Understanding biological regulation through forward synthesis

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ABSTRACT

Engineering synthetic gene regulatory circuits proceeds through iterative cycles of design, building, and testing. Initial circuit designs must rely on often-incomplete models of regulation established by fields of reductive inquiry – biochemistry, molecular and systems biology. As differences in designed and experimentally observed circuit behavior are inevitably encountered, investigated, and resolved, each turn of the engineering cycle can force a re-synthesis in understanding of natural network function. Here we outline research that uses the process of gene circuit engineering to advance biological discovery. Synthetic gene circuit engineering research has not only refined our understanding of cellular regulation, but has furnished biologists with a toolkit that can be directed at natural systems to exact precision manipulation of network structure. As we discuss, using circuit engineering to predictively reorganize, rewire, and reconstruct cellular regulation serves as the ultimate means of testing and understanding how cellular phenotype emerges from systems-level network function.
1. INTRODUCTION: THE DUAL PURPOSE OF SYNTHETIC GENE CIRCUIT ENGINEERING

Synthetic biology is a field of research that aims to establish engineering rules for the forward synthesis of cellular function (43). It was launched in the early 2000s with the publication of two artificial gene regulatory circuits; a genetic toggle switch (39), and a gene expression oscillator (termed repressilator)(29). While the field has diversified in recent years (60, 85), heavy focus continues to be placed on regulatory circuit design and construction as a means to establish user-defined control of cellular function (12). The term ‘circuit’ is used generically, but can be defined as a cellular regulatory program with specific molecular inputs, an intermediate information-processing step, and measurable output. While gene circuit engineering can vary in scope from tinkering with small synthetic test-bed networks, to the construction of complex multi-component circuitry, the process of reaching a desired target behavior is often mired by unpredictable, non-linear interactions both between circuit components and with the encompassing cellular milieu. As a result, gene circuit engineering is typically iterative, and may involve numerous cycles of design, construction, and testing as the bioengineer attempts to move from an initial, abstract description of a circuit’s behavior to its experimentally verified, physical implementation. At each step of this circuit engineering cycle, the bioengineer must continuously engage with the allowances and constraints imposed by the underlying biology of the system. Thus, while realizing designed function is usually the stated goal of a gene circuit engineering project, extracting a deeper understanding of biology is often an unavoidable bi-product (28).

We acknowledge that the motivating vision for synthetic biology has always been centered on creating useful—even transformational—technological applications. Progress in
engineering regulatory circuits to address problems in medicine (91), metabolic engineering (72), and materials science (88) have been well-reviewed elsewhere. Here, we instead highlight research that embraces using synthesis as a tool for learning about biology, and describe how the process of engineering synthetic gene circuits and network connections can enrich our understanding of the organization, function, and evolutionary design of natural regulatory systems.

We begin by discussing how foundational work engineering gene circuits—mostly in prokaryotes—provided quantitative insight into the biology of gene regulation, establishing the circuit engineering cycle as a process that can both complement and motivate basic research. These early lessons have inspired ongoing work on gene and signaling circuit engineering in mammalian systems. Our growing ability to freely manipulate network connections has enabled hypothesis-driven rewiring of native regulatory pathways. As such, circuit engineering tools can be thought of as precision-designed alleles that can be used for making systematic, targeted genetic perturbations. We highlight how synthetic circuits can be fashioned into sophisticated reporter modules, capable of converting obscure biological activity into observable, recordable data. What has emerged is a view of gene circuit engineering as a dynamic and essential partner for systems biology, capable of systematically reconfiguring, augmenting, or even replacing native networks in order to understand the adaptive forces shaping their design. In this capacity, using synthetic systems may prove the best and only way to understand determinants for complex phenotypes like cell differentiation, morphology, and fitness. We close by describing a far-reaching vision for advancing synthetic circuit engineering toward the long-term goal of system-wide synthetic control of complex phenotype.
2. SYNTHETIC GENE CIRCUITS AS PHYSICAL MODELS FOR STUDYING REGULATORY FUNCTION

A key principle to arise in the contemporary, post-genomic era is the idea that biological systems are organized according to a scalable, modular hierarchy (46, 63). Systems biology has revealed that cellular regulatory networks are constructed from a common set of molecular building blocks, and further organized into recurring patterns of connectivity (70). These network ‘motifs’ often appear in analogous physiological contexts, implying a modularity of function (46). Thus, it would appear that regulatory network behavior is ultimately an emergent property of the arrangement of constituent motifs (1).

The gene circuit engineering process has arisen as a way to test this hierarchical view of cellular regulation, by directly evaluating the physical determinants of motif function (Fig 1) (5). A critical first step in this process involves searching for molecular parts, usually by examining native networks to identify molecular components that can be modularized, abstracted, and reconnected (31). Through the process of part identification, a bioengineer implicitly evaluates how regulatory connections may have evolved, since modular, ‘tunable’ circuit components may correspond to evolvable features in natural networks (1). After working out a construction scheme, a motif can be reconstituted by establishing user control over circuit input (e.g., inducing transcription with an exogenously-administered small molecule inducer) and connecting it to a measureable output (e.g., a transcriptional GFP reporter). Function can then be directly tested in quantitative fashion, and the design can be iteratively adjusted until the desired behavior is achieved (Fig 1).

Circuit engineering is typically guided by mathematical models in which parameters represent properties like reaction rate, species concentration, and interaction affinity (58). Model-
based, *a priori* notions of gene circuit behavior may guide initial designs, but are usually adjusted following testing and redesign. Achieving close correspondence between model parameters and circuit components creates a powerful experimental framework where questions regarding motif behavior can be addressed in systematic, quantitative fashion (Fig 1): How robust is behavior to parameter tuning? How do molecular properties of the parts shape or constrain behavior? By expanding parameter space with new parts, or rewiring the circuit with new connections, what new behavior regimes can be accessed? Thus, synthetic gene circuits not only serve as physical models for exploring the limits of motif behavior, but also permit evaluation of the evolutionary logic of motif design through comparison with designs not selected by nature. Once the relationship between design and behavior are well understood, a gene circuit can be embedded within a natural network by wiring it to relevant inputs and outputs (Fig 1). In this context, the tunable circuit becomes means to probe the relationship between network structure and phenotype, and different circuit configurations can be tested for their impact on cellular behavior and fitness.

### 2.1 Insights into prokaryotic gene regulation through iterative circuit engineering

The engineering of small synthetic gene circuits in prokaryotes—an area of both foundational and ongoing research—is particularly illustrative of how iterative cycles of design and construction can yield new biological insight (15). Early progress in this area was facilitated by a detailed understanding of the functional modularity of native gene networks in *E. coli* (84). This provided the field with a ready-made collection of promoters, transcriptional repressors, activators, and copy-number-controlled plasmids in which tunability, composability, and modularity were pre-validated (Fig 2) (68), enabling facile construction, tuning, and modelling.
Design of target behavior for both the toggle and repressilator circuits—bistability and periodic oscillation, for the toggle and represillator, respectively—were model-driven, and inspired by natural behaviors: bistable trans-repression was long understood to regulate the lysogenic cycle in bacteriophage (84), while a ring oscillator architecture is a well-known feature of circadian clocks in photosynthetic bacteria (26). Initial circuit designs functioned poorly, and had to be iteratively adjusted to realize target behaviors; by changing the RBS sequence to balance TF translation for opposing arms of the toggle, and decreasing TF lifetime by appending degradation tags for the represillator. Together, these studies served as proof-of-concept that non-linear behavior could be reconstituted with molecular components not naturally associated with one another, and that an area of productive behavior space could be pin-pointed by rationally tuning component features within a synthetic design framework.

