

Overpowering the component problem

Matthew R Bennett & Jeff Hasty

Synthetic gene networks can be readily redesigned using new libraries of quantitatively characterized promoters coupled with predictive mathematical modeling.

The physicist John von Neumann famously said (as quoted by Enrico Fermi to Freeman Dyson)¹, “With four parameters I can fit an elephant, and with five I can make him wiggle his trunk.” By this, von Neumann meant that if a mathematical model has too many unknown parameters, it can be manipulated to fit almost anything, making the value of the modeling exercise questionable. In synthetic biology, as in physics, such model ‘over-determination’ is crippling, and progress in designing large circuits will hinge on improving our ability to predict the output of networks based on their governing architecture². In this issue, Ellis *et al.*³ describe an important advance toward this goal that relies on the creation and modeling of a large library of promoters with well-quantified inputs and outputs. Using these promoters, the authors created genetic circuits that reliably behaved as predicted by computational modeling.

Among the many challenges that face synthetic biologists, the ‘component problem’—that is, the paucity of well-characterized component parts—is imminently solvable. For example, at the transcriptional level, the most common negative regulators are the Lac and Tet promoters. These workhorses of synthetic biology come ‘as is’—there is only one version of each from which to choose. And if a network requires more than one repressible promoter, the designer has few other options. Despite this lack of components, synthetic biology has had some major successes^{4–6}, and a large fraction of successful synthetic networks have been built using these two promoters and their associated

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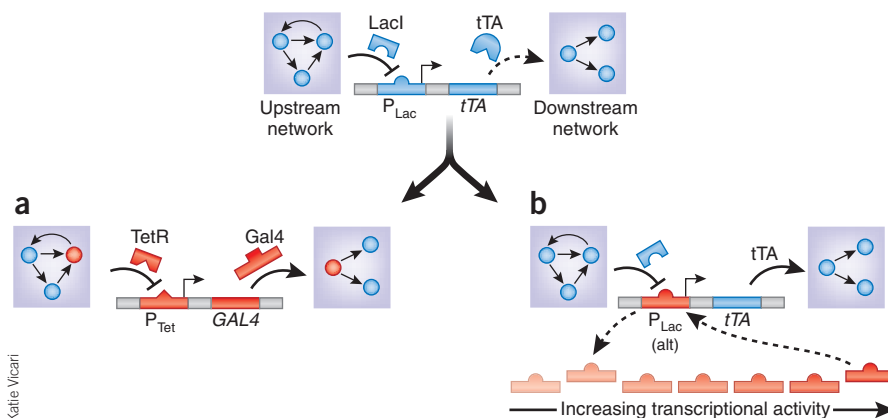


Figure 1 Design and redesign of a synthetic gene network. In this example, the researcher wishes to create an intermediate signaling gene between upstream and downstream transcriptional networks. Initially, tTA is used to upregulate the downstream network. tTA is placed under the control of the P_{Lac} promoter, which is repressed by LacI produced upstream. However, the Lac promoter is too weak to produce the intended signal. (a) One possible solution is to replace the Lac promoter with a stronger promoter. Because of the lack of available transcription factor–promoter combinations, the best option is the repressible Tet promoter. This requires changes not just in the intermediate gene, but also in the upstream and downstream components (red indicates changed components). (b) With a library of Lac promoters of varying strengths, only one component must be changed to produce the desired output.

transcription factors. However, this situation is slowly changing as various groups begin to create libraries of synthetic parts⁷.

A consequence of the component problem is an issue that might be called the ‘impedance problem’. This problem can be defined as the inability to change the dynamic range or regulatory strength of a network component so that it will work with another component without redesigning the entire network. For example, in the situation depicted in Figure 1, the designed network requires a regulatable promoter to drive a downstream signaling protein. The first experimental realization of the network might use the P_{Lac} promoter and the gene encoding the inhibitory transcription factor (LacI) that binds P_{Lac} (Fig. 1). If, after testing, the P_{Lac} promoter is found to be too weak or too noisy, the researcher must find

an alternative transcription factor–promoter pair that will increase the signal or change the frequency spectrum of the stochastic fluctuations⁸.

Because of the small number of existing components, however, this may require a complete redesign of the entire network. For instance, replacing the repressible Lac promoter with the repressible Tet promoter requires changes to the intermediate gene and to both the upstream and downstream networks. The upstream network must be reconfigured to produce the transcription factor TetR instead of LacI, and the downstream network must now receive a new input, in this case Gal4p (Fig. 1a). This is necessary because the Tet repressor (TetR) and the Tet transactivator (tTA) have the same binding domain and therefore interfere with one another.

