Balancing gene expression without library construction via a reusable sRNA pool

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\begin{abstract}
Balancing protein expression is critical when optimizing genetic systems. Typically, this requires library construction to vary the genetic parts controlling each gene, which can be expensive and time-consuming. Here, we develop sRNAs corresponding to 15nt ‘target’ sequences that can be inserted upstream of a gene. The targeted gene can be repressed from 1.6- to 87-fold by controlling sRNA expression using promoters of different strength. A pool is built where six sRNAs are placed under the control of 16 promoters that span a \( \sim 10^3 \)-fold range of strengths, yielding \( \sim 10^7 \) combinations. This pool can simultaneously optimize up to six genes in a system. This requires building only a single system-specific construct by placing a target sequence upstream of each gene and transforming it with the pre-built sRNA pool. The resulting library is screened and the top clone is sequenced to determine the promoter controlling each sRNA, from which the fold-repression of the genes can be inferred. The system is then rebuilt by rationally selecting parts that implement the optimal expression of each gene. We demonstrate the versatility of this approach by using the same pool to optimize a metabolic pathway (\( \beta \)-carotene) and genetic circuit (XNOR logic gate).
\end{abstract}

\begin{introduction}
Expressing proteins at the wrong levels is a common reason why genetic designs fail. For a metabolic pathway, suboptimal enzyme concentrations could lead to problems, such as the accumulation of toxic intermediates, titration of cofactors, or overburdening the host (1–5). Similarly, when building a genetic circuit, the expression of regulatory proteins needs to be balanced in order to perform the desired computational function (6,7). Across these projects, the challenge is that the correct expression levels are often not known \textit{a priori} and, even if they were, there is uncertainty in selecting genetic parts to achieve target expression levels (8,9). Further, the optimal expression of one gene often depends on the expression levels of other genes in the system, thus creating a ‘rugged’ search space where multiple genes have to be changed simultaneously to achieve an improvement (10–13). For these reasons, optimization often requires the creation of libraries by mutagenesis and screening.

Libraries of multi-gene systems can be built using guided and unguided approaches. Many variants of a pathway can be constructed by substituting genetic parts controlling each gene; for example, promoters, ribosome binding sites (RBSs), and RNA stability elements (1,5,14–19). Beyond part substitution, libraries can be built that alter the gene order, orientation, and operon occupancy (20–22). These approaches are often blind searches that require large libraries whose size is limited by assay throughput. To address this, mathematical modeling and combinatorial optimization algorithms have been used to guide the search towards a design objective, such as improved metabolic flux (23–28). In the absence of a model, the construction of pathway variants can be guided by multivariate statistical methods, including multifactor design of experiments and Bayesian approaches (12,29–32). At the other extreme, random approaches (e.g. mutagenesis and recombination) can be applied across the entire construct or targeted to particular regions (14,33–37). These approaches are low cost because they do not require sequence verification of each construct; however, they are limited by low mutagenesis rates that limit the simultaneous optimization of multiple genes (38). Further, the quality of the library needs to be monitored by sequencing to prevent nucleotide substitution biases and mutations as stop codons can cause a large fraction to be non-functional. Whether guided or unguided, the construction and sequence verification of the library is a slow and expensive step that has to be repeated for each system to be optimized.

We sought to develop a method where the expression levels of multiple genes could be simultaneously tuned without the need to rebuild a library for each system. Variation is achieved with a separate library, built once, that contains regulators expressed at different levels. Each regulator

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would control a different gene in the system so that changes in regulator expression lead to changes in the target gene. This approach requires regulators that are:

- **Orthogonal** so that each regulator only controls its target
- **Wide in dynamic range** to be able to sweep across expression levels,
- **Programmable** so that the regulator can be targeted to different genes and
- **Non-toxic** so that the regulators themselves do not influence the system.

We selected sRNAs (39–42) as they satisfy these criteria. sRNAs are potent negative regulators that do not require heterologous proteins to function (43). When expressed, sRNAs bind their cognate mRNA in the 5′ untranslated region (UTR) via Watson–Crick base pairing with assistance from Hfq, a ubiquitous RNA chaperone protein. Once bound, translation initiation is inhibited and RNA degradation by RNase E is accelerated (44,45). Most synthetic sRNAs studied to date have involved fusing new target sequences to natural sRNA scaffolds and minor scaffold modifications (41,46–48). The best-engineered sRNAs have been shown to achieve up to 85-fold repression without cross-reactions (41,46).