One phenomenon underscored by both the toggle and repressilator was the role of molecular noise in limiting circuit performance, as apparent cell-to-cell variations in constituent molecular species resulted in non-uniform circuit behavior, and led to eventual decay of steady-state behavior. Circuit engineering thus presented both the motivation and experimental means to investigate the molecular origins of gene expression noise—a topic given mostly theoretical treatment up until that point (69). One early inquiry into noise in bacterial gene expression utilized a simple genomically-integrated, LacI-regulated GFP reporter in *E. coli* (75). By differentially tuning transcriptional and translational rate (by respectively de-repressing LacI with a small-molecule inducer and tuning RBS strength) the authors determined that variability in protein production was the main source of noise. Subsequent work performed in yeast used a similar approach, but identified transcript initiation was the primary source of noise (9). The use of single-cell fluorescence reporters to separate noise into intrinsic (promoter-to-promoter
fluctuations) and extrinsic (global, cell-to-cell fluctuation) components (30) laid the groundwork for using synthetic circuits to study how noise is propagated within a gene networks (82) and sources of contributions to noise in gene regulation (90).

The idea that feedback and feed-forward circuitry could play an information processing role in regulatory networks was first hypothesized by Jacob and Manod in 1961 (54). Subsequent to this insight, theoretical concepts emerged regarding the role of positive and negative autoregulation in gene expression stability (23) that could only be directly tested with the advent of synthetic gene circuit engineering. Becskei and Serano constructed a simple negative autoregulatory circuit that demonstrated reduced stochastic fluctuations when single-cell expression was measured (7), while positive autoregulatory feedback as a way to achieve population bistability (6, 52). Cumulatively, understanding of the basis for both noise propagation and feedback control yielded models capable of a priori prediction of simple circuit behavior (44).

A number of recent studies have utilized the toggle and the oscillator as testbed circuits for continued exploration of network design principles. These updates illustrate an underappreciated feature of the field: beyond design cycles within a project, insight can be developed between successive projects, as circuit designs are refined over many years (Fig 2). Recently, engineering of a single-copy, genomically integrated version of the toggle switch was reported (65). Using both experiment and modelling, the authors demonstrate the single-copy toggle, which functions at lower component expression levels, is more stable and less prone to mutation—a refinement that highlights design features potentially important for epigenetic state maintenance in natural systems. Another recent study used toggle circuit engineering to demonstrate how promoter position and orientation are important network design feature (107).
Publication of a robust oscillator (97) utilized a design containing opposing positive and negative feedback loops (Fig. 2). As predicted by theory (47), enhanced stability and tunability of this design compared to the repressilator was borne out by experiment. Subsequent work used design principles to program oscillatory behavior at the population (within a biofilm) (22) and community scales (between co-cultured strains) (21). In a more recent study, the repressilator was revisited by simplifying the original circuit design in order to limit sources of noise; by removing degradation tags and limiting plasmid copy number, stable oscillations could be achieved (66). Toggle and repressilator updates also highlight an initially overlooked aspect of circuit engineering: that interactions between circuit and host cell network components can impinge on circuit function in way that initial designs often cannot anticipate. Additionally, components from different circuits can interact with each other through shared host machinery. Hasty and colleagues treated this as a feature, rather than a bug, demonstrating that the period and phase of oscillating circuits can be coupled through shared protein degradation machinery (83).

2.2. Decomposing mammalian gene regulation with synthetic circuits

Transcriptional regulation in eukaryotic cells takes place on multiple organizational scales, with contributions from transcription factor binding dynamics, chromatin regulation, and three-dimensional (3D) genome organization all conspiring to regulate promoter activity. Inability to disentangle the integration of these different regulatory modes precludes a systems-level understanding of complex processes like context-dependent cell state maintenance (108) and cell differentiation during tissue development (33). As a handful of recent studies have
shown, it may be possible to use engineered circuits to decompose overlapping regulatory contributions.

As with prokaryotes, network structure in eukaryotes is an outgrowth of promoter architecture. In both cases, trans regulation is accomplished by TF binding, but eukaryotic promoters feature multivalent, often cooperative binding of TFs to upstream enhancer sequences; a feature that underlies greater network density and regulatory complexity. Recently, a circuit engineering scheme inspired by eukaryotic promoter design was reported in yeast (61). It featured a set of synthetic zinc-finger transcription factor (ZF-TF) activators engineered to bind synthetic, orthogonal enhancer sequences upstream of a generic core promoter. Transcriptional output could be tuned by adjusting either ZF-TF affinity or the number of enhancers. Appending protein-protein interaction domains to tandem-bound ZF-TFs led to cooperative binding, enabling programming of Boolean-like logic behaviors. Since it simulates ubiquitous features of eukaryotic TF-activator regulation, this platform should scale to metazoan, enabling the engineering of multi-node networks approach natural-like sophistication. Furthermore, by leveraging cooperative binding in the upstream promoter complex, it should be possible to program TF binding assemblies capable of performing complex computations (76).

The clearest point of divergence between prokaryotic and eukaryotic gene regulation comes from the organization of eukaryotic DNA into chromatin (59), which is associated with epigenetic memory and the maintenance of heritable, self-reinforcing states of transcriptional activation and repression. A diverse array of chromatin regulators (CRs) add, remove, and are recruited through chemical marks that control histone packing and regulate binding and activity of TFs (55). Because of the number of regulatory states that can arise from combinations of modifications, TF occupancies, and spatial configurations, even a relatively simple eukaryotic
promoter can adopt a myriad of regulatory states. Our current understanding of this variation comes from extensive genome-wide mapping of the concurrence of histone modification with CR, TF, and histone occupancy (32), but details of how these configurations are decoded into a transcriptional output remain obscure.

A pair of recent studies took a bottom-up synthetic approach to understanding the relationship between chromatin regulatory state and transcription, engineering simple test-bed circuits where binding of inducibly expressed CRs were recruited to fluorescent transcriptional reporter loci. The first of these studies, performed in yeast, tested the effects of recruitment of a library of ZF-TF-CR fusions for their ability to either activate or repress reporter transcription (59). When recruited in tandem with a transcriptional activator, CRs repress, enhance, or synergize with the activator. Spatial effects of the CRs were tested: while activating CRs were found to act when recruited to promoters, repressive CRs could regulate at a distance.

In a second study, chromatin regulation dynamics were investigated using transient recruitment of different CR repressors to a reporter locus on a human artificial chromosome (8). The authors found that both repression and re-activation (upon small molecule washout) proceeded in an all-or none-fashion, and the fraction of cells repressed was proportional to recruitment duration. Post-washout time-course measurements revealed behavioral difference between the repressors, leading the authors to propose a simple model explaining the observed dynamics in terms of transitions between one of three states: active, reversibly silenced, and permanently silenced. As the authors demonstrate, some individually recruited repressors can push promoter regulation into distinct states, while others can yield mixed reversible and irreversible, with the ratio dependent on duration of recruitment.
Despite progress, gene circuit engineering strategies commensurate with the sophistication of mammalian transcriptional regulation are still at an early stage of development. To date, most engineered applications have relied on binary regulatory connections, often by superimposing bacterial TFs onto eukaryotic promoters (100). Studies described above suggest a framework more congruent with natural network design could eventually be realized, combining TF activation with chromatin-mediated regulation, integrating the ability to make arbitrary network connections with epigenetic control over expression timescale and chromosome structure.

