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# Quantitative characterization of translational riboregulators using an in vitro transcription-translation system

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Abstract

Riboregulators are short RNA sequences that, upon binding to a ligand, change their secondary structure and influence the expression rate of a downstream gene. They constitute an attractive alternative to transcription factors for building synthetic gene regulatory networks because they can be engineered de novo. However, riboregulators are generally designed in silico and tested in vivo, which provides little quantitative information about their performances, thus hindering the improvement of design algorithms. Here we show that a cell-free transcription-translation (TX-TL) system provides valuable information about the performances of in silico designed riboregulators. We first propose a simple model that provides a quantitative definition of the dynamic range of a riboregulator. We further characterize two types of translational riboregulators composed of a cis-repressed (cr) and a trans-activating (ta) strand. At the DNA level we demonstrate that high concentrations of taDNA poisoned the activator until total shut off, in agreement with our model, and that relative dynamic ranges of riboregulators determined in vitro are in agreement with published in vivo data. At the RNA level, we show that this approach provides a fast and simple way to measure dissociation constants of functional riboregulators, in contrast to standard mobility-shift assays. Our method opens the route for using cell-free TX-TL systems for the quantitative characterization of functional riboregulators in order to improve their design in silico.

## 22 Keywords

in vitro synthetic biology, RNA translational riboregulator, cell-free protein synthesis

During the early wave of synthetic biology, 1,2 known transcription factors were wired to their corresponding promoter sequences to control the expression of other transcription factors or effector proteins. While this approach has been very successful in engineering gene regulatory networks (GRNs)<sup>3</sup> with few nodes, the number of different elements in the majority of synthetic GRNs has stagnated at 5-6, 4,5 although a remarkable example contains 11 elements. Two arguments may explain this limit. First, protein-DNA interactions are very difficult to design, although very promising computational methods are arising, 7 and the engineer must thus choose well-known transcription factor-promoter pairs. Second, the expression of these transcription factors imposes a metabolic burden to the cells.<sup>8</sup> 

Implementing regulatory circuits at the RNA level may help solving these issues essen-tially because RNA-RNA interactions can be predicted from the sequence. 9-11 Moreover, in the case of transcriptional regulators, protein expression is not needed for regulation, which lowers the metabolic burden. 12 The principal component of an RNA-regulated GRN is the riboregulator: an RNA sequence in the 5' untranslated region (UTR) of a gene of interest that has an effect on its expression rate. Since they were first used in synthetic biology more than a decade ago, 13 several riboregulators have been designed and implemented in vivo, both in prokaryotic <sup>14–18</sup> and eukaryotic cells. <sup>19</sup> However, their design remains more difficult than expected and many implementations do not work in vivo. 12 One reason to this is that structure-prediction tools do not yet precisely capture the complexity involved in the folding of RNA species several hundreds of nucleotides long. Furthermore, in silico design relies on a structural model of riboregulation, which needs to be transformed into predictable features in order to generate optimized sequences. Another reason is that it is hard to control and tune the copy number of plasmids or genes in vivo and thus testing new parts in vivo 20,21 often provides information that is difficult to correlate with thermodynamic parameters used in silico. 

Including a phase of in vitro testing in the workflow of engineering riboregulators could potentially solve these problems. Structural characterization of riboregulators <sup>22,23</sup> helps as-

sessing the correctness of the designed structures and has been recently combined with functional information<sup>24</sup> but does not provide quantitative kinetic and thermodynamic data. To overcome these difficulties and accelerate the improvement of in silico designs, cell-free transcription-translation (TX-TL) platforms are an attractive tool for testing genetic regu-latory modules in synthetic biology. <sup>5,25–27</sup> Currently, there are two types of TX-TL systems available, cell-extract-based and PURE (Protein synthesis Using Recombinant Elements) systems. The first one is obtained by recovering the protein fraction from E. coli<sup>26</sup> while the PURE is just composed of individually-purified recombinant elements necessary for expression in vitro. 28,29 TX-TL in vitro testing can be used to qualitatively evaluate the performances of new designs in a faster manner<sup>5,27,30</sup> or to provide quantitative data such as thermodynamic and kinetic rates<sup>31</sup> that are of great value to improve in silico methods. Here we used a PURE TX-TL platform to illustrate the second approach. Briefly, the PURE system includes T7 RNA polymerase (RNAP), an energy-coupling module for NTP regeneration, transfer RNAs, ribosomes and translation initiation, elongation and release factors in a suitable buffer. 29,32 Its composition is well-controlled and it contains low levels of ribonucleases. The PURE system has already been used to characterize transcriptiontranslation dynamics 33,34 and GRNs 55 but has not yet been used to characterize riboregulators. Cell-extract TX-TL systems have been used to study transcriptional 30,31 but not translational riboregulators. To the best of our knowledge, TX-TL systems have so far investigated GRNs that mix both transcriptional and translational dynamics. In this work, we characterize the dynamics of translational riboregulators at the DNA and RNA level, which allows to independently study transcription and translation and clearly pinpoint de-sign shortcomings. The simplicity of the PURE system allows us to propose an analytical model of riboregulation that fits our data. The proposed model, together with the in vitro experiments, show that: i) TX-TL linearly amplifies the concentration of active RNA and quadratically amplifies the concentration of coding DNA, ii) we can provide a quantitative definition of the dynamic range of a riboregulator,  $\rho_{ON/OFF}$ , iii)  $\rho_{ON/OFF}$  is a bell-shaped

function of the concentration of regulatory DNA, iv) the relative values of  $\rho_{ON/OFF}$  measured in vitro coincide with published data in vivo for four riboregulators out of five and v) we can reliably use TX-TL to measure the dissociation constant,  $K_d$ , between the two RNA species. Although these results were facilitated by the simplicity of the PURE system, they could be extended to other cell-free TX-TL systems and possibly in vivo. These quantitative insights on translational riboregulators may also help improving in silico design routines.

## 84 Results and discussion

## 85 Translation rate vs. structure as the optimization goal for a ri-

Our study focuses on translational riboregulators, which are composed of two RNA strands

(Figure 1A). One of them, called cis repressed RNA, noted  $R_{cr}$ , about 800 nucleotides (nt)

long, codes for a gene but bears a hairpin in its 5'-untranslated region (5'-UTR) that pre-

### $_{86}$ boregulator

vents the ribosome to start translating the downstream gene. The other one, a small transactivating RNA, about 100 nt long, noted  $R_{ta}$ , hybridizes to the 5'-UTR of  $R_{cr}$ , opens up the hairpin and forms an active complex,  $R_{act}$ , increasing the translation rate. Ultimately, the riboregulator engineer is interested in controlling the rate of translation for  $R_{cr}$  and  $R_{act}$ , noted respectively  $r_{tl}^{cr}$  and  $r_{tl}^{act}$ , and seek the objective  $r_{tl}^{act} \gg r_{tl}^{cr} \approx 0$  for an activator (Figure 1B). For convenience we assign a species name to an RNA sequence, but one must bear in mind that a given RNA sequence, for instance  $R_{cr}$ , may fold in an ensemble of different structures  $\{R_{cr}^i\}$ , with different translation rates  $\{r_{tl}^{cr,i}\}$ . Current in silico design methods 11,36 compute the ensemble of secondary structures  $\{R_{cr}^i, R_{ta}^j, R_{act}^k\}$  that minimizes free energy. However, the structure-to-function relationship that associates an RNA conformation with its translation rate is hard to establish. Thus, a set of heuristic rules attributes low values of translation rates  $r_{tl}^{cr,j}$  to structures where the RBS or the start codon are buried in a hairpin (Figure 1), and high values of  $r_{tl}^{act,k}$ , to structures where 