Since each RNA element targets a sequence specific to its cognate gene, the RNA must be designed ad hoc for each target and screening of multiple designs is often required (49,50). This is usually successful because of the programmability of nucleotide interactions, however, it requires optimization for each gene (48,51). We wanted to simplify this process so that sRNAs could be optimized once and then used to control arbitrary genes. To this end, we designed sRNA ‘target sequences’ that are placed in the 5′ untranslated region (5′-UTR) of the gene to be tuned (Figure 1A). Fixing the sRNA target site avoids the need to redesign the sRNA for each target gene. The sRNA is optimized to increase the fold-repression against this target and verified with multiple genes. This is repeated to create a set of six sRNA:target pairs that are orthogonal to each other and do not impact the host even when highly expressed. Further, it is demonstrated that the repression can be controlled in a graded manner by using constitutive promoters of different strengths to control the expression level of the sRNA (Figure 1B). All six sRNAs are used to build a combinatorial pool of ~10^7 differentially expressed sRNAs (Figure 1C). This pool is capable of tuning genetic systems with up to six different genes over a wide expression space (Figure 1D). As a proof of concept, we applied the same sRNA pool to tune very different systems: a metabolic pathway (β-carotene) and a genetic circuit (XNOR logic). This only requires building a single construct for each system, where the target sequence is placed upstream of each gene to be tuned. Once the optimal construct is identified, the reliance on sRNA expression can be removed by performing limited part substitution in order to recover the same expression levels discovered with the sRNA screen.

**MATERIALS AND METHODS**

**Strains and media**

*Escherichia coli* DH10B [F- merA Δ(mrr-hsdRMS-mcrBC) F80lacZΔM15 ΔlacX74 recA1 endA1 araD139 Δ(ara leu) 7697 galU galK rpsL mup ] (52) was used in all cloning procedures and experiments unless stated otherwise. *Escherichia coli* K-12 MG1655 [ F- k-12G rfb-50 rph- Δ(arabCBAD) Δ(lacI)] (53) was used for iron starvation assays. *Escherichia coli* K-12 JW4130 [ F- Δ(arad-DaraB)567 ΔlacZ4787::rbrB-3 ] rph- Δ(aradD-araB)568 Δhsq-722::kan, hsdR514] and BW25113 [ F- Δ(arad-DaraB)567 ΔlacZ4787::rbrB-3, rph- Δ(aradD-araB)568 hsdR514] from the KEIO collection (54) were used for hfq knockout experiments. MegaX DH10B T1 Electrocomp Cells (ThermoFisher, C640003) were used for sRNA pool construction. ‘Clonetegation’ as described by St-Pierre et al. (55) was used to integrate sRNA-A-tagged RFP at attP21. The following *Pseudomonas* and *Salmonella* strains were used: *Pseudomonas protegens* Pf-5 (ATCC BAA-477), and *Salmonella typhimurium* LT2 (ATCC 700720). Cells were grown in LB Miller broth (Difco, 9003-350) or SOC (SOB (Teknova, S0210) supplemented with 0.4% glucose) for cloning. Supplemented M9 minimal media (1x M9 Salts (Sigma-Aldrich, M6030), 2 mM MgSO₄, 100 μM CaCl₂, 0.4% glucose, 0.2% casamino acids, 340 mg/l thiamine) was used for knockdown assays and iron starvation assays. Chloramphenicol (34 μg/ml) (Alfa Aesar, AAB20841–14), kanamycin (50 μg/ml) (GoldBio, K-120-10) or spectinomycin sulfate (50 μg/ml) (MP Biomedicals LLC, 158993) was supplemented where appropriate. IPTG (Sigma-Aldrich, I6758) and aTC (37919) were used as inducers. Three fluorescence proteins, GFPmut3 (56), mRFP1 (57) and YFP (56), were used as reporters.

**sRNA repression assay**

Colonies were picked and inoculated into 300 μl LB with antibiotics and grown 16 h at 37°C and 900 RPM in a Multitron Pro incubator shaker (In Vitro Technologies). Three microilters of this overnight culture were then added to 147 μl of M9 media with antibiotics and grown for 3 h in an ELMi shaker (Elmi Ltd.) at 37°C and 1000 RPM, resulting in an OD₆₀₀ of 0.2–0.4. Twenty microilters of the cultures were diluted into 180 μl 1× phosphate buffered saline (PBS) with 2 mg/ml kanamycin. For sRNA optimization experiments, sRNAs were expressed using promoter BBa_J23119 and terminator BBa_B0010 from a pBR322 plasmid (pRNA) while the reporter (GFP or RFP) was expressed from a p15A plasmid using promoter BBa_J23101, RBS BBa_B0032, terminator BBa_B0012, and RiboJ, an insulating ribozyme (pREP, Supplementary Figure S1) (58).

