an energy solution (HEPES pH 8 700 mM, ATP 21 mM, GTP 21 mM, CTP 12.6 mM, UTP 12.6 mM, tRNA 2.8 mg/mL, CoA 3.64 mM, NAD 4.62 mM, cAMP 10.5 mM, folinic acid 0.95 mM, spermidine 14 mM, and 3-PGA 420 mM) and amino acids (leucine, 5 mM, all other amino acids, 6 mM).

Extract and buffer were aliquoted in separate tubes (volume appropriate for seven reactions) and stored at -80 °C.

TX-TL Experiment. TX-TL buffer and extract tubes were thawed on ice for approximately 20 min. Separate reaction tubes were prepared with combinations of DNA representing a given circuit condition. Appropriate volumes of DNA, buffer, and extract were calculated using a custom spreadsheet developed by Sun et al.²⁸ Buffer and extract were mixed together and then added to each tube of DNA according to the previously published protocol.²⁸ Each TX-TL reaction mixture (10 μ L each) was transferred to a 384-well plate (Nunc 142761), covered with a plate seal (Nunc 232701), and placed on a Biotek SynergyH1m plate reader. We note that special care is needed when pipetting to avoid air bubbles, which can interfere with fluorescence measurements. Temperature was controlled at either 29 or 37 °C. SFGFP fluorescence was measured (485 nm excitation, 520 emission) every 1-5 min depending on the experiment. Spike experiments at 29 °C were paused after 25 min at which point solutions containing DNA, theophylline, or controls were added to the appropriate wells, and then placed back on the plate reader for fluorescence monitoring. Spike experiments at 37 °C were paused after 20 min. The spike time for each temperature was set to the start of constant protein production determined by preliminary experiments with our Att-1-SFGFP construct. In general, fluorescence trajectories were collected for 2 h, and each experiment lasting a total of 2-3 h.

Strains, Growth Media and *In Vivo* Gene Expression. All experiments were performed in *E. coli* strain TG1. Plasmid combinations were transformed into chemically competent *E. coli* TG1 cells, plated on Difco LB+Agar plates containing 100 μ g/mL carbenicillin, 34 μ g/mL chloramphenicol, and 100 μ g/mL kanamycin and incubated overnight at 37 °C. Plates were taken out of the incubator and left at room temperature for approximately 7 h. Four colonies were picked and separately inoculated into 300 μ L of LB containing carbenicillin, chloramphenicol, and kanamycin at the concentrations above in a 2 mL 96-well block (Costar 3960), and grown approximately 17 h overnight at 37 °C at 1,000 rpm in a Labnet Vortemp 56 benchtop shaker. This overnight culture (20 μ L) was then added to separate wells on a new block containing 930 and 980 μ L (1:50 dilution) of M9 minimal media (1xM9 minimal salts, 1 mM thiamine hydrochloride, 0.4% glycerol, 0.2% casamino acids, 2 mM MgSO₄, 0.1 mM CaCl₂) containing the selective antibiotics and grown for 4 h at the same conditions as the overnight culture. Two wells were used for the M9 growth to represent (+) and (-) theophylline conditions for the same colony. At this point, 50 μ L of a 40 mM theophylline solution was added to the wells containing 930 μ L of M9. The 96-well block was placed back on the shaker. Every 10 min for the next 90 min, 50 μ L of the cultures with and without theophylline was removed from the block and transferred to a 96-well plate (Costar 3631) containing 50 μ L of phosphate buffered saline (PBS). SFGFP fluorescence (485 nm excitation, 520 nm emission), mRFP fluorescence (560 nm excitation, 630 nm emission), and optical density (OD, 600 nm) were then measured at each time point using a Biotek Synergy H1m plate reader.

Response Time Calculation. TX-TL Experiments. To quantify the circuit response time, we calculated τ using data from multiple replicates in three individual experiments by first normalizing trajectories to the end point fluorescence of the L1+L2 condition to account for variation in fluorescence output between experiments (Supporting Information Figure S6). Normalized fluorescence distributions from all replicates, between each condition, were compared using Welch's t test at each time point to determine the time at which the L1+L2 and L1+L2+L3 data sets were statistically different from each other. The difference in average normalized fluorescence at this point was used to set a threshold, which was then used in each individual data set to determine the time at which each spiked trajectory differed from the average of the L1+L2 curves of that experiment (Supporting Information Figure S6). These times were then used to calculate reported τ with error.

In Vivo Experiment. To quantify the circuit response time of the SIM, we calculated τ using data from multiple replicates in three individual experiments by first normalizing trajectories to the average of the t = 0 fluorescence of each individual colony's with and without theophylline condition. Normalized fluorescence distributions from all replicates, between each condition, were compared using Welch's t test at each time point to determine the time at which the with and without theophylline data sets were statistically different from each other. The difference in average normalized fluorescence at this point was used to set a threshold that was then used in each individual data set to determine the time at which each spiked trajectory differed from its corresponding no-theophylline trajectory. These times were then used to calculate reported τ with error.

ASSOCIATED CONTENT

S Supporting Information

Tables S1–S4, Figures S1–S12, additional methods and Appendix 1. This material is available free of charge *via* the Internet at http://pubs.acs.org.

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Author Contributions

M.K.T., J.C., C.A.H., Z.Z.S., J.K., V.S., K.J.S., S.A.K., and C.P.F. performed the experiments. M.K.T., J.C., C.A.H., and J.B.L. designed the experiments and wrote the manuscript. V.S.,

K.J.S., S.A.K., C.P.F., Z.Z.S., J.K., V.N., and R.M. designed the experiments and edited the manuscript.

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Notes

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ABBREVIATIONS

Transcription-translation, TX-TL; single input module, SIM; super folder green fluorescent protein, SFGFP; red fluorescent protein, RFP; ribosome binding site, RBS

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