2.3. Using synthetic circuits to understand post-translational signaling

Cells—especially those of higher-order taxa—process external signals using networks of signaling proteins that connect upstream receptor activation to downstream outputs including gene expression, metabolic regulation, and cytoskeletal rearrangement. Designing synthetic signaling circuits is challenging, requiring precise accounting of expression levels, interactions, and activities of freely diffusing pathway components (40). As a result, post-translational circuit engineering has moved at a much slower overall pace than for gene circuits, with modular parts collection and generalizable design principles still under development.

As with gene circuits, a prerequisite activity for signaling circuit engineering is assessing what commonly repeated, potentially engineerable parts, modules, and motifs can be found to comprise native networks. Reversible chemical modifications are the primary currency of information transfer in most signaling networks. Phosphorylation is the most common, but other examples including acetylation, methylation, and ubiquitination (102). Post-translational modifications carry information to targets by altering their activity, localization, stability, or
interaction specificity. Both addition and removal of marks are catalyzed by enzymes arranged in opposing, ‘push-pull’ fashion (Sidebar 1). Signaling proteins typically contain multiple domains that fall into one of two categories: catalytic domains harbor the activities that make or remove modifications, while interaction domains account for pathway wiring by specifying localization and protein-protein interactions (81). The latter include domains that mediate unregulated interactions (e.g., SH3, LIM and PDZ), but also those where binding is regulated by modifications (e.g., SH2, 14-3-3, and WW that specifically recognize phospho-peptides) (81).

A number of pioneering studies have demonstrated that domain recombination can be used as a design principle for creating new connections in signaling networks, in much the same way that modular promoter elements wire gene networks together through TF- trans regulation. Through novel combinations of catalytic and interaction domains, protein switches have been engineered to accept novel inputs and perform elementary Boolean computations (25), and even establish synthetic control over cytoskeletal morphology (106). Adapters and scaffolds are proteins composed of interaction domains that regulate signaling by organizing pathway components into complexes. Several studies have shown that pathway input/output relationships can be reprogrammed by engineering these proteins to accommodate new interactions. One study rewired scaffold-regulated osmolarity and mating response pathways in yeast (80). In another, a normally proliferative, EGF receptor-mediated input was re-routed to a cell death output via an engineered adapter protein (50).

The relative ease with which pathway connectivity can be synthetically altered suggests that domain recombination may have driven diversification of signaling pathway connectivity during evolution. This possibility was reinforced in a study where response dynamics were reshaped using engineered scaffold recruitment to overlay synthetic feedback circuitry onto the
yeast mating pathway (4). This was accomplished by coupling pathway induction to the transcription of positive and negative modulators recruited to the Ste5 scaffold through synthetic interactions. By tuning a limited set of parameters (transcription strength, recruitment affinity, and the sign and order of the effector recruitment), the authors made feedback circuit connections that radically altered pathway behavior. In a follow-up study, synthetic phospho-regulated feedback (fast) positive feedback was combined with positive transcriptional feedback (slow) to engineer temporally sensitive switching behavior – short-duration inputs were amplified by the fast loop and converted into a permanent memory state by the slow one (41).

Continued progress of post-translational circuit engineering will be driven by development of diagnostic biosensing (94) and cell-based therapeutics (36). By coupling synthetic signaling circuitry to gene expression, secretion, and cell migration, cells could be engineered to sense features of their chemical and mechanical environment, and respond in a therapeutically relevant manner through movement, secretion, or differentiation. As the repertoire of parts and design practices used for post-translational circuit engineering mature, two general strategies could be adopted: (1) a bottom-up strategy, where engineered protein components are wired together to construct signaling pathways de novo; while an enabling part toolkit has yet to be developed, adopting such a ‘model-pathway’ approach would grant precise control of molecular composition and network structure, allowing exploration of the molecular determinants underlying signaling behavior; and (2) a more top-down approach, where whole pathway modules, with known signal processing attributes, are appropriated via rewiring of inputs and outputs; such a strategy could be used to investigate the degree to which modules retain their signal processing features when placed in a different context. In an early example of this method, an scaffolded human MAP kinase cascade was orthogonally expressed in yeast (74).
Adjusting stoichiometry of both kinases and scaffold, and introducing negative regulation, allowed for tuning of pathway transfer function in model-predicted fashion, suggesting that transplanted modules placed under synthetic control are capable of a high degree of behavioral plasticity.

3. SYNTHETIC GENE CIRCUITS AS TOOLS FOR PERTURBING AND MONITORING NATURAL SYSTEMS

Extensive mapping of cellular networks using the tools and methods of systems biology has resulted in a near-complete inventory of cellular contents, and an increasingly accurate map of the interactions between them (3). Unfortunately, our ability to identify network connections has dramatically outpaced our understanding of how connectivity gives rise to function. In the following section, we discuss how the techniques of network construction and manipulation that have been developed for synthetic circuit engineering can be appropriated as tools for studying native cellular regulation. This includes the ability to wire in new connections, tune existing ones, or integrate entire circuit modules in cellular networks to report on their behavior.

3.1 Synthetic rewiring of natural regulatory networks

Traditionally, the study of regulatory network function has been conducted through reverse engineering. Understanding of how a specific motif or pathway works has often superseded the equally important question of why evolution may have chosen a particular design. Using circuit engineering tools to modify or tune network connectivity precisely and in-place provides an experimental means to test both the quantitative behavior and phenotypic consequences of evolutionarily plausible, alternative network topologies (Fig 3a). Furthermore,
assessing the behavior and fitness of rewired variants can offer clues as to why certain topologies evolved, and even what features of the environment may have driven their selection.

The use of synthetic rewiring to understand natural network design was superbly illustrated in a series of studies examining the gene circuitry underlying transient switching between competence and vegetative growth states in *B. subtilis* (13, 98, 99) (Fig 3b). Based on data from time-resolved fluorescence microscopy analysis of single cells, the authors proposed a simple model for switching in which a master regulator, ComK, is activated by an excitatory positive feedback loop, leading to a transient ‘on’ state that slowly relaxes through subsequent action of a refractory negative feedback loop. Fluctuations in gene expression can stochastically trigger activation, and also create variability in the distribution of relaxation rates. To verify their model, the authors use built several circuit variants: (1) adding a positive feedback loop that circumvents the negative feedback loop results in permanent activation (98), (2) addition of a stronger negative feedback loop results in a shorter refractory period and reduced population variation in the relaxation time (99), and (3) changing the regulatory details of the negative feedback loop (swapping transcriptional repression for protein degradation), reducing variability in competence duration (13). Cumulatively, these results argue that competence switching is determined by the intrinsic timing and noise of the network, and while alternate circuit wirings may be adaptive under particular conditions, noise in the wildtype network probably evolved as bet-hedging strategy for balancing growth with DNA absorption.

For many years, it was hypothesized that the dynamics of re-activation of latent HIV was driven by positive feedback of the virally encoded transcriptional activator tatA (Fig 4c). To investigate this possibility, Weinberger *et al.* engineered a YFP transcriptional reporter driven by a synthetic tatA autoregulatory circuit (104). Following genomic integration of the circuit using a
lentiviral vector, a combination of fluorescent microscopy and modelling was used to
demonstrate that fluctuations between ‘on’ and ‘off’ states in the circuit are sufficient to drive a
phenotypic bifurcation between latent and active HIV. In a subsequent study, the authors used
the same approach to investigate whether activation is triggered by sensing the state of the host
cell (87). By tuning their synthetic latency circuit in a manner that decoupled its dynamics from
endogenous cellular machinery, the authors showed transactivation dynamics are, in fact, an
intrinsic property of the circuit itself. This surprising result suggests that HIV may have evolved
to regulate latency using a bet-hedging strategy, the dynamics of which are agnostic to the
physiological state of its host.