**sRNA growth assay**

Plasmid pMAX, in which all sRNAs are expressed from the strongest promoters used in the pool (PA₁₆, PB₁₆, PC₁₆, PD₁₆, PE₁₆ and PF₁₆), was used to quantify growth defects from sRNA overexpression (Supplementary Figure S2 and Table S1). *Escherichia coli* DH10B cells harboring control plasmid pEMPTY or pMAX were grown for 18 h in LB,
normalized to an OD$_{600}$ of 1, then diluted 1:200 in LB and grown at 37°C for 3 h. Cultures were then diluted 1:200 again in LB into microtiter plates and growth was measured for 16 h. OD$_{600}$ measurements were performed on a Cary 50 Bio UV-Vis spectrophotometer (Varian). Cultures were diluted with sterile water until the OD$_{600}$ fell between 0.10 and 0.60. The same protocol was used for iron starvation growth assays with the exception that _E. coli_ MG1655 cells and supplemented M9 media without iron were used.

**Promoter strength measurements**

The strengths of all constitutive promoters were characterized by using them to drive YFP expression from a reporter plasmid (pPROM, Supplementary Figure S1). Reported promoter strengths are the geometric mean of the measured fluorescence output. The sRNA repression assay was followed with the exception that cells were grown in LB instead of M9 media. T7 promoter characterizations were performed by co-transforming the T7RNAp expression plasmid N249 (59) with the reporter plasmid. For these assays, colonies were grown overnight and diluted 1:200 into LB with antibiotics and 1 mM IPTG and grown for 6 h at 37°C, then analyzed using the flow cytometer.

**Flow cytometry**

Fluorescence was measured using a MACSQuant VYB (Miltenyi Biotec) with a 488-nm laser for GFP and YFP excitation and 561-nm for RFP excitation. For each sample at least 5 × 10⁴ events were recorded using a flow rate of 0.5 μl/s. FlowJo v10 (TreeStar Inc.) was used to analyze the data. All events were gated by forward scatter and side scatter. Fluorescence values are shown as the geometric mean. Fold-repression is calculated as the geometric mean of the fluorescence measured with an sRNA expression vector.

**Computational methods**

All Random DNA sequences were generated using a random DNA sequence generator (http://www.faculty.ucr.edu/~mmaduro/random.htm), with a GC content probability parameter of 0.5. All sRNA tags were evaluated against the _E. coli_ genome using BLASTn (http://www.ecogene.org/ecoblast) (60); sRNA tags exhibiting more than 11bp chromosomal homology or homology within 200 bp of annotated coding sequences on either strand were omitted. The search was performed on the _E. coli_ K-12 MG1655 U00096.3 genome database with an expect threshold of 10 and the BLOSUM62 matrix. Nupack (www.nupack.org) (61) was used to predict RNA secondary structures. The following parameters were used for all RNA folding simulations: Nucleic acid type: RNA; temperature: 37°C; number of strand species: 1; maximum complex size: 1; RNA energy parameters: ‘Mathews et al., 1999’; Dangle treatment: some. RBS sequences for the pathway and circuit were generated using the RBS Calculator (https://www.denovodna.com/software/) (62). For all calculations, the following parameters were used unless otherwise noted: pre-sequence: (insulating ribozyme sequence followed by an sRNA tag); protein coding sequence: (full coding sequence from start to stop codon); target translation initiation rate: 50 000; free energy model: version 2.0; organism: _E. coli_ str. K-12 substr. MG1655 (ACCTCCTTA). RBSs for BetI and HlyRII were designed with the Target Translation Initiation Rate set to 25 000.

**sRNA library construction and sorting**

All degenerate nucleotide libraries were built using circular polymerase extension cloning (CPEC) (63) with oligos...
containing N's (Integrated DNA Technologies) at appropriate positions. The CPEC product was column purified (DNA Clean & Concentrator-5, ZymoResearch) and transformed into cells harboring the appropriate reporter plasmid. After 1 h outgrowth in SOC, cells were diluted in to 5 ml LB with antibiotics and grown overnight at 37°C. The sRNA repression assay was performed with the following modifications: (i) 100 μl of culture was used instead of 3 μl, (ii) after 3 h of growth in M9 media cells were pelleted and resuspended in 2 ml 1× PBS with 1 mg/ml chloramphenicol. Cells were sorted on a FACS Aria II (BD Biosciences). Sorting gates were set to sort cells with lower GFP expression than that achieved by the previous sRNA scaffold design. Sorting was stopped after 5 × 10^4 cells were collected. Sorted cells were allowed to recover in SOC for 2 h then plated and allowed to form colonies. Colonies were then streaked out and individually assayed by flow cytometry.

sRNA pool construction

The CombiGEM (64) protocol was used to assemble the sRNA pool (pPOOL, Supplementary Figure S1) with the following modifications. (i) The AvrII restriction site was replaced with XbaI. (ii) Ligation products were transformed into MegaX E. coli DH10B T1 Electrocomp Cells and after 1 h outgrowth in SOC, cells were diluted in to 5 ml LB with antibiotics and grown overnight at 37°C; the following morning, the culture was miniprepped and the process was iterated to until all six sRNAs were cloned.