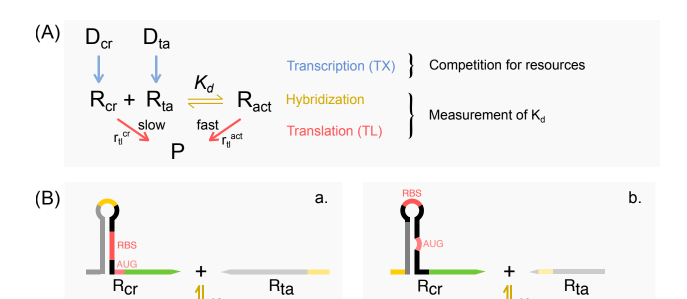


Figure 1: Principle of a translational riboregulator and of its characterization using a cell-free transcription-translation system (TX-TL). (A) Mechanism of transcription, riboregulation through RNA hybridization and translation used in this work. DNA sequences  $D_{cr}$  and  $D_{ta}$  are transcribed into a cis-repressed,  $R_{cr}$ , and a trans-activator,  $R_{ta}$ , RNA strands.  $R_{cr}$  may be slowly translated into protein P or hybridize with  $R_{ta}$  to form  $R_{act}$  that is translated more rapidly into P. Measuring the dynamics of fluorescence production by a fluorescent protein P provides information about resource competition when evaluating the system at the DNA level and quantitative values of dissociation constants  $K_d$  when RNA concentration is fixed. (B) Sketches of the two operation modes of translational riboregulators functioning as an activator. The 5'-UTR of  $R_{cr}$  RNA, forms a hairpin that hides either the ribosome binding site (RBS, a.) or the start codon (AUG, b.) away from the ribosome.  $R_{ta}$  hybridizes with  $R_{cr}$ , unwinding the hairpin and liberating the RBS and/or the AUG promoting translation.

Ract

Ract

these are accessible. However, these heuristic rules often fail. Moreover, minimizing the free energy of the RNA structures implies that the hypothesis of thermodynamic equilibrium holds, which is far from being true in vivo in the presence of co-transcriptional folding and RNA chaperones. <sup>37,38</sup> To shed light into this problem we measured translation and expression (transcription and translation) dynamics of recently in silico designed riboregulators <sup>20,21</sup> in the PURE system. <sup>28</sup>

## The TX-TL system linearly amplifies the concentration of RNA and quadratically amplifies the concentration of DNA

We first characterized the translation and expression reactions of the PURE system in the absence of riboregulation. To do so, we prepared by PCR a linear DNA fragment coding for a green fluorescent protein (GFP) with no upstream regulatory region, called cr<sup>-</sup>DNA. It is composed of a T7 RNAP promoter, a ribosome binding site, the GFP-coding sequence, and a T7 terminator. In addition, we prepared by in vitro transcription the corresponding messenger RNA, cr<sup>-</sup>RNA, from cr<sup>-</sup>DNA. We successively used cr<sup>-</sup>RNA and cr<sup>-</sup>DNA as the coding nucleic acid input of the TX-TL system. We varied the concentration of the input and we measured the fluorescence emitted by the GFP produced over time (Figure 2). Starting from cr<sup>-</sup>RNA, the translation module of the TX-TL system actively produced GFP during 2 hours. The translation kinetics displayed three different phases: during about 5 min no signal was discernable from the background level, then followed a phase of quasi-linear increase during 100 min, that slowed down until a plateau was reached (Figure 2A). In the range 0-80 nM of cr<sup>-</sup>RNA, both the final intensity and the maximum rate of fluorescence growth,  $v_{tl}^{max}$ , increased linearly with the initial quantity of coding RNA (Figure 2B). For higher concentrations there was a saturation: putting more RNA template did not increase significantly the final yield or the maximal production rate. When using cr<sup>-</sup>DNA as the initial input, the dynamics of the fluorescence intensity showed both common and contrasting features with the previous case (Figure 2C). Three phases were still observed: delay, growth

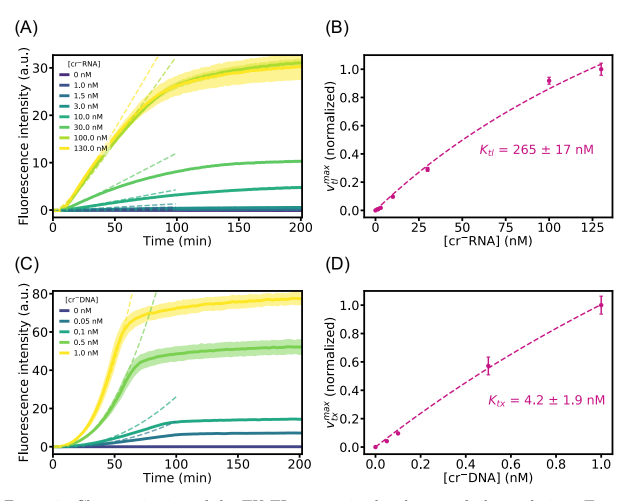


Figure 2: Characterization of the TX-TL system in the absence of riboregulation. Translation dynamics (A) and maximum fluorescence production rate (B) for increasing concentrations of an unregulated mRNA fragment coding for GFP. Expression (transcription and translation) dynamics (C) and maximum fluorescence production rate (D) for increasing concentrations of an unregulated linear DNA fragment coding for GFP. Solid lines (A,C) and disks (B,D) represent data, dotted lines are fits to the model. All experiments were performed in triplicate. Shading around the lines and error bars correspond to one standard deviation.

and a plateau. However, the delay observed before an increase of fluorescence was now of 15 min. Finally, the quantity of DNA required to saturate the maximum rate of fluorescence growth,  $v_{tx}^{max}$ , was almost two orders of magnitude lower than the quantity of RNA that saturated translation (Figure 2D).

We propose a simple analytical kinetic model that fits our data. To take into account the saturation of the production rates we assigned Michaelis-Menten kinetics to the transcription and the translation reactions. As a plausible source of the initial delay in the translation reaction, we included a first-order step of maturation of the non-fluorescent GFP protein, noted P, into the functional fluorescent protein  $P^*$ . This is in accordance to published maturation times.<sup>39</sup> We neglected DNA and RNA degradation and we did not take into consideration the depletion of resources because we analyzed our data between 0 and 50 min. For these reasons, our model did not reach a plateau in  $P^*$  concentration (Figure 2A,C). These approximations are valid as long as the RNA molecules do not deteriorate and the enzymatic resources, more specifically the ribosomes, are not depleted. We thus write the following mechanism 

$$D_{act} \stackrel{r_{tx}}{\to} D_{act} + R_{act} \tag{1}$$

$$R_{act} \xrightarrow{r_{tl}} R_{act} + P$$
 (2)

$$P \stackrel{r_m}{\to} P^* \tag{3}$$

where  $D_{act}$  and  $R_{act}$  are, respectively, cr<sup>-</sup>DNA and cr<sup>-</sup>RNA and  $r_{tx}$ ,  $r_{tl}$  and  $r_{m}$  are, respectively, the transcription, translation and maturation rates. With the aforementioned
hypotheses, this mechanism is associated with the rate equations

$$\frac{dR_{act}}{dt} = r_{tx} = \frac{k_{tx} \cdot D_{act}}{K_{tx} + D_{act}} \tag{4}$$

$$\frac{dP}{dt} = r_{tl} - r_m = \frac{k_{tl} \cdot R_{act}}{K_{tl} + R_{act}} - k_m \cdot P \tag{5}$$

$$\frac{dP^*}{dt} = r_m = k_m \cdot P \tag{6}$$

where  $k_x$  and  $K_x$  are, respectively, the rate and the Michaelis-Menten constants of reaction xand species concentrations are noted in italics. Equations (4-6) have exact solutions both for initial conditions corresponding to the translation  $(D_{act}(0) = 0, R_{act}(0) \neq 0)$  and expression experiments  $(D_{act}(0) \neq 0, R_{act}(0) = 0)$  (SI Section 3). For translation we obtain (SI Section 3.1)

$$P^*(t) = \frac{R_{act}(0)}{K_{tl} + R_{act}(0)} \frac{k_{tl}}{k_m} \left( e^{-k_m t} + k_m t - 1 \right)$$
 (7)