Using synthetic rewiring approaches to study natural regulatory networks will become
increasingly popular as familiarity of the biological research community with circuit engineering
tools grows. Refactoring, an extreme form of rewiring that involves complete removal of native
regulatory control over a set of related genes (e.g., a gene cluster or signaling pathway), has been
largely application-driven, but could be used as a tool for decoupling the phenotypic
contributions of regulation and expression (Sidebar 2). In recent years, one notable addition to
the circuit engineering toolkit is CRISPR-Cas9, which has been used as tool to recruit
transcriptional machinery to specific genomic loci (20, 48, 86), and to construct multi-node
circuitry (57, 62). In the future, CRISPR-based regulation could be used to wire arbitrary
linkages between native loci, permitting systematic exploration of alternate network topologies.
Since many complex metazoan phenotypes, like cell differentiation and morphology, are thought
to evolve from adaptive variation within a defined molecular and regulatory framework (17),
network rewiring could be used to address questions regarding the regulatory basis for
phenotypic plasticity and evolutionary optimality.
3.2 Constructing circuit engineering tools for precision genetic screening

To a great extent, confidence in what can be learned from reverse genetic screening—observing phenotype arising from targeted genotypic perturbation—is a matter of precision in both the method used to generate alleles, and the resolution of the screening readout. When implementing genetic screens, well-constructed controls and careful interpretation of results may be required to weed out false positive or negative hits. Generally, the more precise or targeted the allele, the less likely that data are clouded by pleiotropic effects. Traditionally, genetic screens that utilize gene knockouts, targeted mutations, or over-expression may fail to decouple an allele’s phenotype from the secondary effects of genotype creation. Use of RNAi is one classic solution to this problem – knocking down transcripts decouples expression from the consequences of physically disrupting a gene’s expression locus (45). Synthetic circuitry could be used to improve readouts for genetic screens by suppressing noise, or improving resolution of allelic effects by amplifying small changes in output.

Through precise and quantitative control over gene expression, circuits can be used to control the timing and dose of allele presentation. For example, expression of an unstable or toxic allele can be kept ‘caged’ until the user is ready to assess phenotype. In one recent implementation of this strategy, an orthogonal, inducible protein degradation system was developed for use in bacteria (16). This was accomplished by appropriating Lon protease-specific tmRNA degradation tags from *M. florum*. Screening of a small library of tags yielded variants that were recognized by *mf*-Lon, but not by *E. coli* ClipXP/AP and native Lon. As the authors convincingly demonstrate, induction of *mf*-Lon facilitated rapid and specific degradation
of endogenous proteins bearing the tag. Moreover, degradation was tunable, allowing for tightly controlled titration of protein expression levels. A library was composed by appending degradation tags to 238 of 305 of essential *E. coli* genes, and screened for antibiotic sensitivity, identifying several potential new drug targets.

Network wiring variants can be systematically generated and screened to address important evolutionary questions (Fig 3c). For example, gene duplication and recombination events are thought to give rise to new regulatory connections (73), but it is unknown if intermediate alleles provide any adaptive advantages. Serrano addressed this issue by rewiring the *E. coli* transcriptional regulatory network (53) (Fig 3d). To generate new network linkages, the authors generated an ~600-member plasmid library containing interchanged combinations of promoter and coding sequences selected from genes in which one TF regulates the transcription of another. When introduced into cells, a majority of library members were tolerated; few showed growth defects, while several actually demonstrated increased fitness under certain selective pressures. Thus, the *E. coli* transcriptional network appears robust to individual rewiring events, suggesting recombination through duplication may allow gene networks to explore new, potentially adaptive configurations while incurring little fitness cost.

3.3. Using synthetic circuits to reporting on natural systems

Few tools in biological research have had as much impact as the voltage clamp, which was used to study the physical and molecular basis of electrical conductance across cell membranes. In its original implementation, the device worked by immersing an axon in a saline bath and using an electronic control circuit to maintain (‘clamp’) a constant transmembrane
voltage (49). By measuring current necessary to sustain the target voltage, the device could record results of changes to membrane potential, enabling decoupling of action potential into the movement of Na+ and K+ during respective depolarizing and rectifying phases. We envision a potential role for engineered circuits similar in impact to voltage clamping – by using them to interface with and report on difficult-to-observe events occurring with or experienced by cells.

Synthetic gene circuitry can be used to convert signals of rare or transient molecular species into readable outputs. For example, synthetic riboregulators—mRNA secondary structure elements that block RBS access—can be disengaged in the presence of trigger RNA, permitting translation (14, 51). A versatile riboregulator-based circuit engineering platform was recently reported in E. coli (42) capable of gating by any RNA sequence, and was used to detect endogenous mRNAs. Toehold detection has also been used to construct highly sensitive cell-free, paper-based detection circuits that can be configured to convert the presence of diverse nucleic acid species into a macroscopic readout (77). Potential applications for cell-free diagnostic circuits are numerous, including detection of rare transcripts or proteins in biological or environmental samples (78, 79). In another recent example of circuit-based RNA detection, Benenson and colleagues used intracellular miRNA detection circuitry to reports on cell state in mammalian cells (105). By configuring logic gates to detect sets of miRNA species, they engineered a ‘classifier’ circuit that could identify a cancer cell phenotype and trigger apoptosis.

In one recent example of circuit-mediated interconversion of a molecular signal, circuitry was engineered to detect intracellular DNA sequences (93). The system featured a sensor module comprised of two ZF arrays programmed to bind to adjacent target sequences. Tandem sensor binding triggers an intein trans-splicing event, linking together associated TF and transcriptional activator domains that can trans activate a GFP reporter module. Sequence sense and response
worked effectively in mammalian cells, and the authors demonstrated the ability to couple
detection of viral infection to apoptosis. In principle, detection capabilities of the system could
be expanded by swapping the ZF arrays for TALEN or CRISPR, enabling versatile real-time
detection of the occurrence or movement of rare nucleic acid species (e.g., a latent virus) within
a cell population or tissue, or as a live readout of specific mutations or chromosomal
rearrangements.

One emerging application space is the use of molecular memory circuits to ‘record’ the
occurrence of discrete cellular or environmental events. Recombinase-based switches, where
DNA segments flanked by recombination sites are either excised or inverted upon recombinase
expression, can be used as digital memory elements (10). Recombinase-based circuitry has been
configured to perform simple computations, including Boolean logic (11, 92, 103) and event
counting (38). Analog recording of transient input has been realized using an engineered retron
to generate ssDNA that can introduce genomic sequence mutations through strand invasion (34).
When distributed across a cell population, such a ‘tape recorder’ circuit can integrate input
duration and intensity based on the proportion of cells that undergo recombination. More
recently, a system that records successive events using CRISPR-based mutagenesis coupled to a
fluorescence in situ hybridization readout was used to track mammalian cell lineage (37).

Combining computation with memory could become a powerful tool for understanding
the order and timing of cellular events. One recent demonstration utilized recombinase switches
to construct state machine circuitry capable of computing the order of any combination of three
distinct inputs (89). Synthetic ‘reporter circuits’ that combine compute and record functions with
a programmed sense and response could be used for a variety of applications involving
extraction of conditional or time-resolved information from complex environments. For example,
cells introduced into natural ecosystems could be programmed to detect, amplify, and record rare ecological events. Reporter circuits introduced into cells at early stages of embryonic development could be used to record the order and magnitude of events encountered by different cell lineages during tissue development.