High-throughput sequencing

sRNA pool sequencing barcodes were PCR amplified using 100 ng of the pool as template. Barcode PCR products were sent to the Massachusetts General Hospital DNA Core where the samples were processed and run on a Mi-Seq producing ~200 000 reads. Custom software was written to search the raw sequencing FASTQ file against the list of all 96 sequencing barcodes (Supplementary Table S1) to count the number of occurrences of each corresponding promoter in the pool.

β-carotene production quantification

The β-carotene pathway construct (pCAROTENE, Supplementary Figure S1) was co-transformed with plasmid N249 (59). Cells were grown overnight, made electro-competent (65) and transformed with 100 ng of the sRNA pool and plated. Colonies harboring all three plasmids were grown in 1 ml LB with antibiotics at 30°C for 18 h. Cultures were then diluted to an OD_600 of 0.2 into 1 ml Terrific Broth (TekNova, T7060) with appropriate antibiotics and 1 mM IPTG and grown for 48 h in a Multitron Pro at 30°C in 96-well deep plates (USA Scientific, 1896-2000). Cells were harvested for quantification by centrifugation at 4000g for 10 min. They were then washed with 0.6 ml water, and re-centrifuged. Cell pellets were then resuspended in 0.5 ml acetone and incubated at 55°C for 30 min with frequent vortexing. Pigment extraction mixtures were then centrifuged at 4000g for 15 min. 0.2 ml of supernatant was transferred to a polypropylene 96-well plate (Greiner, 655201) for quantification. Samples were measured using a H1 Synergy plate reader (BioTek Instruments) at an absorbance wavelength of 470 nm. A standard curve (Supplementary Figure S3), using purified β-carotene (Sigma-Aldrich, 22040), was established to correlate absorbance to amount of β-carotene produced. The following equation was used: \( \text{titer} = \frac{120.6 \times \text{measured absorbance}}{\text{background absorbance}} \) . Background absorbance was measured from a strain harboring pEMPTY (Supplementary Figure S1) in place of the pathway construct. Samples from the top producing strains were further analyzed by liquid chromatography (1260 Infinity LC System, Agilent Technologies) using a reversed phase C18 column (Phenomenex, 00A-4462) to confirm absence of lycopene and other precursor metabolites and confirm sample purity (Supplementary Figure S3).

XNOR circuit assay and sorting

Cells harboring the XNOR circuit (pCIRCUIT) and output (pOUT) plasmids (Supplementary Figure S1) were grown overnight at 37°C in 1 ml LB then diluted 1:200 into 200 μl M9 with antibiotics and grown for 3 h at 37°C. The culture was then diluted 1:700 into 200 μl M9 with antibiotics and inducers and grown for 6 h. Twenty microliters of the cultures were diluted into 180 μl 1× phosphate buffered sulfate (PBS) with 2 mg/ml kanamycin and run on the cytometer. For the first input 1 mM IPTG was used for a ‘1’ input and no IPTG for a ‘0’ input. For the second input, 2 ng/ml aTC was used for a ‘1’ input and no aTC for a ‘0’ input. The geometric mean of fluorescence in each state was used as the output. For XNOR optimization, cells harboring the XNOR circuit and output plasmid were grown overnight, made electro-competent (65), and transformed with 100 ng of the sRNA pool. After 1 h of outgrowth in SOC, cells were diluted in 5 ml LB with antibiotics and grown overnight at 37°C. The following day the culture was diluted 1:200 into 5 ml M9 with antibiotics and grown for 3 h at 37°C. Induction was commenced at this point by diluting cultures 1:700 into 3.5 ml M9 with antibiotics and appropriate inducers (1 mM IPTG and/or 2 ng/ml aTC). After 6 h of growth, cells were pelleted and resuspended in 2 ml PBS supplemented with 1 mg/ml chloramphenicol and sorted. Sort gates were selected according to the desired output (YFP-high or YFP-low) for each input condition and were designed to include 5% of all cells. Sorting was stopped after 5 × 10^4 cells were collected. Sorted cells were allowed to recover in SOC for 2 h then diluted into 5 ml LB with antibiotics and grown overnight. The induction and sorting process was repeated for all input conditions. After the final sort, cells were allowed to recover in SOC for 2 h then plated and allowed to form colonies. Colonies were then streaked out and individually assayed by flow cytometry.