The term  $(e^{-k_m t} - 1)$ , due to protein maturation, makes the kink of the curves in Figure 2A at t = 10 min, while the linear term in time dominates for t = 20 - 50 min. Note that when the ribosome is not saturated,  $R_{act}(0) \ll K_{tl}$ , and for  $t \gg k_m^{-1}$  we can write

$$P^*(t) \approx \frac{k_{tl}}{K_{tl}} \cdot R_{act}(0) \tag{8}$$

explicitly showing that translation acts as a linear amplifier of the initial concentration of active RNA.

For expression, the exact solution is given in SI Section 3.3. Here we provide an approximated solution when  $R_{act}(t) \ll K_{tl}$  (SI Section 3.2),

$$P^*(t) \approx \frac{D_{act}(0)}{K_{tx} + D_{act}(0)} \frac{k_{tx}k_{tl}}{2K_{tl}} \left( t^2 - \frac{2}{k_m}t + \frac{2}{k_m^2} (1 - e^{-k_m t}) \right)$$
(9)

Again, if  $D_{act}(0) \ll K_{tx}$  and  $t \gg k_m^{-1}$ , expression quadratically amplifies  $D_{act}(0)$  into a fluorescence signal.

Considering that the fluorescence intensity is proportional to  $P^*$  we fitted (7) and (9) to the data in Figure 2. We obtained  $K_{tx}=4.2\pm1.9$  nM,  $K_{tl}=265\pm17$  nM and  $k_m=1.0$  $0.10 \pm 0.01 \ \mathrm{min^{-1}}$ , in fair agreement with previous measurements reporting  $K_{tx} = 4 - 9 \ \mathrm{nM}$ for T7 RNAP,  $^{33,34,40}$   $K_{tl}=66$  nM  $^{33}$  and  $k_m=0.2$  min $^{-1}$ .  $^{33,41}$  Note that although cell-extract TX-TL uses E. coli instead of T7 RNAP, the reported 41 value of  $K_{tx}$  is similar, 1-10 nM. In summary, the saturation of transcription by DNA occurs at a concentration two-orders of magnitude lower than the saturation of translation by RNA. Below saturation, the TX-TL system acts as a linear amplifier of the concentration of active RNA,  $R_{act}$ , and as a quadratic amplifier of  $D_{act}$  with a readout of intensity fluorescence. As a result we can use GFP fluorescence as a measure of the concentration of  $R_{act}$ . 

## Analytical model of translational riboregulation and quantitative definition of the dynamic range of a riboregulator

When riboregulators are used in vivo the DNA sequences  $D_{cr}$  and  $D_{ta}$ , respectively coding for the cis-repressed and trans-activator RNA  $R_{cr}$  and  $R_{ta}$ , can either be inserted in the chromosome, in the same plasmid or in two different plasmids. The performance of a riboregulator in vivo is assayed by fusing  $D_{cr}$  with a GFP and measuring the dynamic range, defined as

$$\rho_{ON/OFF} = \frac{\text{GFP fluorescence in the presence of D}_{ta}}{\text{GFP fluorescence in the absence of D}_{ta}}$$
(10)

In vivo it is common to use a two-plasmid strategy,  $^{20,42}$  trying to improve  $\rho_{ON/OFF}$  by inserting  $D_{ta}$  in a high-copy plasmid. The simplicity of in vitro TX-TL allows to provide a quantitative definition of  $\rho_{ON/OFF}$  and to test the effect of  $D_{ta}$  concentration on  $\rho_{ON/OFF}$ . We start by writing the simplest model of riboregulation dynamics from DNA that is consistent with the results of the previous section. Within the TX-TL system the two DNA molecules,  $D_{cr}$  and  $D_{ta}$ , are transcribed into the corresponding RNA strands, which associate into a

coding RNA,  $R_{act}$ . The production of P mainly comes from the translation of  $R_{act}$  but also may come from  $R_{cr}$ , when cis-repression is not very effective. We thus write the following mechanism,

$$D_{cr} \stackrel{r_{tx}^{cr}}{\rightarrow} D_{cr} + R_{cr} \quad , \qquad D_{ta} \stackrel{r_{tx}^{ta}}{\rightarrow} D_{ta} + R_{ta} \tag{11}$$

$$R_{cr} + R_{ta} \stackrel{K_d}{\rightleftharpoons} R_{act}$$
 (12)

$$R_{cr} \stackrel{r_{tl}^{cr}}{\to} R_{cr} + P \quad , \qquad R_{act} \stackrel{r_{tl}^{act}}{\to} R_{act} + P$$
 (13)

$$P \stackrel{r_m}{\to} P^* \tag{14}$$

We model reactions (11-14) with the following set of ODEs, that takes into account the competition for transcriptional resources,

$$\frac{dR_{cr}}{dt} = r_{tx}^{cr} = \frac{k_{tx}^{cr} \cdot D_{cr}}{K_{tx} + D_{cr} + D_{ta}}$$
(15)

$$\frac{dR_{ta}}{dt} = r_{tx}^{ta} = \frac{k_{tx}^{ta} \cdot D_{ta}}{K_{tx} + D_{cr} + D_{ta}}$$
(16)

$$R_{act} = \frac{R_{cr}R_{ta}}{K_d} \tag{17}$$

$$\frac{dP}{dt} = r_{tl}^{act} + r_{tl}^{cr} \approx \frac{k_{tl}^{act}}{K_{tl}} R_{act} + \frac{k_{tl}^{cr}}{K_{tl}} R_{cr}$$

$$\tag{18}$$

$$\frac{dP^*}{dt} = r_m = k_m \cdot P \tag{19}$$

where we have assumed that  $D_{cr}$  and  $D_{ta}$  may have different transcription rate constants  $k_{tx}^i$ .

We have also assumed, as previously, that transcription follows Michaelis-Menten dynamics, that translation can be considered a non-saturated Michaelis-Menten  $(R_{cr}, R_{act} \ll k_{tl}^{act})$  and further that the hybridization reaction (12) is fast compared with the others and thus can be considered at equilibrium. We have seen in the previous section that maturation reaction (14) introduces an additional term  $(e^{-k_m t} - 1)$  that vanishes when  $t \gg k_m^{-1} = 10$  min. To

facilitate subsequent calculations we will suppose  $t \gg k_m^{-1}$  and thus  $P = P^*$ .