4. FUTURE: USING SYNTHETIC GENE CIRCUITS TO UNDERSTAND THE REGULATORY BASIS OF COMPLEX CELLULAR BEHAVIOR

Circuit engineering offers a formidable set of tools for construction, rewiring, and refactoring regulatory networks. As these techniques continue to advance, we may begin to view synthetic biology as a way to deconstruct the complex biological phenotypes that emerge from regulatory network structure. Below we discuss several areas where we believe synthetic circuit engineering will make an impact in the years to come.

4.1. Synthetic biology as in vivo biochemistry: the cell as a test tube

Much of our understanding of the behavior of biological macromolecules comes from examining them in isolation from their native cellular context. By reconstituting macromolecular species and manipulating their content and concentration in vitro, interaction affinity, complex stoichiometry, and induced changes in activity can be measured (Fig. 4a). While using this approach to examine regulatory network components on an individual basis offers mechanistic insight, it yields little understanding of how systems-level behavior emerges from collections of components in vivo. A rich history of reconstituting complex, multispecies systems using cell extracts has served to partially bridge this gap (67). For example, reconstitution of transcription
and translation apparatuses, involving study of both purified species and in cell lysate, revealed many essential mechanistic details of the central dogma (64).

Traditionally, the goals of in vitro reconstitution have been similar in spirit to synthetic biology: determine functional sufficiency by rebuilding a biochemical system from the bottom-up. However, living cells maintain their physical organization, metabolism, and regulatory dynamics far from thermodynamic equilibrium, and provide a chemical and physical environment fundamentally unattainable through reconstitution. One possible strategy to overcome this gap is to use circuit engineering tools to reconstitute biochemical systems in a non-native, in vivo environment, effectively using cells as living test tubes (Fig 4a). In this capacity, synthetic gene circuitry could be used to titrate component expression levels or assemble complexes at a specified rate or in a defined order. Reconstitution using circuits could give the user a degree of control on par with in vitro reconstitution, but with the added possibility of directing assembly to a particular subcellular location, perturbing behavior by changing the intracellular environment, or connecting input and output of a biochemical system to endogenous networks.

This approach is exemplified by two recent studies in which synthetic circuitry was used to reconstitute systems. In one study, the authors sought to demonstrate that gene expression noise in E. coli can be tuned by repressor-operator binding interactions (56) (Fig 4b). Using FISH to assay mRNA output from a transcriptional reporter, and adjusting repressor expression and affinity, the authors were able to demonstrate that expression noise could be modeled as a function of regulatory configuration. In another study, a minimal biochemical circuit yielding robust cell polarization was reconstituted in yeast (19) (Fig 4c). The circuit motif, which the authors computationally demonstrated could produce a polarized intracellular distribution,
featured reciprocally two positive autoregulatory nodes that engage in reciprocally negative regulation. This motif was realized using elements that coupled recruitment to activation of the small GTPase Cdc42 and synthesis of PIP3. Model predictions were largely accurate: cells harboring a precisely tuned version of the motif demonstrated individual puncta, indicative of localized circuit activation. Both studies featured circuitry-enabled titration of in vitro-characterized components. More meaningfully, while both systems could have been reconstituted in vitro, using circuitry to reconstitute them in an in vivo context was essential for observing their behavior.

‘In vivo reconstitution’, as this approach may come to be known, could be used to study systems with complexity too unwieldy for in vitro work, or behaviors that only manifests in an intracellular setting (e.g., modules that exert physical force on cellular shape or 3D organization). By leveraging the rich history of biochemical research derived from buffer and extract systems, this approach could be used to compare in vitro and in vivo properties of reconstituted systems. One intriguing possibility lies in the development of parts frameworks amenable to testing in both settings. Candidate systems might consist of well-behaved, easy-to-purify proteins that could be readily studied in isolation, interfaced with extract systems, and also deployed in vivo using a circuit engineering scheme. This would allow for decoupling of component biophysical properties from the effects of both the intracellular chemical environment and the physical organization of intact cells.

4.2. Evolving circuits to explore adaptive fitness

Despite circuit engineering advances, minimizing interactions with endogenous host cell systems remains an outstanding challenge. As discussed above, circuits often impose a fitness
cost on the host, either through depletion of resources, or by interference with native network function. As a result, loss-of-function variants that ameliorate cost may spontaneously arise and rapidly sweep through a cell population. To avoid this, circuit engineering will need to identify and incorporate design features that maximize stability, especially for applications requiring engineered cells to retain circuit function over many generations. One study investigating the molecular determinants of fitness burden in *E. coli* tested a small library of circuits by simultaneously measuring fitness cost and resource depletion using a ‘capacity monitor’ circuit (18). By examining the relationship between diminished fitness and capacity for different circuit designs, the authors implicated the depletion of free ribosomes as a primary source of fitness burden.

Further investigation into long-term circuit stability may involve using experimental evolution to monitor cultures of circuit-bearing cells for the acquisition of adaptive mutations. By observing how circuits ‘break’ upon mutation, specific circuit failure modes could be uncovered. By the same token, adaptive mutations that relieve fitness burden in the face of intact circuit function could be identified. Indeed, it may be possible to produce selection schemes that stipulate circuit function as a selective pressure, thereby forcing the host cell to adapt. To achieve this, automated continuous growth platforms designed to maximize throughput, while allowing for automated manipulation and live readout of culture conditions (e.g., growth rate, GFP fluorescence) could be employed to simultaneously track culture fitness and circuit function (101), while NGS could be used to examine the resulting mutational landscape in both the circuit and host genome.

Far from viewing it as merely a factor complicating engineering, we see evolutionary instability of synthetic regulatory circuits as an opportunity to investigate the relationship
between network structure and cell fitness. Using experimental evolution, synthetic circuit-host interactions can be used to understanding how cells incorporate adaptive regulatory network connections under diverse selective pressures. The consequences of new circuitry or rewired linkages could be studied by examining fitness tradeoffs between the cost of regulatory reconfiguration and fitness advantages conferred by newly acquired phenotypes. Evolution experiments could be used identify environmental conditions under which rewiring is rejected or accepted, while observing adaptive or compensatory mutations that arise to maximize fitness. One long-term goal might be creating regulatory systems that use novel molecular currencies (e.g., new transcription factors or post-translational modifications) and operate orthogonally to natural ones. Experimental evolution could be used to assess to what extent integrated parallel programs would interact with native regulatory machinery.

4.3. Scaling up circuit engineering to accelerate discovery

Synthetic circuit construction has been traditionally conducted at relatively low throughput, essentially operating at the pace and scale of an artisanal craft (Fig 5). Recent work has focused on accelerating the engineering cycle in order to move beyond testing only a handful of circuits at each turn (15). DNA assembly techniques have enabled rapid, combinatorial construction of circuit variant libraries (27), while bioinformatics-driven ‘parts mining’ efforts have expanded parts availability (96). Both of these developments have synergized with rapidly decreasing gene synthesis costs, allowing circuit engineering projects to more rapidly move from design to construction. In conjunction with updated modelling techniques that use context-based part compatibility and assembly rules (2), these developments have set the stage for exploration
of larger, more complex circuitry, and the field appears poised to enter a more industrial phase where automated, predictive design becomes regular practice (71) (Fig 5).