sRNA-mediated knockdown in other Gram-negative bacteria

The sRNA expression cassette from sRNA-A and from its corresponding reporter plasmid were cloned into a pBBR1 vector (66) (pBBR-RNA, Supplementary Figure S1). The plasmid was electroporated into Pseudomonas and...
Salmonella cells. Pseudomonas cells were subjected to the sRNA repression assay with the exception that cells were grown at 30°C in LB. Salmonella cells were subjected to the sRNA repression assay with the exception that cells were grown in LB. Fold-repression is calculated as the geometric mean of the fluorescence measured with a vector without the sRNA expression cassette (pBBR-EMPTY) divided by the geometric mean of the fluorescence measured with pBBR-RNA.

RT-qPCR
Probe-based RT-qPCR was performed on a LightCycler 480 (Roche) using PrimeTime Gene Expression Master Mix (Integrated DNA Technologies, 1055772). Custom PrimeTime qPCR assays with a 6-FAM dye and ZEN/IBFQ quencher were designed for each gene using the PrimerQuest Design Tool (https://www.idtdna.com/PrimerQuest/). The promoter measurement assay described above was followed. RNA was isolated immediately after growth using the Quick-RNA Fungal/Bacterial Micro-prep kit (ZymoResearch, R210). DNA was removed by DNase treatment with the DNA-free kit (ThermoFisher, AM1906). The ProtoScript First Strand cDNA Synthesis Kit (New England BioLabs, E6300S) was used to generate cDNA. The cysG, hcaT and idhT genes were used as reference genes based on literature (67).

RESULTS
Optimization of sRNA: target pairs
Our approach is based on the ability to design a target sequence that can be inserted upstream of any gene to make it sensitive to a cognate sRNA. This required first identifying the optimal region within the 5’-UTR to place the target. Then, the sRNA was optimized to maximally reduce target gene expression. Multiple parameters, including sRNA tag length (41), Hfq scaffold sequence, terminator strength, and 5’ standardization (68) and RNA stabilization (15) were optimized. Finally, a set of orthogonal sRNA:target pairs were created so that multiple genes can be simultaneously controlled.

To evaluate sRNA designs, we developed an assay involving two plasmids (Supplementary Figure S1). The first contains a constitutive promoter (J23119) driving the transcription of the sRNA. The second contains a constitutively expressed fluorescent protein in which an sRNA target sequence is inserted upstream of a strong RBS (B0032) (Materials and Methods). The plasmids are co-transformed and fluorescence quantified by flow cytometry (Materials and Methods). Fold-repression is reported by comparing the fluorescence measured with a vector with an empty vector.

We first identified the optimal region in the 5’-UTR to target with an sRNA. For our initial tests, ten 20nt sRNA tags with 50% GC content were generated using a random DNA sequence generator (Materials and Methods). Each tag was evaluated for sequence identity with the E. coli genome using EcoBlast (60) and a tag with no match to the genome was selected. This tag was fused to the wild-type MicC sRNA scaffold as described by Na et al. (41) (sRNA-AG1) while its cognate targeting sequence (reverse complement of tag) was inserted at multiple positions in the 5’-UTR of GFP and RFP (Supplementary Figure S4). We found that placing the target sequence closer to the RBS on the 5’ side improved gene repression while inserting it on the 3’ of the RBS resulted in reduced repression. These results corroborate similar studies and are consistent with the mechanism of sRNA-mediated gene repression (44,69).

We avoided directly targeting the RBS or coding sequences to ensure that the target can be reused to control different genes.

Once the optimal target position was identified, we tested different tags and target lengths to further improve repression. For E. coli, which has a genome size of ∼4.6 Mb, 12nt is the minimum number nucleotides necessary to ensure perfect orthogonality to the genome. Though targeting sequences short as 6nt have been observed in natural sRNAs, shorter target sequences can lead to sRNA promiscuity requiring very high sRNA expression, and off-target effects (45). We designed and tested tags and corresponding target sequences ranging from 12nt to 20nt in length. Repression efficiency was similar across all tag lengths (Supplementary Figure S4). As tag length decreases, RNA hybridization is less energetically favorable (41). However, as tag length increases the probability of alternate secondary conformations for the sRNA and target mRNA increases, which may reduce repression (70). Therefore, we standardized sRNA tag/target length at 15nt as a compromise between these constraints.