We define  $\alpha = \frac{D_{ta}}{D_{cr}}$  and integrate (15-18) to obtain (SI Section 4)

$$P^*(\alpha) = \frac{k_{tl}^{act} \alpha}{3K_d k_{tl}^{act}} \frac{k_{tx}^{ta} k_{tx}^{cr}}{\left(1 + \alpha + \frac{K_{tx}}{D_{cr}}\right)^2} t^3 + \frac{k_{tl}^{cr}}{2k_{tl}^{act}} \frac{k_{tx}^{cr}}{1 + \alpha + \frac{K_{tx}}{D_{cr}}} t^2$$
(20)

where we have indicated explicitly that  $P^*$  is a function of  $\alpha$ . We can naturally define the dynamic range of the riboregulator as

$$\rho_{ON/OFF}^{th} = \frac{P^*(\alpha)}{P^*(\alpha = 0)} = \frac{1 + \frac{K_{tx}}{D_{cr}}}{\left(1 + \alpha + \frac{K_{tx}}{D_{cr}}\right)} \left[1 + \frac{2}{3} \frac{k_{tx}^{ta} k_{tl}^{act}}{K_d k_{tl}^{cr}} \frac{\alpha}{\left(1 + \alpha + \frac{K_{tx}}{D_{cr}}\right)} t\right]$$
(21)

where the superscript th indicates that this is an theoretical quantity defined in the framework of model (15-18). This equation reveals three important points. Firstly,  $\rho_{ON/OFF}^{th}$ depends linearly on time and thus it is difficult to compare  $\rho_{ON/OFF}$  between two experi-ments, in vivo or in vitro, if they have not been calculated at the same time. This linear dependence comes from the fact that protein production from the riboregulator is cubic in time, while the leak production from  $D_{cr}$  alone is quadratic (20). In vivo one may expect that  $\rho_{ON/OFF}^{th}$  reaches a plateau due to degradation. However a linear increase of  $\rho_{ON/OFF}$ was observed over 4 h for riboregulator 7 in reference 20. Secondly,  $\rho_{ON/OFF}^{th}$  is proportional to the aggregate factor  $\beta = \frac{k_{tx}^{ta}k_{tl}^{act}}{K_dk_{tl}^{cr}}$ . Thus,  $\rho_{ON/OFF}^{th}$  is proportional to the transcription rate constant  $k_{tx}^{ta}$ , and thus will differ between different RNAPs and promoters, it is also propor-tional to the ratio between the translation rate constant of the active and the inactive state, which is intuitive, and it is inversely proportional to the equilibrium constant of dissociation between  $R_{cr}$  and  $R_{ta}$ . Finally,  $\rho_{ON/OFF}^{th}$  is strongly and non-trivially dependent on the concentration of  $D_{cr}$  and  $D_{ta}$  (through  $\alpha = D_{ta}/D_{cr}$ ). This is, to the best of our knowledge, the first rigorous definition of the dynamic range of a translational riboregulator. The maximum of  $\rho_{ON/OFF}^{th}$  is obtained for (SI Section 4.2)

$$D_{ta} = D_{cr} + K_{tx} (22)$$

## The dynamic range of a riboregulator strongly depends on the concentration of $D_{ta}$

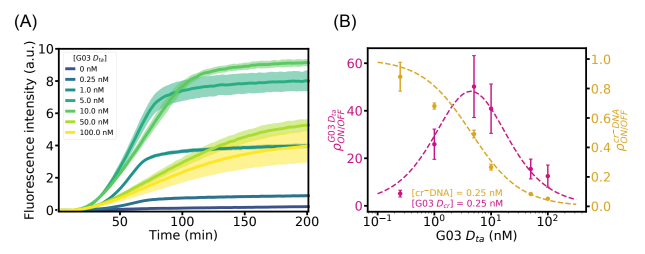


Figure 3: The dynamic range of a riboregulator strongly depends on the concentration of  $D_{ta}$ . (A) Fluorescence intensity vs. time for the in vitro expression of 0.25 nM of  $D_{cr}$  DNA, coding for GFP, with increasing concentrations of  $D_{ta}$  DNA, for riboregulator G03. (B) Dynamic range  $\rho_{ON/OFF}$  at time 75 min for a  $D_{cr}$  with, G03 (pink disks, left axis), or without, cr<sup>-</sup> (yellow disks, right axis), cis regulatory region as a function of the concentration of  $D_{ta}$  from riboregulator G03. The dashed lines correspond to a fit to equation (21) with a single free parameter  $\beta = (k_{tx}^{ta}k_{tt}^{act})/(K_dk_{tt}^{cr})$  (pink) and to SI equation (30) without fit (yellow), both using  $K_{tx} = 4.2$  nM as measured in Figure 2D. All experiments were performed in triplicate. Shading around the lines and error bars correspond to one standard deviation.

To test the model's prediction that  $\rho_{ON/OFF}$  strongly depends on  $\alpha$  and thus on  $D_{ta}$ , we titrated riboregulator G03 (Table S1) by keeping  $D_{cr} = 0.25$  nM constant, varying  $D_{ta}$ in the range 0-100 nM and recording GFP fluorescence over time (Figure 3). Increasing  $D_{ta}$  in the range 0-5 nM resulted in an increased fluorescence signal. However, for  $D_{ta}$ 5 nM the fluorescence signal dramatically decreased until reaching 10% of the maximum production rate at  $D_{ta} = 100$  nM. Equation (21) fits the data with a single free parameter  $\beta = 286$  (Figure 3B, pink line), indicating that the bell-like shape of  $\rho_{ON/OFF}$  arises from the competition of  $D_{ta}$  and  $D_{cr}$  for transcriptional resources. To further test this interpretation we titrated cr<sup>-</sup>DNA, which lacks the cis-regulatory region, with the  $D_{ta}$  of riboregulator G03. The data were quantitiatively predicted by equation (21) taking the limit  $K_d \to \infty$ (SI equation (30)) without fitting parameters. Importantly, the addition of non-transcribing DNA had little effect on the expression dynamics (SI Figure S3).

The observation that an increase in non-coding DNA concentration reduces protein ex-pression in TX-TL systems has already been reported <sup>25,30,43,44</sup> and it has been modeled <sup>45</sup> in the absence of riboregulation, although a quantitative comparison between the model and the data has not been reported. Our model (15-18) explicitly takes into account the competition between the two DNA substrates  $D_{ta}$  and  $D_{cr}$  and the predicted dependence of  $\rho_{ON/OFF}^{th}$  on  $D_{ta}$  is in agreement with the data. Hu et al. recently proposed a kinetic model for transcriptional riboregulators<sup>31</sup> and compared their model with in vitro TX-TL experiments. Our models are of the same type in the sense that they describe the kinetics with a set of ODEs at the level of concentrations. In contrast, the model of Hu et al has significantly more parameters than ours, 13 instead of 5, and takes into account the degra-dation of both RNA and protein. As a result, the authors cannot provide analytical results that clearly pinpoint the important parameters to design functional riboregulators, such as equation (21). 

Saturation of transcriptional resources is particularly important in the context of riboregulators, where the non-coding DNA produces a regulatory RNA that has an important effect
in the GRN. To the best of our knowledge, the bell-like curve in Figure 3B has not been
reported before. The similar value of  $K_{tx}$  for T7 and E. coli RNAP, together with previous
observations of transcriptional saturation in E. coli-based TX-TL systems, suggests that
this behavior is not due to a particular property of the T7 RNAP. Our model and in vitro
results thus predict that inserting  $D_{ta}$  in a high-copy plasmid will decrease the performance
of the riboregulator activator and suggests a trade-off between resource competition and the
over-expression of antisense RNAs. This prediction shall be tested in a future work.