In recent years, NGS technology has radically enhanced our ability to obtain genome-scale information describing cellular content and organization. Application of NGS to synthetic circuit engineering remains largely unexplored, but could dramatically enhance engineering cycle throughput. By combining NGS with high-throughput single-cell analysis, barcoded circuit libraries could be synthesized in batch, subjected to screening or selection, and then sequenced. Creating a diversity-based engineering workflow that combines NGS with automation—both for circuit synthesis and testing—would generate tremendous quantities of data on circuit behavior, part performance, and context rules in each turn of the engineering cycle, and permit comprehensive mapping the relationship between circuit function and design (Fig 5).

Transforming gene circuit engineering into a data-driven, post-industrial process would dramatically broaden the scope of all research areas discussed in this review. For example, expanding diversity used for circuit construction could redefine our understanding of the relationship between sequence features and regulatory behavior. Massive diversification of part sets and circuit topologies would permit testing a design space that approaches the mutational landscape available to natural systems during evolution. Up until this point, circuit modelling has described essential features of circuit behavior in terms of part-specific parameters. For a diverse circuit library, since precise mechanistic relationships between sequences variants and circuit behavior are difficult to discern, it may be necessary to build rules-based models to augment traditional approaches. Machine learning algorithms could be used to train a model on a discreet area of design space, predict a more expansive space, and then iteratively refined using subsequent rounds of high-throughput design and testing. Upon model convergence, interesting
or unusual areas of the space could be examined to understand the molecular basis for model-delineated rules, possibly leading to discovery of previously unknown regulatory phenomena.

5. CONCLUSION

Historically, our understanding of cellular regulation has come from perturbation and measurement working interdependently, leveraging classical tools of genetics and molecular biology to map network connectivity and information flow. In recent years, with the advent of genomics, high-throughput single-cell analysis, and multi-scale imaging, our measurement and mapping capabilities have raced far ahead of our ability to make the proportional perturbations needed to understand their function. We foresee the tools of synthetic gene circuit engineering providing the means to close this gap. In this capacity, the role of synthetic biology may be one of final arbiter for models forwarded by systems biology. At present, our ability to rewire and recapitulate natural network design is at different stages for different systems. While constructing complex, ‘natural-like’ circuitry in prokaryotic gene networks is well-developed, the same level of bottom-up control in eukaryotic and metazoan systems remains a work in progress. Combining the ability to make new connections with an accelerated engineering cycle should facilitate rewiring, refactoring, and reconstitution of increasingly complex regulatory systems. To this end, rather than studying discreet circuit function using a single type of network connection (e.g., a circuit composed of entirely transcriptional network connections), focus should turn to obtaining comprehensive multi-scale engineering control over entire cellular subsystems in a manner that subsumes native behavior, while granting the ability to systematically test alternate, unnatural designs. Such a ‘synthetic regulatory capture’ approach could be used to refactor an organism’s stress response program in order to understand the
operational design rules underlying fitness, or to acquire engineered control over an animal developmental module to investigate the regulatory determinants of morphology. To this end, other branches of synthetic biology, like synthetic genome engineering, could merge with gene circuit engineering to enable large-scale re-organization of regulatory networks at the whole-organism level. One medium-term step in this direction might involve layering regulatory machinery over a minimal bacterial genome to assess fitness advantages gained through increasing degrees of regulatory control. As our ability to reconstruct network function becomes more sophisticated, we anticipate that understanding of cellular regulatory network design will increasingly come from using forward engineering and manipulation, causing meaningful distinctions between synthetic biology and systems biology to gradually disappear.

**DISCLOSURE STATEMENT**

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

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REFERENCES

KEY WORDS

**Synthetic biology** - engineering discipline that uses forward engineering to establish control over cellular behavior

**Regulatory network** – a physically interlinked collection of molecules used to transmit information intracellularly

**Synthetic gene circuit** – a unit that features an identifiable molecular input, intermediate information processing step, and a molecular output

**Engineering cycle** – a single iteration in the circuit engineering process whereby a synthetic circuit design is constructed, tested, and evaluated for its ability to achieve a desired target behavior

**Motif** – a recurring pattern of molecular connectivity found within networks

**Refactoring** – for a set of related genes, engineering the replacement native regulation with user defined, synthetic control

ABBREVIATIONS

**RBS** – ribosome binding site

**TF** – transcription factor

**SH3** – Src homology 3
LIM – Lin11, Isl-1, and Mec-3.

PDZ - post synaptic density protein, Drosophila disc large tumor suppressor, and zonula occludens-1 protein

SH2 – Src homology 2

EGF – epidermal growth factor

MAPK – mitogen activated protein kinase

FISH – fluorescence in situ hybridization

PIP3 - phosphatidylinositol (3,4,5)-trisphosphate

NGS – next generation sequencing

CRIB – Cdc42 and Rac interactive binding

PH – pleckstrin homology

GAP – GTPase activating protein

SUMMARY POINTS

1) Designing, building, and testing synthetic gene circuits can be used to evaluate hypotheses regarding the modular organization and function of regulatory networks.

2) As the most developed area of the field, synthetic gene circuits in prokaryotes exemplify how biological understanding can be acquired through iterative engineering cycles.

3) While part sets and strategies for circuit engineering in metazoan systems are still emerging, testbed circuits can be used to decompose and understand the modular design principles of both transcriptional and post-translational regulation.
4) By furnishing biologists with a toolkit for precise manipulation of regulatory network connectivity, gene circuit engineering enables the creation of sophisticated genetic perturbations for investigating systems-level function.

5) Circuit engineering can be used to construct reporter circuits capable of converting obscure molecular species or cellular events into measureable, recordable data.

6) By scaling up circuit engineering approaches it may be possible to modify or reconstruct regulatory networks in a way that allows us to understand how network structure shapes complex phenotype.

FUTURE ISSUES

1) Emphasis should be places on developing techniques that accelerating the gene circuit engineering cycle. Automated combinatorial library assembly, high-throughput screening, and NGS should be incorporated. A diversity-based workflow could be used to comprehensively map the relationship between network structure and function.

2) An engineering grammar should be devised for bottom-up construction of post-translational signaling networks—similar to modular building blocks used for transcriptional networks. Signaling motifs reconstituted with such a system could be used as physical models for studying signal transduction.
3) Focus in prokaryotic gene circuit engineering should migrate towards using experimental evolution of synthetic gene networks to study the relationship between regulation and fitness. Experimental tools will need to be developed to assess the relationship between environmental pressure and circuit-host coevolution.

SIDEBAR 1. The push-pull cycle as the fundamental engineering unit of post-translational signaling circuits.

In protein signaling pathways, addition and removal of chemical modifications are catalyzed by pairs of antagonistic enzymes. Phosphorylation, the most common modification, is catalyzed in the forward direction by a kinase, and by a phosphatase in the reverse direction. Other examples include acetylation and methylation which play a role in regulating chromatin state. Opposing enzymes, along with their mutual target, comprise a ‘push-pull’ motif (24). Changes in forward and reverse reaction rates within a push-pull (e.g., resulting from regulatory input that alters either activity) result in a rapid shift in equilibrium between modified and unmodified target. Both the rate and magnitude of this shift, as well as the rate at which the original equilibrium is restored upon input removal, are dependent upon the molecular features of a push-pull, which include component concentration, interaction affinity, and activity. From an engineering perspective, this correspondence between molecular configuration and quantitative behavior makes the push-pull attractive as a modular building block, and we argue that it should be considered the fundamental unit for post-translational circuit construction. Circuits built from interconnected phosphorylation push-pulls could be configured to receive upstream input from a receptor, or connected to an output by activating transcription, or inducing cytoskeletal
rearrangement, or targeted secretion. Synthetic push-pulls could be linked together to form higher-order information processing circuitry; they could be vertically arranged by coupling target modification of an upstream unit to forward or reverse activities of a downstream unit or, if multiple pairs acted on a single target, it may be possible to program sophisticated logic functions by coupling combinations of phosphorylation events to an output.