Next, we evaluated different Hfq scaffolds and sought to engineer one for strong repression. Six well-studied Hfq scaffolds from natural sRNAs in E. coli were fused to the 15nt tag described above and assayed: DsrA (71), GcvB (72), MicC (73), MicF (74), SgrS (75) and Spot42 (76). All tested sRNAs exhibited activity, of which the Spot42-derived sRNA was the strongest achieving 68-fold repression (Figure 2A).

Most natural sRNAs include a rho-independent terminator in their Hfq scaffolds as it is essential for Hfq recruitment (77,78). Prior studies have shown that this terminator can be exchanged for other rho-independent terminators without abrogating sRNA-mediated repression (47,79). Since the secondary structure of sRNAs is essential for Hfq-binding (80), and consequently gene repression, we hypothesized that replacing the natural terminator found in the Spot42 Hfq scaffold with a stronger terminator would improve repression efficiency as this would result in fewer sRNAs with extended 3’ ends that could interfere with sRNA folding. The best, T(BF1176), was selected as the core scaffold for further optimization (Figure 2C).

A prior structural study showed that all but six nucleotides (not including the terminator stem-loop) of the Spot42 scaffold are involved in Hfq-binding, and consequentially essential for sRNA-mediated repression (81). Therefore, we created a targeted mutagenesis library in which these six nucleotides were randomized resulting in a 4096-variant library (Figure 2C, arrows). Potent scaffold variants were identified by fluorescence-activated cell sorting (FACS) (Materials and Methods). The best one (sRNA-AG4) yields 115-fold repression with all positions mutated (Figure 2A). Folding predictions of these high-
cleotides are essential for Hfq-binding (81). Red nucleotides indicate the terminator stem loop of terminator BF1176. Arrows indicate nucleotides that were randomized to screen for improved folding predictions after adding the 5′ overhang to the sRNA and found that 10/16 disrupted folding. Ribozymes were chosen to standardize the 5′ end of the sRNA. Using a hammerhead ribozyme (HHR) scaffold (83), we designed a random library in which 18 non-essential nucleotides of the ribozyme (Figure 2D, purple nucleotides with arrows) were replaced with degenerate bases resulting in a theoretical library of $10^{10}$ variants fused to sRNA-AG4. This library was then screened using FACS. Approximately 3% of the screened variants exhibited improved repression from which the top-performing HHR/sRNA design (sRNA-A), capable of 175-fold repression, was discovered. The repression achieved with this variant was found to strongly correlate with promoter strength (Figure 2B, lower panel).

We then created a set of sRNAs that target different tags to enable the control of multiple genes within a cell. Redesigning the sRNA scaffold for a new target simply requires changing the 15nt tag on the sRNA and inserting the reverse complement of the cognate target sequence preceding the Shine Delgarno sequence on the gene to be controlled. A bioinformatic approach was pursued to generate a list of tags that avoid off-target effects in E. coli (Materials and Methods). In addition, secondary structure predictions were performed to ensure that the addition of the target sequence to the sRNA would not disrupt folding of the Hfq-binding region. From this list, 25 tags were selected, fused to sRNA-A, HHR appropriately modified for each, and tested for repression against corresponding reporter constructs. Those were eliminated that exhibited toxicity (9), crosstalk (3), or produced less than 100-fold repression (7). This led to a set of 6 sRNAs that efficiently target different sequences without impacting the host (Supplementary Table S2).

The optimized sRNA scaffold (without the tag) is 55nt performing scaffold variants revealed that the predicted secondary structure of the Spot42 Hfq-binding region was preserved (Materials and Methods). As expected, this repression is Hfq-dependent (Supplementary Figure S5).

A challenge with using promoters to control sRNA expression is that they can generate a distribution of transcription start sites (14,82). As a result, the initial attempt to control gene expression using different strength promoters produced a poor correlation between promoter strength and repression (Figure 2B, top panel). Using the published transcription start sites of the promoters (14), we performed folding predictions after adding the 5′ overhang to the sRNA and found that 10/16 disrupted folding. Ribozymes were chosen to standardize the 5′ end of the sRNA. Using a hammerhead ribozyme (HHR) scaffold (83), we designed a random library in which 18 non-essential nucleotides of the ribozyme were replaced with degenerate bases resulting in a theoretical library of $10^{10}$ variants fused to sRNA-AG4. This library was then screened using FACS. Approximately 3% of the screened variants exhibited improved repression from which the top-performing HHR/sRNA design (sRNA-A), capable of 175-fold repression, was discovered. The repression achieved with this variant was found to strongly correlate with promoter strength (Figure 2B, lower panel).