Fortunately, the work of Ellis *et al.*³ solves

the impedance problem by solving the component problem. Using established experimental techniques, the authors screened numerous promoter variants with randomized, non-essential sequences to create a library of Lac and Tet promoters with different levels of transcriptional activity. They then fit these experimental data to computational models, which enabled the promoters to be incorporated into synthetic networks in a quantitative, predictable manner. By creating a library of promoters that are functionally the same but quantitatively different, they allowed the precise tuning of promoter strength without the need for network redesign. Therefore, researchers are no longer confined by the inherent strength of the standard Lac or Tet promoters. Instead, they can choose from a large array of variants of these promoters, each with a well-quantified transcriptional strength (Fig. 1b). Rather than redesigning the entire network shown in Figure 1, they can simply replace the weak promoter with a stronger version.

Ellis *et al.*³ went on to show that simple transcriptional control systems can be tuned by promoter choice. They used Lac and Tet promoters from their library to create a negative feedforward loop and a co-repressive toggle switch. In both circuits, they were able to use mathematical modeling to correctly predict how the behaviors of the networks would change when the promoters were swapped for others in the library. They also showed that these promoters can be used to create timers that are programmable by component choice. Their co-repressive toggle switch is used to drive the induction of *FLO1*, which is responsible for the initiation of yeast sedimentation. By changing the relative strengths of the two promoters, the timing of the event can be changed predictably.

Of course, this work is only one step toward true ‘tunability’ of synthetic gene networks. There are many factors that go into the proper control of gene activity, and many avenues through which regulation occurs^{9,10}. For instance, it may be possible to engineer libraries of transcription factors that are qualitatively the same (that is, they target the same promoter) but that vary quantitatively. With such a transcription factor library, synthetic biologists would be able to choose the regulatory strength, Hill coefficient or binding affinity of the transcription factor before network construction begins. One can also imagine tunable post-transcriptional regulators, such as kinases or proteases of differing efficiencies. Previous tunable synthetic systems have often involved small-molecule inducers that must be present in precise

concentrations in the growth medium. Libraries of interchangeable parts of differing performance provide a much-needed alternative for synthetic biologists.

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Expanded CAG repeats in the crosshairs

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Antisense oligomers targeted to CAG repeats allow allele-specific knockdown of the gene that causes Huntington’s disease.

A normal version of *HTT*, which encodes the huntingtin protein, contains 10 to 35 serial CAG triplets, but a mutant *HTT* with 36 or more repeats predicts susceptibility to Huntington’s disease¹. Such trinucleotide repeat disorders pose a special challenge for the design of oligonucleotide therapeutics as it is difficult to silence the mutant allele while preserving expression of its normal counterpart. In this issue, Corey and colleagues² show that selective knockdown of the mutant allele can be achieved using antisense oligomers targeted to the CAG repeat region. Their approach is the first to selectively silence a mutant triplet-repeat gene without relying on single-nucleotide polymorphisms. As such polymorphisms are restricted to subsets of patients, therapeutics that target the CAG region itself should be more broadly effective.

In developing therapies for autosomal dominant diseases, it seems prudent to avoid interfering with expression of the normal allele. In the case of CAG repeat expansion disorders, unselective targeting of CAG carries additional risk as the human genome contains ~80 genes with this triplet in series. Although indiscriminate knockdown of both *HTT* alleles has been advocated, this approach remains a leap of faith. Lack of normal huntingtin protein in the mammalian brain has not been shown to be safe, and treatment for Huntington’s disease, which often manifests in the fourth decade of life, generally continues for decades.

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Corey and colleagues² begin with the hypothesis that an mRNA with expanded CAG repeats might be selectively silenced based on structural differences from the wild-type mRNA. Using oligomers of peptide nucleic acid (PNA) and locked nucleic acid (LNA), they target CAG

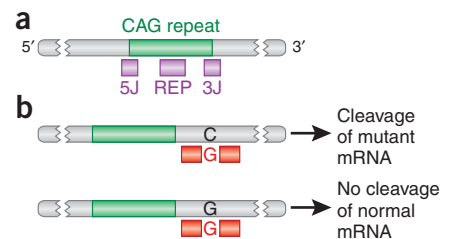


Figure 1 Two nucleic acid-based strategies for allele-specific silencing of mutant genes containing expanded triplet repeats. (a) In an antisense approach developed by Corey and colleagues², peptide nucleic acid and locked nucleic acid oligomers (purple) are targeted to regions within the tract of CAG repeats (REP) or spanning its 5′ or 3′ junctions (5J and 3J, respectively). Both classes of oligomer can discriminate a normal CAG tract from an expanded CAG tract, although the REP oligomers show greater silencing efficacy than the 5J and 3J oligomers. The authors hypothesize that allele-specific knockdown depends on differences in the three-dimensional structure of mutant and wild-type mRNAs. (b) In RNA interference approaches, double-stranded siRNAs achieve allele-specific knockdown by recognizing polymorphisms. Complementarity of the siRNA guide strand to the mutant mRNA at the SNP initiates nuclease activity by the RNAi silencing complex, whereas a selective, mismatched sequence at the SNP does not induce silencing.