SIDEBAR 2. Refactoring regulatory systems to probe and regulation

Refactoring is the process by which a set of related genes are removed from their native regulatory context and placed under synthetic expression control. Refactoring a system establishes the equivalent of a molecular tuning board by eliminating pre-translational regulation of gene expression, recoding constituent genes for optimized translation, and placing them under control of orthogonal promoters of adjustable expression strength. Using gene synthesis combined with combinatorial assembly techniques dramatically augments exploration of regulatory space for a refactored system, allowing for simultaneous tuning of component expression levels and introduction of alternate or mutant alleles. Additionally, it becomes possible to wire user-defined regulatory links specifying input control, intra-system regulatory connections, or orthogonal output. Bacterial gene clusters, which can encode related sense-and-response elements, transcription factors, and biosynthetic enzymes, have been an early target for synthetic refactoring (35). Gene clusters are good targets due to often opaque regulatory features that result from close proximity and potential co-regulation of constituent ORFs. Recently, the Nif operon, which encodes nitrogen-fixing machinery from Klebsiella oxytoca was refactored and expressed orthogonally in E. coli (95). After placing all 16 genes under synthetic expression control, the authors tested several hundred variants, finding that their best was able to recover
most of the nitrogen fixing capacity of the WT cluster. Using a similar approach, it may be possible to study mammalian signaling using pathway refactoring to encode a post-translational network.
FIGURE LEGENDS

Figure 1. The synthetic gene circuit engineering cycle. Synthetic circuit engineering involves iterative cycles of design, building, and testing, where insight from one cycle is incorporated into the next as a bioengineer attempts to achieve a target behavior. Inception of a circuit engineering project is inherently hypothesis-driven; circuits are designed with specific target behaviors in mind, often using natural circuit motifs as inspiration. By attempting to create a synthetic version of the motif, functional modularity is implicitly evaluated. Theory of motif function may inform initial circuit designs, which are built from sets of molecular parts culled from native networks. Initial testing of circuit input/output function may involve connecting to a user-defined input and measurable output, permitting evaluation of both dose- and time-dependent behavior. Convergence of circuit design and model prediction results in a model system for studying regulatory properties of the motif; tuning the circuit through changing the strength of network, or rewiring with new links, allows systematic assessment of the range of behavioral available to the circuit. Embedding a tunable synthetic circuit within a native network grants insight into the relationship between circuit tunability and phenotype, and can be used to investigate relationships between network structure and cell behavior. In aiming to fulfill target behavior, different modes of experimental testing can feed back (black lines) to reshape circuit design, challenging and growing our understanding of regulatory network function.

Figure 2. Learning through iterative engineering of model prokaryotic gene circuits. Successive efforts to construct bistable genetic toggle switch and oscillator circuits have resulted
in progressive refinement of gene circuit engineering, leading to concomitant insight into design principles of prokaryotic gene regulation. For both circuits, initial designs drew heavily from historical research into gene regulation in *E. coli*. Deep understanding of the molecular basis of TF-mediated promoter regulation and theory of gene expression dynamics enabled gathering of parts that could be used to design circuitry embodying simple non-linear behavior: inducible bi-stability and stable, periodic oscillations. Inaugural designs (toggle and repressilator) both successfully captured target behaviors by part tuning, but failed demonstrate to robust, long-lived steady-state behavior. Success (and shortcomings) of these initial designs motivated inquiry into diverse topics, including sources of gene expression noise and the role of feedback in network dynamics. Next-generation designs used more precise approaches; utilization of more varied parts and updated modeling approaches improved behavioral robustness. These advances included engineering a single-copy toggle switch and, in the case of the oscillator, utilization of an alternative circuit design. More recent design iterations have highlighted the role of circuit context and host-circuit interactions in regulatory network behavior.

**Figure 3. Rewiring native regulatory networks.** A. By wiring new linkages, circuit engineering tools can be used to explore the functional consequences of alternate network wiring topologies. B. Rewiring the transcriptional network in *Bacillus subtilis* that mediates competence. Spontaneous, noise-driven elevation in the activity of the master transcriptional regulator ComK results in activation of competence, followed by slow, relaxation back to basal activity (sporulation). Dynamics of these excitatory and refractory phases are mediated by positive feedback (blue) and negative feedback (yellow) circuits respectively. Various rewiring schemes were used to determine the relationship between circuit architecture and function; rewiring to bypass the negative feedback loop eliminated the refractory phase, while adding in a
new negative feedback loop to weaken the positive loop resulted in a faster refractory phase while diminishing the variability in the normally noisy refractory phase. Replacing the negative feedback loop with a simpler, protease-mediated loop resulted in similar dynamics, but diminished variability. C. Circuit engineering tools can be scaled to phenotypically screen network wiring variants. D. Systematic rewiring of transcriptional regulatory network in *E. coli*. A library of novel network linkages created by shuffling coding sequences of various transcription factors and sigma factors against corresponding promoters. Library members showed few fitness defects and minimal changes in genome-wide transcription. For some members, enhanced fitness was demonstrated under selective conditions, suggesting that the native *E. coli* transcriptional network may be generally robust to spontaneous rewiring that yield specific adaptive advantages.

**Figure 4. Using synthetic circuitry to reconstitute biochemical systems in vivo.** A. Gene circuit-based reconstitution can be placed on a continuum with traditional techniques of buffer- and extract-based reconstitution. Using in vitro methods, control over the composition and concentration of the components of a biochemical system permit quantitative determination of its biophysical characteristics (e.g., interaction affinity, complex stoichiometry, and activity). Extract systems allow in vitro reconstitution of higher complexity systems. While enabling interaction with intracellular components, extracts are unable to fully replicate the physical and regulatory features of an intact cell. In vivo reconstitution uses synthetic circuitry to precisely control the relative stoichiometry, timing, and localization of component expression. Additionally, circuits can be used to integrate biochemical systems into cellular networks to assess effects on cellular function. B. Testing effects of promoter regulatory architecture on
transcriptional noise. Using a simple testbed circuit, the possibility that transcriptional noise can be tuned by altering promote architecture was tested by counting single cell transcripts using fluorescence in situ hybridization (FISH). Tuning both expression level and affinity of RNA polymerase and the LacI repressor, various promoter configurations were tested against a quantitative model, and shown to harbor different characteristic expression noise. C. Synthetic reconstitution of a minimal cell polarization circuit. A self-organizing cell polarization circuit was described in yeast featuring components expressed at defined concentrations and subcellular localizations. A mutual inhibition circuit was created using protein domain fusions; PIP3 production creates localized positive feedback through PH (binds to PTP3) domain-mediated recruitment PI3 kinase, while PH-domain recruitment of a GAP inactivates Cdc42 locally. Active Cdc42 globally inhibits PIP3 production through recruitment CRIB (binds Cdc42) domain-phosphatase fusion. The circuit demonstrated spontaneous formation of individual, long-lived poles as visualized by localized recruitment of a PH-GFP fusion.