## Comparison of the dynamic range of riboregulators in vivo and in vitro

To evaluate how dynamic ranges determined in vitro compared with in vivo measurements, we investigated five riboregulators of two different types, two loop-mediated 21 and three toehold-mediated <sup>20</sup> (Table S1). In the former, the RBS is buried inside the hairpin and the  $R_{ta}$  binds first to the loop on the hairpin. In the later, the start codon is protected by the hairpin and the  $R_{ta}$  binds to a toehold sequence on the 5' side of the hairpin (Figure 1B). We performed in vitro GFP expression experiments at 1 nM  $D_{cr}$  in the presence and in the absence of 5 nM of the corresponding  $D_{ta}$ . We chose  $D_{cr}$  and  $D_{ta}$  that verified (22) to determine the maximum dynamic range. In vivo, the dynamic range is generally defined without subtracting the autofluorescence of the cells. <sup>20,21</sup> In the previous section, to compare with  $\rho^{th}_{ON/OFF}$ , we computed the experimental  $\rho_{ON/OFF}$  by dividing fluorescence signals that had been subtracted from the autofluorescence of the PURE system. In this section, to compare with in vivo measurements, we computed  $\rho'_{ON/OFF}$ , where the prime indicates that autofluorescence was not subtracted.

Table 1: Comparison of the performance of five riboregulators in vivo and in vitro. Dynamic range calculated without subtracting the autofluorescence,  $\rho'_{ON/OFF}$ , in vivo and in TX-TL, ON and OFF raw fluorescence signals,  $I_{ON}$  and  $I_{OFF}$ , in TX-TL, and ratio of  $\rho'_{ON/OFF}$  in vivo relative to in vitro. In vivo data were extracted from ref. 20 for GXX and from ref. 21 for RAJXX. TX-TL data were measured at  $D_{cr} = 1$  and  $D_{ta} = 5$  nM at t = 75 min. The typical value of autofluorescence was  $0.07 \pm 0.01$  a.u.. Error bars correspond to one standard deviation of a triplicate experiment.

	in vivo	TX-TL	TX-TL	TX-TL	ratio $\rho'_{ON/OFF}$
Device	$ ho'_{ON/OFF}$	$ ho'_{ON/OFF}$	$I_{ON}$ (a.u.)	$I_{OFF}$ (a.u.)	in vivo/ TX-TL
G01	$290 \pm 20$	$37 \pm 10$	$116 \pm 25$	$3.1 \pm 0.5$	8±2
G03	$260 \pm 30$	$26 \pm 6$	$81 \pm 10$	$3.1 \pm 0.6$	$10\pm 3$
G80L18	$500\pm150$	$23 \pm 3$	$93 \pm 8$	$4.1 \pm 0.4$	$22 \pm 8$
RAJ11	$11 \pm 2$	$1.9 \pm 0.3$	$2.1 \pm 0.3$	$1.1 \pm 0.1$	$6\pm1$
RAJ12	$8 \pm 1$	$1.2\pm0.1$	$1.2 \pm 0.2$	$1.1\pm0.1$	$7\pm1$

The agreement between the values obtained in vivo and in vitro is remarkable. Of course, the absolute values of  $\rho'_{ON/OFF}$  in vivo and in vitro are different, which is expected because

 $\rho_{ON/OFF}^{th}$  is proportional to time and in vivo and in vitro data were obtained at different times (Table 1). In contrast, the relative order of  $\rho'_{ON/OFF}$  is similar in vivo and in vitro. Moreover the ratio between the two is constant for all riboregulators except for G80L18 that is twice more active in vivo, indicating that TX-TL experiments predict well  $\rho'_{ON/OFF}$ in vivo. The measured value of  $\rho'_{ON/OFF}$  for RAJ12 is close to unity. However, increasing the DNA concentrations to  $D_{cr} = D_{ta} = 50$  nM, which increases protein production (20) while respecting (22), demonstrated that RAJ12 was indeed functional and we obtained  $\rho'_{ON/OFF} = 7$  at these concentrations. Comparing the values of  $I_{OFF}$  shows that RAJXX leaked significantly less than GXX while G80L18 leaked slightly more than G01 and G03. Finally, our experiments show that the remarkable  $\rho'_{ON/OFF}$  values of GXX devices come from their high  $I_{ON}$ , and thus a very active ON state. We thus conclude that in vitro TX-TL provides values of  $\rho'_{ON/OFF}$  that correlate well with in vivo measurements, in agreement with previous reports comparing protein expression in vivo and in vitro. <sup>30,45,46</sup> 

## Translation from RNA characterizes the reaction between the cisrepressed and the trans-activator RNA

The regulatory step of translational riboregulators takes place when the two RNA fragments,  $R_{cr}$  and  $R_{ta}$ , hybridize and thereby change the accessibility of the ribosome to a site needed for initiating translation (RBS or AUG). The core of the riboregulation process can thus be described with reactions (12) and (13), where the first one involves the hybridization of  $R_{cr}$  with  $R_{ta}$  to form an active RNA complex,  $R_{act}$ , that can be translated, and the second being the translation of  $R_{act}$  into protein P. We have seen that the thermodynamics of the first reaction play an important role in  $\rho_{ON/OFF}^{th}$  through  $K_d$  (21). However  $K_d$  is not straightforward to determine. One possibility is to use an electrophoretic mobility shift assay in a polyacrylamide gel. Another way uses the property of a reverse transcriptase to terminate on stable RNA duplexes.  $^{13}$  In both cases these assays characterize the species  $R_{act}$ for being a duplex RNA but they are not sensitive to its translational activity. Here, instead,

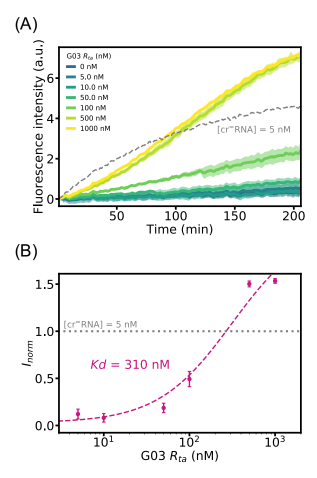


Figure 4: Titration of a riboregulator at the RNA level measures the dissociation constant of the riboregulator complex. GFP fluorescence produced over time (A) and normalized maximum fluorescence (B) for different trans-activator concentrations,  $R_{ta}$  for riboregulator G03. As a control, panel A shows the fluorescence intensity produced by the translation of 5 nM of an unregulated cr<sup>-</sup>RNA (grey dashes). In (B) disks correspond to experimental data and the dashed line is a fit of (23) to the data. All experiments were performed in triplicate. Shading around the lines and error bars correspond to one standard deviation.

we probed the equilibrium concentration of  $R_{act}$  that is active for translation. Our method is thus more meaningful to evaluate the design performances of a riboregulator.

To characterize reaction (12) we in vitro transcribed the five riboregulators described previously (Figure S1). We studied their translation dynamics by titrating 5 nM  $R_{cr}$  with increasing concentrations of its corresponding  $R_{ta}$  in the range 0 – 1000 nM (Figure 4 and SI Figure S4). Because translation linearly amplifies  $R_{act}$  (Figure 2B and (8)), measuring the GFP intensity at a given time is directly proportional to the concentration of  $R_{act}$  that is translationally active. We thus plotted the normalized GFP fluorescence at 200 min as a function of the log of  $R_{ta}$  concentration. For a bimolecular equilibrium such as (12) one expects these plots to be described by

$$I_{norm} \sim \bar{R}_{act} = \frac{1}{2} R_{cr}^{0} \left( \frac{K_d + R_{cr}^{0} + R_{ta}^{0}}{R_{cr}^{0}} - \sqrt{\left( \frac{K_d + R_{cr}^{0} + R_{ta}^{0}}{R_{cr}^{0}} \right)^2 - 4 \frac{R_{ta}^{0}}{R_{cr}^{0}}} \right)$$
(23)

where  $\bar{R}_{act}$  is the equilibrium concentration of  $R_{act}$  and superscript 0 indicates initial concentrations (SI Section 6). Our experimental data followed this trend (Figures 4 and S4).