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The optimized sRNA scaffold (without the tag) is 55nt performing scaffold variants revealed that the predicted secondary structure of the Spot42 Hfq-binding region was preserved (Materials and Methods). As expected, this repression is Hfq-dependent (Supplementary Figure S5).
pREP was integrated at site attP21 in *E. coli* DH10B cells (Supplementary Figure S1 and Materials and Methods). Once constructed, the reporter strain was tested against a wide range of RNA levels spanning 18–21,000 au from promoters P*PA1*-P*PA16* with the same expression plasmids used in Figure 2B (Supplementary Table S1). Even though the sRNA was characterized with a target expressed from a multi-copy plasmid (pREP and Figure 2B), target repression on the genome, constituting a drastic change in target copy number, was predictable. However, since the starting level of the reporter was low, repression saturates as reporter copy number, was predictable. However, since the starting level of the reporter was low, repression saturates as reporter levels approach background (Figure 2F).

The sRNAs were then evaluated in other Gram-negative bacteria where Hfq is conserved (81) (Figure 2G). A single broad host range plasmid was constructed, from which sRNA-A and the corresponding GFP reporter gene are expressed (pBBR-RNA). The same vector without the sRNA expression cassette was also constructed and used for fold repression calculations (pBBR-EMPTY). Though the parts in pBBR-RNA are native to *E. coli*, they were previously evaluated and observed to exhibit activity in *Pseudomonas protegens* PF-5 and *Salmonella typhimurium* LT2. The sRNA repression assay was performed for both *Pseudomonas* and *Salmonella* cells harboring pBBR-RNA (Materials and Methods). sRNA activity was observed in both organisms (Figure 2G).

### Combinatorial sRNA pool construction and validation

A library was built that varies the expression levels of each of the sRNAs. Ninety-six promoters (16 different promoters for each of 6 sRNAs) spanning strengths of 18–40 000 au were gleaned from a published set (Supplemental Table S1) (8). Each of the 96 promoter–sRNA combinations were evaluated for repression of both GFP and RFP to capture any variability due to the target gene sequence (Figure 3A and Supplementary Figure S7). For each sRNA, there is a strong correlation between promoter strength and repression (Figure 3A). Intermediate promoter strengths lead to a graded response. There is only a slight reduction in growth rate when all six sRNAs are maximally expressed (Supplementary Figure S2 and Materials and Methods).

The next step was to build the sRNA pool by randomly combining the promoter–sRNA pairs so that each member of the pool expresses the six sRNAs at different levels. The sRNA pool was constructed using CombiGEM assembly (64). Each of the 96 sRNA assembly plasmids contain strategically placed restriction sites that encompass a 10nt sequencing barcode associated with a specific promoter/sRNA pair, unique constitutive promoter, sRNA, and a strong terminator (85) (Figure 3B, Supplementary Table S1). The assembly reactions are then performed, the result of which is a barcode string that can be used to determine the promoter identity at each sRNA position with a 96 bp sequencing read (Figure 3C, Materials and Methods). Deep sequencing of the final pool resulted in the recovery of 80/96 promoters from ~200 000 reads and an even distribution of promoter strengths (Figure 3D).

All of the sRNAs are oriented in the same direction on the construct and thus could be influenced by transcriptional readthrough. To evaluate this effect, we built constructs that permuted the order of the sRNAs. No effect of ordering the sRNA genes on repression was observed, likely due to the use of strong terminators (Supplementary Figure S8).

### Application to a metabolic pathway: optimization of β-carotene

The β-carotene pathway was selected as a model system to test our sRNA pool because it is known that enzyme balancing is important to increase titer (5,37,86–88). A single construct was built that includes the four-gene pathway to β-carotene (crtE, crtB, crtI, crtY) as well as the *E. coli* genes *dxs* and *idi*, which have been shown to improve titers when overexpressed (Figure 4A and B) (37,89). Each enzyme gene was placed under the control of a strong T7 promoter and is preceded by an insulating ribozyme, strong RBS designed for that gene (Materials and Methods), and sRNA target sequence (A–F) (Figure 4B and Supplementary Table S3). A unique strong terminator was placed after each gene. Transcription is induced from the promoters by expressing T7 RNA polymerase (RNAP) from a separate plasmid under IPTG-inducible control (59).