**Figure 5. Evolution of the gene synthetic circuit engineering cycle.** The scale and scope of inquiry into regulatory network design and function will depend on continued progress in circuit engineering techniques. The field began as an artisanal craft, relying on limited part sets, traditional techniques of molecular cloning, and low-throughput building and testing. While critical for proof-of-concept engineering of simple circuits, insight gained during each cycle was largely intuitive, as non-standardization of parts and techniques made design rules difficult to generalize. Engineering approaches are beginning to transition into a more industrial mode in which the acquisition of larger collections of better characterized parts permit more ambitious designs. This work is bolstered by low-cost DNA synthesis and development of circuit assembly
techniques, both of which enable automation of building and testing phases. Higher throughput, more systematic data acquisition in each cycle affords detailed, parameter-oriented profiles of individual parts. This, in turn, enables the engineer to pinpoint more robust, precisely-tuned circuit behavior within a multi-dimensional design space. The throughput and precision of this workflow could be used to address the complexity presented by larger networks where part compatibility and circuit-host interaction are at issue (e.g. rewiring large regulatory modules or refactoring gene clusters). The field awaits transition to a post-industrial, information-based mode, where fully integrated automation of design, build, and test phases are driven by AI and machine learning algorithms. By leveraging large-scale library synthesis and NGS as a means to scale input and output, large circuit design libraries can be tested at each turn, gathering data that establish rules-based relationships between sequence-level features and synthetic network behavior. Comprehensive, closed-loop characterization of circuit design space could be realized when data collected at each iteration is used to both refine models, and suggest subsequent design spaces for testing. This mode of engineering could be used to probe design spaces of the highest complexity, including genome-scale refactoring and perturbation of regulatory networks.
**Figure 1**

**GENE CIRCUIT ENGINEERING CYCLE**

**DESIGN**
- Identify motif
- Target behavior
- Theory of motif function

**BUILD**
- Encode circuit
- PART TOOLKIT
- Gene circuit parts taken from native networks
  - TFs (repressors, activators)
  - Modular promoters elements

**TEST**
- Test circuit I/O
- Test for phenotype
- Host cell
- Reporter output
- GFP luminescence
- Quantitative behavior
- Investigate I/O expression, affinity, activity
- Theoretical space

**TUNE CIRCUIT BEHAVIOR**
- Tune circuit behavior
- Explore behavior space
- Alternative designs, evolutionary logic?
- Test temporal and dose-response behavior
- Iteratively refine behavior, converge with model
- Use engineering tools to further tune and rewire circuitry

**EMBED CIRCUIT**
- Embed circuitry in a native network
- Study relationship between network structure and cellular behavior
- Test for phenotype
- Cell function
- Cell fitness
E. COLI

UNDERSTANDING of GENE REG.
- TF structure & function
- understanding of regulation
- theory of regulatory dynamics

TOOLS for CIRCUIT ENGINEERING
- defined parts set
- theory of circuit dynamics
- small molecule induction
- tuning expression regulation
- GFP trans. readout

TARGET BEHAVIORS

BISTABLE SWITCH
- IN1 IN2
- multi-copy
- bistability
- decay of stable steady-state

GENE OSCILLATOR
- tuned transcription strength
- tuned component interactions
- limited oscillation
- decay of steady-state
- stable oscillations

TOGGLE
- Gardner et al. (2000)
- reciprocal repression
- IN1 IN2
- multi-copy

SINGLE COPY
- Lee et al. (2014)
- single-copy
- tuned transcription strength
- tuned component interactions
- improved stability
- diminished load

PART CONTEXT
- Yang et al. (2017)
- improved stability
- greater sensitivity

REPRESSILATOR
- Elowitz & Leibler (2000)
- 3-node ring (repressors)
- multi-copy
- limited oscillation
- decay of steady-state

2-NODE
- Stricker et al. (2008)
- alternate design (repressor + act.)
- detailed model
- stable oscillations

LIMITING NOISE
- Potvin-Trottier et al. (2016)
- insulate repressilator from sources of noise
- lower copy number
- remove degradation tags
- stable oscillation
- limited cell-to-cell variability

TARGET BEHAVIORS

- stable oscillation
- limited cell-to-cell variability
- detailed model
- understanding of modularity
- sources and propagation of gene expression noise

- effect of context on network behavior
- effect of host-cell interaction on behavior

- basis of robust behavior
- relationship between part molecular properties and regulatory behavior

- understanding of modularity
Figure 3

A) **NATIVE PATHWAY REWIRING**

**WT PATHWAY** → **REWIRED VARIANTS**

- **REWIRE PATHWAY**
  - create new links/nodes
- **ASSAY PATHWAY**
  - pathway output
  - cell function

B) **B. subtilis**

**REWIRE COMPETENCE CIRCUITRY**

- Excitatory +fb
- Refractory -fb
- bypass neg fb
- ComK pos fb
- ComS neg fb
- weaken pos fb
- replace neg fb
- no relaxation
- reduced duration
- reduced variability

C) **SYSTEM-WIDE REWIRING**

**WT NETWORK** → **REWIRED NETWORK**

- systematic regulatory network rewiring
- pathway output
- fitness

D) **E. coli**

**REWIRE TRANSCRIPTIONAL NETWORK**

- Generate library 600 TF-TF regulatory connections
- Asses fitness defects, adaptations

**SHUFFLE SEQUENCE: NEW CONNECTION**

- **NATIVE NETWORK**
  - A 1
  - B 2
  - C 3

- **SHUFFLE SEQUENCE**
  - A 1
  - B 2
  - C 3
**Figure 4**

A. **MULTISCALE BIOCHEMISTRY**

- **TUNING**
  - Composition
  - Species concentration
  - Mutant alleles
  - Chemical environment
  - Physical environment
  - Subcellular localization
  - Interaction w/ cellular networks

- **IN VITRO**
  - Extract
  - Reconstitute with purified components

- **IN VIVO**
  - Reconstitute with synthetic circuitry

B. **TUNING NOISE IN TRANS. REG.**

- **TESTBED CIRCUIT**
  - TUNE copy no. & affinity
  - RNA polymerase
  - LacO • assess transcripts cell-1 by FISH
  - Noise is tunable with promoter architecture

- **mRNA**
  - Measure noise

C. **CELL POLARITY CIRCUIT**

- **MUTUAL INHIBITION MOTIF**
  - PIP3 production
  - Cdc42 activation
  - PIP3 deactiv. Cdc42 dephos.

- **Measure localization**
  - GFP localization
SCALING UP THE CIRCUIT ENGINEERING CYCLE

PAST ARTISINAL
- small, specialized parts sets
- conventional molecular cloning
- nonsystematic design process

10^5 - 10^6 per cycle

NOW INDUSTRIAL
- large, paramaterized parts collections
- discontinuous, partially automated workflow
- each cycle: acquire parts and context data

10^7 - 10^8 per cycle

FUTURE INFORMATION
- relate sequence elements to network features
- ultra HT library design and testing (NGS)
- AI-driven determination of library content

10^9 - ? per cycle

TARGET BEHAVIOR

DESIGN
- simple quant. models
- anecdotal 'design rules'
- design intuition

BUILD
- parts collection
- manual combination
- low throughput

TEST
- context, part compatibility rules
- DNA synthesis
- HT circuit assembly
- HT circuit analysis

LEARN
- parameters, context rules
- DNA synthesis
- HT circuit assembly
- HT circuit analysis

- closed-loop: automated design
- AI
- paramaterized parts repository
- bulk library construction
- next-gen sequencing

- Genome-scale regulatory network rewiring, refactoring
- Systematic regulatory control over complex cellular behavior

- Proof-of-concept, toy circuit engineering
- Elementary network rewiring

- Complex synthetic circuit engineering
- Synthetic gene cluster or regulatory pathway refactoring

10^0 - 10^2 per cycle

10^2 - 10^4 per cycle

10^4 - ? per cycle