We thus fitted (23) to the data and found dissociation equilibrium constants in the range 10 - 2000 nM (Table 2), in agreement with  $K_d$  values of the order of 100 nM that have already been reported for loop-mediated activators. Values of  $K_d$  obtained from different batches of PURE were within 50 % (Figure S5).

In the case of G01, however, after a normal sigmoidal increase of  $I_{norm}$  vs.  $R_{ta}$ ,  $I_{norm}$  decreased for  $R_{ta} > 200$  nM (Figure S4). To evaluate why in this particular case high  $R_{ta}$  inhibited translation, we performed a control experiment where a well-behaved regulator, G80L18, activated with 50 nM of its corresponding  $R_{ta}$ , was titrated with increasing concentrations of  $R_{ta}$ -G01 (Figure 5). We observed again that very high concentrations of  $R_{ta}$ -G01 significantly reduced the final GFP concentration. In contrast, similarly high concentrations of  $R_{ta}$ -RAJ11 did not have a significant effect in translation. We thus concluded that  $R_{ta}$ -G01 poisoned the translation machinery, which could occur by nonspecific binding to other RNA

components, including tRNAs, ribosomes or mRNA, with about 1  $\mu$ M affinity. However, a sequence alignment between riboregulator's sequence and tRNA and rRNA from E.~coli did not show significant differences among riboregulators (Figure S6).

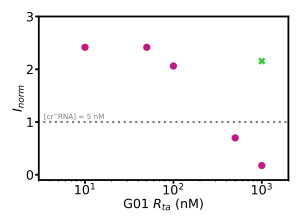


Figure 5: Titration of activated riboregulator G80L18 ( $R_{cr} = 5 \text{ nM } R_{ta} = 50 \text{ nM}$ ) with increasing concentrations of  $R_{ta}$  from riboregulators G01 (disks) and RAJ11 (green cross).

To assess the performance of our method for measuring  $K_d$ , we independently measured it with a standard mobility-shift assay performed with capillary gel electrophoresis. We used the same purified  $R_{cr}$  and  $R_{ta}$  that we mixed together at 37°C in a buffer with identical salt composition than the TX-TL system during 10 min before performing the electrophoresis assay.  $R_{cr}$  concentration was 8.3 nM and the  $R_{ta}$  concentration was ranging from 0 to 200 nM. Figures 6 and S7 show the electropherograms for riboregulator G03, where a peak in intensity at a given time point corresponds to an RNA structure. We detected three main peaks corresponding to  $R_{ta}$  at 28 s (Figure S7) and  $R_{cr}$  and  $R_{act}$  complex between 37 and 40 s (Figure 6A). Interestingly, species  $R_{cr}$  and  $R_{act}$  yielded well-resolved peaks for toehold-mediated but not for loop-mediated riboregulators (Figure S8), which suggests a structural difference between the two. As a result this method only provided  $K_d$  for some but not all of the tested riboregulators, in contrast with the TX-TL method. The values obtained were of the same order of magnitude of those obtained by TX-TL. However, mobility-shift assay yielded  $K_d$  in a narrower range of 100 – 250 nM, while TX-TL was able to better discriminate  $K_d$  for the same species and provided values in the range 15 – 2200 nM (Table 2).

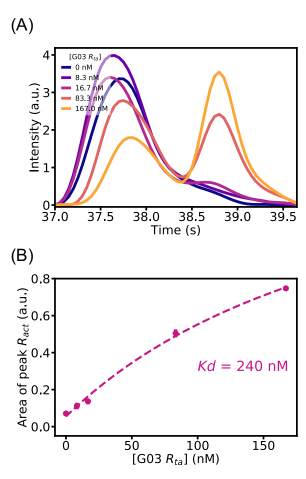


Figure 6: Titration of translational riboregulator G03 by mobility-shift capillary electrophoresis. (A) Corrected electropherograms vs. elution time and (B) peak area for different concentrations of  $R_{ta}$ . Experiments were performed in triplicate. Error bars correspond to one standard deviation. Dashed line is a fit of (23) to the data.

Table 2: Dissociation constants  $K_d$  at 37°C for the studied riboregulator devices.  $K_d$  was measured using the cell-free translation method (txtl) and the mobility-shift method (ms). N.M. indicates that the electropherogram showed ill-defined peaks from which  $K_d$  could not be extracted. Error bars correspond to one standard deviation of the fit. Values for G03 and RAJ11 were fitted to data in triplicate.

Device	$K_d^{txtl}$ (nM)	$K_d^{ms}$ (nM)
G01	$46 \pm 56$	$180 \pm 20$
G03	$310 \pm 154$	$240 \pm 110$
G80L18	$31 \pm 19$	$110 \pm 90$
RAJ11	$15 \pm 14$	N.M.
RAJ12	$2220\pm950$	N.M.

## **Conclusion**

We have demonstrated that in vitro transcription-translation (TX-TL) systems are an at-tractive platform to quantitatively characterize translational riboregulators. To do so we have taken advantage of the ribosome as a molecular machine that not only recognizes RNA complexes that are translationally active but also measures their concentration. The sim-plicity of the TX-TL system allowed us to propose an analytical expression for the dynamic range of a riboregulator. In quantitative agreement with this model we have shown that increasing the DNA concentration of the trans-activating species first promotes and later in-hibits expression. This result suggests that inserting trans-activating elements in high-copy plasmids in vivo could limit the efficiency of translational activators, a prediction that shall be tested in future work. Furthermore, relative dynamic ranges measured in vitro were in agreement with those reported in vivo for four out of five measured riboregulators. Finally, by titrating the cis-repressed gene with the trans-activating species at the RNA level we could determine dissociation constants,  $K_d$ , for the RNA hybridization reaction in a very simple manner. In particular, we could obtain  $K_d$ 's for riboregulators that could not be resolved by mobility-shift assays. Our method thus provides a simple and rapid way for the quantitative characterization of riboregulators. 

Combined with other biomolecular techniques such as molecular beacons<sup>34</sup> and automatedbased designs, <sup>47</sup> cell-free transcription-translation systems are becoming essential for a wide
brand of applications. They allow to verify theoretical predictions on both RNA structures
and behaviour of large scale regulatory networks. Their versality is a real asset for conceiving
new synthetic biological features <sup>48</sup> and creating innovative biomolecular tools. <sup>49</sup> The use of
an in vitro step in the design and elaboration of complex synthetic regulatory networks will
maximise the chance of expected in-vivo performances.

## Methods

## $_{\scriptscriptstyle 61}$ DNA and RNA preparations

DNA templates were prepared by PCR amplification of plasmids encoding for the RNA translational regulators, followed by affinity column purification using Monarch PCR Pu-rification Kit (New England BioLabs) or PureLink PCR Purification Kit (Thermo Fisher Scientific). Primers used for PCR amplification contained a T7 promoter or a T7 terminator (Biomers). RNA templates were prepared by in vitro transcription followed by purification using MEGAclear Transcription Clean-Up Kit (Ambion). The DNA and RNA integrity was determined by a 1.5% agarose gel (Figure S1) and the concentrations were determined by absorbance at 260 nm using a NanoDrop 2000 UV-Vis spectrophotometer. The sequences of the riboregulator domains (Table S1), of the PCR primers (Table S2) and of the plasmids are compiled in the SI. 

