Synthetic biology aims to provide an engineering-driven approach towards building biological entities with complex and novel functionality using well-characterized modular parts. Precise and programmable control of gene expression is becoming increasingly important for many synthetic biology applications as well as biotechnology in general. The ability to control the expression of multiple genes, for example, will aid in the optimization of biosynthetic pathways for industrial chemical production while maximizing productivity and minimizing host toxicity. Over the years, synthetic biology approaches have yielded increasingly sophisticated means of controlling gene expression, including synchronized oscillators, logic gates, memory devices, analog signal processors and state machines. However, many of the regulatory elements in previous work overlap or are incompatible with each other, thereby limiting the integration of such diverse components in more complex circuits.

A basic requirement for engineering complex systems is a large repertoire of regulators that are modular, programmable, homogeneous, predictable and easy to compose. The idiosyncratic nature of many protein regulators presents challenges for their use in circuits, but is being addressed using insulation, computer-aided design strategies and orthogonal proteins and enzymes. RNA molecules provide an alternative to proteins for constructing genetic circuits with excellent programmability and composability due to their predictable base-pairing rules and well-characterized thermodynamics. An assortment of RNA-based regulators have been developed using interaction strategies from nature, with the hok/sok, pT181 and IS10 antisense systems being used for translational activation, transcriptional attenuation and translational repression, respectively.

To expand the potential of RNA-based regulation, we have recently harnessed de novo RNA design to develop toehold switch riboregulators that detect trigger RNAs with virtually arbitrary sequences and repress gene expression by up to 300-fold and yield orthogonal sets of up to 15 devices. Automated forward engineering is used to improve toehold repressor dynamic range and SHAPE-Seq is applied to confirm the designed switching mechanism of 3WJ repressors in living cells. We integrate the modular repressors into biological circuits that execute universal NAND and NOR logic and evaluate the four-input expression NOT ((A1 AND A2) OR (B1 AND B2)) in Escherichia coli. These capabilities make toehold and 3WJ repressors valuable new tools for biotechnological applications.
can decrease gene expression in excess of 100-fold, a substantial improvement over previous RNA-based translational repressors and comparable to protein repressors. Thermodynamics-based forward engineering is used to enhance the performance of the toehold repressors. In-cell SHAPE-Seq is used to directly confirm the formation of 3WJ structures in the 3WJ repressors. Validated repressors are integrated into ribocomputing devices to achieve NOR and NAND logic with up to four sequence-independent input RNAs, providing universal building blocks for logical computation.

Results
Design of synthetic translational repressors. Previously, we developed toehold switches that inhibit translation using a hairpin secondary structure that sequesters the ribosome binding site (RBS) and start codon within a hairpin loop and stem, respectively (Fig. 1a). A single-stranded toehold domain \( a^* \) at the 5’ end of the switch RNA hairpin provides the initial binding site for a single-stranded trigger RNA strand, which has a complementary domain \( a \). On binding of the cognate trigger molecule to the switch hairpin and completion of a toehold-mediated branch migration process, the RBS and start codon are available for ribosomal access and translation of the downstream gene. The lack of sequence constraints in designing trigger RNA molecules greatly expands the orthogonality of toehold switches and the use of thermodynamically and kinetically favorable toehold-mediated interactions provides wide dynamic range.

We sought to obtain a library of programmable, wide-dynamic-range translational repressors analogous to the toehold switches and devised two types of repressor inspired by the design principles of these earlier riboregulators. The first repressor employs a switch RNA with a 5’ toehold domain and is referred to as a toehold repressor (Fig. 1b; see Methods and Supplementary Fig. 1a for details). The 15-nt toehold domain of the switch RNA is followed by a hairpin structure and a single-stranded expression region containing an RBS, start codon and the coding sequence of the output gene. Without the trigger RNA, the exposed RBS and start codon enable active translation of the output gene. The trigger RNA of the toehold repressors is a 45-nt single-stranded RNA sequence that is complementary to the toehold and stem of the switch RNA.

After binding of the trigger to the switch RNA toehold, the ensuing
branch migration process unwinds the hairpin stem and releases the domains b' and c'. Domain b' is complementary to the sequences upstream and downstream of the start codon, and thus forms a hairpin structure with these domains. This newly formed hairpin recapitulates the repressed structure of the toehold switch and thus represses translation upon trigger binding. The toehold repressor trigger sequence does not possess bases complementary to the RBS or the start codon, which allows an arbitrary choice of potential trigger sequences. If a trigger RNA sequence leads to in-frame stop codons in the expression region, bulges can be introduced or shifted in the b' domain of the switch RNA hairpin to compensate.