The sRNA pool was transformed into a strain harboring the pathway, generating a library of variants. More than 1200 colonies were picked, induced with 1 mM IPTG, and assayed for β-carotene production (Materials and Methods). Approximately half of the variants improved yield
Figure 3. Pool assembly from differentially expressed sRNAs. (A) Repression response for all six sRNAs against GFP (circles) and RFP (triangles) genes containing targets in their 5'-UTR. Each sRNA is expressed from a set of 16 promoters unique to each sRNA. Sequences of all promoters are provided in Supplementary Table S1. Error bars correspond to the standard deviation of three experiments performed on different days. (B) Example of pool assembly plasmids with restriction sites for CombiGEM assembly. The barcode symbol depicts a unique 10nt sequencing barcode corresponding to each promoter in the pool. Sequences for promoters and barcodes are provided in Supplementary Table S1. (C) Iterative assembly example of a six sRNA pool variant. (D) Deep sequencing results for promoter coverage in the assembled pool containing six RNAs (1% pool coverage with ~200000 reads); *' signifies that the promoter was not detected. In the promoter names, X refers to the promoter number above the horizontal line.

Application to a genetic circuit

A genetic circuit was chosen to demonstrate that the same pool could be applied to the optimization of a very different type of system. A two-input XNOR circuit was constructed using repressor-based NOT and NOR gates (Figure 5). Previously, we have demonstrated that we can computationally match the response functions of the gates to build this circuit (7). However, as a proof-of-principle, we took a different approach here, where strong RBSs were used to control the expression of each repressor (Materials and Methods).

The sRNA target sequences (targets A–D) were placed between the ribozyme insulators and the RBS of each gene. As would be expected, this ‘blindly’ constructed XNOR circuit does not produce the correct logic operation (Figure 5D).

Because the circuit output is a fluorescent protein, fluorescence-activated cell sorting (FACS) can be used to rapidly screen for improvements. The sRNA pool was co-transformed with the circuit construct and then the library was serially sorted for correct output under the different combination of inputs (1 mM IPTG and 2 ng/ml aTc) (Materials and Methods). After sorting, 96 colonies were individually assayed for the XNOR function, and 21 exhibited proper functionality. The best performing circuit exhibited the correct XNOR logic (Figure 5F). The corresponding sRNA construct was sequenced to determine the barcodes that identify the promoters controlling each sRNA, from which the fold-repression could be inferred (Figure 5G).

While the circuit now works, it is not ideal to have to continue to transcribe the sRNAs for proper function. In contrast to the β-carotene pathway, the circuit’s promoters cannot easily be changed to different strengths because they contain operators that bind to the repressors. Therefore, RBS substitutions were used to recapitulate the opt-
DISCUSSION

We developed a method to tune genetic systems by exploring large gene expression spaces without having to build ad hoc libraries. This method can be used to tune a variety of multi-gene systems, and presents a novel approach to genetic system optimization by using trans-acting RNA regulators that target standardized target sequences. Further, once a functional variant is discovered from the sRNA pool, this can be used to determine the optimal gene expression levels, which can then be hard-coded by selecting corresponding promoters or RBSs to build the final construct. This speaks to the value of having libraries of standardized genetic parts that have been sufficiently characterized such
that they can be rationally selected to achieve a target design objective.

One limitation of our approach is that genes can only be repressed, as opposed to being up-regulated. This constrains the expression space to the maximum level of the initial promoters. This can be problematic if any of the genes are toxic. This was averted in the β-carotene pathway by using T7 promoters and the expression of T7 RNAP on a separate plasmid. This avoids the problem of toxicity because the genes will not be expressed until T7 RNAP is induced. Up-regulating sRNAs have been discovered (91, 92) and may be amenable to a tuning pool; however, systematic characterization and engineering of these sRNAs is still required.

The reusable sRNA pool presented in this work establishes a new paradigm for genetic system optimization. Our approach is to build a library of regulators once and then use this same library to rapidly optimize different systems. Combinatorial optimization in genetic engineering has proven very successful, but the corresponding libraries are expensive, technically difficult to build, and require extensive sequence verification to ensure quality. It is particularly challenging if the constructs are large, reducing the efficiency of transformation, or have to be inserted chromosomally. Our approach only requires that the construct be built—including the sRNA target sequences—and introduced into the cell once. This one construct can be co-transformed with the plasmid-borne pool and screened for the optimal construct. It is sufficiently simple to be routinely or systematically applied when building constructs for new pathways, for example when mining pathways from metagenomic datasets. Often, the necessary expression levels in a heterologous host are unknown and the pooled approach allows many “shots on goal” to be taken rather than risking the selection of potentially suboptimal parts for the control of each gene. Further, this method can be applied simply, without the need for DNA construction robotics and automation platforms.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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