## Preparation of the PURE TX-TL system

The PURE TX-TL system was prepared according to reference 50 to reach the following composition: 1 units/ $\mu$ L of RNase inhibitor Murine (New England Biolabs), 50 mM Hepes-KOH pH 7.6, 13 mM magnesium acetate, 100 mM potassium glutamate, 2 mM spermidine, 1 mM dithiothreitol (DTT), 2 mM of each ATP and GTP, 1 mM of each CTP and UTP, 20 mM creatine phosphate, 0.3 mM 20 amino acids, 56 A260/ml tRNA mix (Roche), 10 μg/mL 10-formyl-5, 6, 7, 8-tetrahydrofolic acid, 0.1 mM each of amino acids, and factor mix. The factor mix contained 1.2  $\mu$ M ribosome, 10  $\mu$ g/ml IF1, 40  $\mu$ g/ml IF2, 10  $\mu$ g/ml IF3, 50  $\mu$ g/ml EF-G, 100  $\mu$ g/ml EF-Tu, 50  $\mu$ g/ml EF-Ts, 10  $\mu$ g/ml RF1, 10  $\mu$ g/ml RF2,  $10 \mu \text{g/ml RF3}$ ,  $10 \mu \text{g/ml RRF}$ , 600-6000 U/ml of each ARS and MTF  $4.0 \mu \text{g/ml}$  creatine kinase (Roche), 3.0  $\mu$ g/ml myokinase (Sigma), 1.1  $\mu$ g/ml nucleoside-diphosphate kinase, 1.0 U/ml pyrophosphatase (Sigma), and 10  $\mu$ g/ml of T7 RNAP.

### Fluorescence measurements in real-time PCR machine

Rotor-GeneQ real-time PCR (Qiagen) was used to record fluorescence from GFP expression (excitation  $470\pm10$  nm, emission  $510\pm5$  nm) in an 8 or 15  $\mu$ L volume. The temperature was set to  $37^{\circ}$ C and fluorescence recorded every minute for at least 3 h. In some experiments (Figure 5) we used PURExpress in vitro protein synthesis kit (NEB).

## 389 Data processing

Data were processed using in-house Python routines. For each condition of template —DNA or RNA— concentration, fluorescence intensity plots were shifted to the origin by removing the mean value of the three first minutes and by subtracting the fluorescence due to the PURE TX-TL system without any template.  $I_{norm}$  was computed by dividing this corrected fluorescence by the final intensity of the cr<sup>-</sup>RNA control. Corrected data were filtered using a Savitzky–Golay filter (window length: 21, polynomial order: 3) to remove residual noise before being derived to compute  $v^{max}$ .

## $_{\scriptscriptstyle 997}$ Electrophoretic mobility shift assays

Electrophoretic mobility shift assays were performed with a 2100 Bioanalyzer System (Agilent Technologies) and an RNA Nano chip Kit. Samples were prepared by mixing RNA
strands in 50 mM Hepes-KOH pH 7.6, 13 mM magnesium acetate, 100 mM potassium glutamate, 2 mM spermidine, 1 mM DTT and nuclase free water. They were incubated at 37°C
for 10 min before being loaded into the electrophoresis chip. Electropherograms were manually aligned along the time axis. Affine curves corresponding to the backgrounds of zones of
interest were subtracted. Areas under peaks were determined by numerical integration and
were normalized using an RNA marker provided in Agilent's kit.

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## Supporting Information Available

- Sequences for the RNA regulators, the PCR primers and the plasmids used, additional figures S1–S8, tables S1–S2, the detailed solution of translation and expression kinetics in the absence and in the presence of regulation, derivation of the equation used for Kd determination, supplementary data on RNA titration and details on the sequence alignment.
- References
- 1. Gardner, T. S., Cantor, C. R., and Collins, J. J. (2000) Construction of a genetic toggle switch in Escherichia coli. Nature 403, 339–342, 10.1038/35002131.
- 2. Elowitz, M. B., and Leibler, S. (2000) A synthetic oscillatory network of transcriptional regulators. *Nature* 403, 335–338.
- 3. Cameron, D. E., Bashor, C. J., and Collins, J. J. (2014) A brief history of synthetic biology. Nat Rev Micro 12, 381–390.
- 4. Purnick, P. E. M., and Weiss, R. (2009) The second wave of synthetic biology: from modules to systems. Nat Rev Mol Cell Biol 10, 410-422, 10.1038/nrm2698.
- 5. Niederholtmeyer, H., Sun, Z. Z., Hori, Y., Yeung, E., Verpoorte, A., Murray, R. M., and Maerkl, S. J. (2015) Rapid cell-free forward engineering of novel genetic ring oscillators. Elife 4.

- 6. Moon, T. S., Lou, C., Tamsir, A., Stanton, B. C., and Voigt, C. A. (2012) Genetic programs constructed from layered logic gates in single cells. *Nature* 491, 249.
- 7. Das, R., and Baker, D. (2008) Macromolecular Modeling with Rosetta. *Annual Review of Biochemistry* 77, 363–382.
- 8. Schaerli, Y., Munteanu, A., Gili, M., Cotterell, J., Sharpe, J., and Isalan, M. (2014) A unified design space of synthetic stripe-forming networks. *Nat Commun* 5, 10.1038/ncomms5905.
- 9. Zuker, M. (2003) Mfold web server for nucleic acid folding and hybridization prediction.

  Nucleic Acids Res. 31, 3406–3415, 10.1093/nar/gkg595.
- 10. Lorenz, R., Bernhart, S. H., Höner zu Siederdissen, C., Tafer, H., Flamm, C.,
   Stadler, P. F., and Hofacker, I. L. (2011) ViennaRNA Package 2.0. Algorithms for Molecular Biology 6, 1–14.
- 11. Zadeh, J. N., Steenberg, C. D., Bois, J. S., Wolfe, B. R., Pierce, M. B., Khan, A. R.,
   Dirks, R. M., and Pierce, N. A. (2011) NUPACK: Analysis and design of nucleic acid
   systems. Journal of Computational Chemistry 32, 170–173.
- Chappell, J., Watters, K. E., Takahashi, M. K., and Lucks, J. B. (2015) A renaissance in
   RNA synthetic biology: new mechanisms, applications and tools for the future. Current
   Opinion in Chemical Biology 28, 47–56.
- 13. Isaacs, F. J., Dwyer, D. J., Ding, C., Pervouchine, D. D., Cantor, C. R., and Collins, J. J.
   (2004) Engineered riboregulators enable post-transcriptional control of gene expression.
   Nat Biotech 22, 841–847, 10.1038/nbt986.
- 14. Win, M. N., and Smolke, C. D. (2007) A modular and extensible RNA-based generegulatory platform for engineering cellular function. *Proc. Natl. Acad. Sci. U. S. A.* 104, 14283–14288.

- 15. Salis, H. M., Mirsky, E. A., and Voigt, C. A. (2009) Automated design of synthetic ribosome binding sites to control protein expression. *Nat. Biotechnol.* 27, 946–950.
- 454 16. Qi, L., Larson, M. H., Gilbert, L. A., Doudna, J. A., Weissman, J. S., Arkin, A. P., and
   Lim, W. A. (2013) Repurposing CRISPR as an RNA-Guided Platform for Sequence Specific Control of Gene Expression. Cell 152, 1173–1183.
- 17. Chang, A. L., Wolf, J. J., and Smolke, C. D. (2012) Synthetic RNA switches as a tool
   for temporal and spatial control over gene expression. Current opinion in biotechnology
   23, 679–688.
- 18. Chappell, J., Takahashi, M. K., and Lucks, J. B. (2015) Creating small transcription
   activating RNAs. Nat. Chem. Biol. 11, 214–20.
- 19. Rinaudo, K., Bleris, L., Maddamsetti, R., Subramanian, S., Weiss, R., and Benenson, Y.
   (2007) A universal RNAi-based logic evaluator that operates in mammalian cells. Nat.
   Biotechnol. 25, 795–801.
- <sup>465</sup> 20. Green, A. A., Silver, P. A., Collins, J. J., and Yin, P. (2014) Toehold switches: de-novodesigned regulators of gene expression. *Cell* 159, 925–39.
- 21. Rodrigo, G., Landrain, T. E., and Jaramillo, A. (2012) De novo automated design of
   small RNA circuits for engineering synthetic riboregulation in living cells. *Proceedings* of the National Academy of Sciences 109, 15271–15276.
- 22. Greenleaf, W. J., Frieda, K. L., Foster, D. A. N., Woodside, M. T., and Block, S. M.
  (2008) Direct Observation of Hierarchical Folding in Single Riboswitch Aptamers. Science (80-.). 319, 630–633.
- 23. Ge, P., and Zhang, S. (2015) Computational analysis of RNA structures with chemical probing data. *Methods* 79, 60–66.

- <sup>475</sup> 24. Watters, K. E., Abbott, T. R., and Lucks, J. B. (2016) Simultaneous characterization
- of cellular RNA structure and function with in-cell SHAPE-Seq. Nucleic Acids Research
- 44, e12–e12, 10.1093/nar/gkv879.
- <sup>478</sup> 25. Noireaux, V., Bar-Ziv, R., and Libchaber, A. (2003) Principles of cell-free genetic circuit
- assembly. Proc. Natl. Acad. Sci. U. S. A. 100, 12672–7.
- 26. Garamella, J., Marshall, R., Rustad, M., and Noireaux, V. (2016) The All E. coli TX-TL
- Toolbox 2.0: A Platform for Cell-Free Synthetic Biology. ACS Synth. Biol. 5, 344–55.
- <sup>482</sup> 27. Sun, Z. Z., Yeung, E., Hayes, C. A., Noireaux, V., and Murray, R. M. (2014) Linear
- DNA for Rapid Prototyping of Synthetic Biological Circuits in an Escherichia coli Based
- TX-TL Cell-Free System. ACS Synth. Biol. 3, 387–397.
- 28. Shimizu, Y., Inoue, A., Tomari, Y., Suzuki, T., Yokogawa, T., Nishikawa, K., and
- Ueda, T. (2001) Cell-free translation reconstituted with purified components. Nat Biotech
- , 751–755.
- 29. Shimizu, Y., Kanamori, T., and Ueda, T. (2005) Protein synthesis by pure translation
- systems. Methods 36, 299–304.
- 490 30. Takahashi, M. K., Chappell, J., Hayes, C. A., Sun, Z. Z., Kim, J., Singhal, V.,
- Spring, K. J., Al-Khabouri, S., Fall, C. P., Noireaux, V., Murray, R. M., and Lucks, J. B.
- (2015) Rapidly Characterizing the Fast Dynamics of RNA Genetic Circuitry with Cell-
- Free Transcription-Translation (TX-TL) Systems. ACS Synthetic Biology 4, 503–515.
- 494 31. Hu, C. Y., Varner, J. D., and Lucks, J. B. (2015) Generating Effective Models and
- Parameters for RNA Genetic Circuits. ACS Synth. Biol. 4, 914–926.
- 496 32. Whittaker, J. W. (2013) Cell-free protein synthesis: The state of the art. Biotechnol.
- 497 Lett. 35, 143–152.

- 33. Stogbauer, T., Windhager, L., Zimmer, R., and Radler, J. O. (2012) Experiment and
   mathematical modeling of gene expression dynamics in a cell-free system. *Integrative Biology* 4, 494–501.
- 34. Niederholtmeyer, H., Xu, L., and Maerkl, S. J. (2013) Real-Time mRNA Measurement
   during an in Vitro Transcription and Translation Reaction Using Binary Probes. ACS
   Synthetic Biology 2, 411–417.
- 35. Niederholtmeyer, H., Stepanova, V., and Maerkl, S. J. (2013) Implementation of cell-free
   biological networks at steady state. Proceedings of the National Academy of Sciences
- 36. Gruber, A. R., Lorenz, R., Bernhart, S. H., Neuböck, R., and Hofacker, I. L. (2008) The
   Vienna RNA websuite. Nucleic Acids Res. 36, 70–74.
- 37. Herschlag, D. (1995) RNA Chaperones and the RNA Folding Problem. Journal of Bio logical Chemistry 270, 20871–20874.
- 510 38. Schroeder, R., Barta, A., and Semrad, K. (2004) Strategies for RNA folding and assem-511 bly. Nat Rev Mol Cell Biol 5, 908–919.
- ${\tt 39. \ Bionumbers, http://bionumbers.hms.harvard.edu//bionumber.aspx?id=102972\&ver=8.}$
- 40. Maslak, M., and Martin, C. T. (1993) Kinetic analysis of T7 RNA polymerase transcription initiation from promoters containing single-stranded regions. *Biochemistry 32*,
   4281–4285, PMID: 8476857.
- 41. Karzbrun, E., Shin, J., Bar-Ziv, R. H., and Noireaux, V. (2011) Coarse-grained dynamics
   of protein synthesis in a cell-free system. *Phys. Rev. Lett.* 106, 1–4.
- 518 42. Urban, J. H., and Vogel, J. Methods in Molecular Biology; Humana Press, 2009; pp 519 301–319.
- 520 43. Shin, J., and Noireaux, V. (2010) Efficient cell-free expression with the endogenous E.
  521 Coli RNA polymerase and sigma factor 70. Journal of Biological Engineering 4, 8.

- 44. Karig, D. K., Iyer, S., Simpson, M. L., and Doktycz, M. J. (2012) Expression optimization
   and synthetic gene networks in cell-free systems. *Nucleic Acids Research* 40, 3763–3774,
   10.1093/nar/gkr1191.
- Siegal-Gaskins, D., Tuza, Z. A., Kim, J., Noireaux, V., and Murray, R. M. (2014) Gene
   Circuit Performance Characterization and Resource Usage in a Cell-Free "Breadboard".
   ACS Synthetic Biology 3, 416–425.
- 46. Chappell, J., Jensen, K., and Freemont, P. S. (2013) Validation of an entirely in vitro
   approach for rapid prototyping of DNA regulatory elements for synthetic biology. Nucleic
   Acids Res. 41, 3471–3481.
- 531 47. Espah Borujeni, A., Mishler, D. M., Wang, J., Huso, W., and Salis, H. M. (2016) Automated physics-based design of synthetic riboswitches from diverse RNA aptamers.

  Nucleic Acids Res. 44, 1–13.
- 48. Iwane, Y., Hitomi, A., Murakami, H., Katoh, T., Goto, Y., and Suga, H. (2016) Expanding the amino acid repertoire of ribosomal polypeptide synthesis via the artificial division of codon boxes. *Nat. Chem.*
- 49. Pardee, K., Green, A. A., Ferrante, T., Cameron, D. E., DaleyKeyser, A., Yin, P., and
   Collins, J. J. (2014) Paper-Based Synthetic Gene Networks. *Cell* 159, 940–954.
- 539 50. Shimizu, Y., and Ueda, T. In *Cell-Free Protein Production: Methods and Protocols*; 540 Endo, Y., Takai, K., and Ueda, T., Eds.; Humana Press: Totowa, NJ, 2010; pp 11–21.

## 1 Graphical TOC entry