The second repressor adopts a 3WJ structure to suppress translation and is referred to as a 3WJ repressor (Fig. 1c; see Methods and Supplementary Fig. 1b for details). Here, the switch RNA employs an unstable hairpin secondary structure that contains an RBS in the loop region and a start codon in the stem region. Despite its high secondary structure, this unstable hairpin was previously demonstrated to be translationally active in toehold switch mRNA sensors11. On either side of the unstable hairpin are single-stranded domains a* and b *. We hypothesized that transient formation of the bottom stem domain of the hairpin would co-localize these two domains to provide an effective binding site for a complementary trigger RNA. To take advantage of this design feature and improve repressor orthogonality, we designed cognate triggers where domain b is mostly contained in a hairpin secondary structure and a toehold composed of domain a and part of domain b is located at the 3' end. When the trigger RNA is expressed, the toehold binds to the a* domain and part of the b* domain of the switch RNA. The switch RNA b' domain then completes a branch migration to unwind the trigger RNA stem. The resulting trigger–switch complex has a stable 3WJ structure that effectively sequesters the RBS and start codon within the loop and stem of the switch RNA, respectively, and strongly represses translation. Despite the use of a trigger with a hairpin structure to improve device orthogonality, the 3WJ repressors can also detect nearly arbitrary trigger RNAs provided that the trigger RNA sequence does not lead to an in-frame stop codon in domain b'.

In silico design of repressors and in vivo validation. We generated libraries of both translational repressors de novo using the NUPACK sequence design package12 (see Methods for details). A total of 44 toehold repressors and 48 3WJ repressors were designed and validated in vivo (see Supplementary Tables 1–3 for sequence information). Members of the toehold repressor library were selected to reduce the potential for a non-cognate trigger RNA to disrupt the switch RNA stem. Members of the 3WJ repressor library were selected to minimize the potential for the non-cognate trigger RNAs to interact with the switch RNA. The Escherichia coli BL21 Star DE3 strain with an IPTG-inducible genomic T7 RNA polymerase and decreased RNase activity was used for repressor characterization. A medium-copy plasmid containing the switch RNA regulating GFP and a high-copy plasmid encoding the trigger RNA were transformed into E. coli. For measurements in the absence of a cognate trigger, a non-cognate RNA strand with high secondary structure was transcribed from the high-copy plasmid.

Figure 1d shows the fold reduction of GFP fluorescence observed for the toehold repressor library. The GFP fold reduction was measured from the geometric mean fluorescence of GFP obtained from flow cytometry for cells in the on state expressing the non-cognate trigger RNA and in the repressed off state expressing a cognate trigger RNA (see Supplementary Fig. 2a for on- and off-state GFP expression levels). Cell autofluorescence was not subtracted from either the on- or off-state fluorescence for determination of the GFP fold reduction. Although the toehold repressor devices show wide variations in performance, 48% (or 21 out of 44) provide at least a 10-fold change in gene expression upon detection of the trigger RNA. Five devices or 11% exhibit a GFP fold reduction of at least 50-fold, corresponding to over 98% repression of GFP signal. The 3WJ repressors overall provided improved performance compared to the toehold repressors (Fig. 1e). A substantially higher fraction of these devices at 71% (or 34 out of 48) provided at least 10-fold reduction of GFP expression, while a smaller fraction (8%, or 4 out of 48) yielded exceptionally high 50-fold or more reduction in GFP (see Supplementary Fig. 2b for the on- and off-state GFP expression levels).

We also tested the repressors in a variety of different conditions to determine their effects on performance (see Methods for details). Although library screening was conducted in BL21 Star DE3, an RNase-deficient strain, we found that both types of repressor provided greater than 20-fold reduction of GFP in E. coli BL21 DE3 cells with wild-type RNase levels (Supplementary Fig. 3). We observed that significant repression occurs within an hour of induction and that the fold reduction of both repressors increased over time (Supplementary Fig. 4). The repressors also functioned well when transcribed using the endogenous E. coli RNA polymerase with inducible promoters in E. coli MG1655/Marionette-Wild12 and provided stronger repression as the trigger RNA concentration increased (see Supplementary Fig. 5 and Supplementary Table 4 for sequence information). In cell-free in vitro translation reactions, the systems achieved 10-fold reductions in expression when supplied with as low as a twofold excess of the trigger RNA over the switch RNA (Supplementary Fig. 6). Variant 3WJ repressors designed with different stem sequences also operated successfully in E. coli when their secondary structures were sufficiently weak to allow translation to occur (see Supplementary Fig. 7 and Supplementary Table 5 for sequence information). Finally, we found that the dynamic range of the 3WJ repressors could be increased by roughly an order of magnitude using a faster-degrading GFP variant or decreased 1.2- to 1.7-fold using a more stable GFP (Supplementary Fig. 8).

Automated forward engineering of toehold repressors. To generate higher-performance toehold repressors, we implemented an automated strategy for ranking putative riboregulator devices. We first compiled a set of 114 thermodynamic parameters that could be computed rapidly from the sequence information of the trigger and switch RNAs (see Supplementary Note and Supplementary Figs. 9 and 10 for details). This set of thermodynamic parameters and experimental GFP fluorescence data from the toehold repressor library were then used in linear regressions to generate a scoring function for the devices (Supplementary Fig. 10).

NUPACK was used to generate an additional set of 265 toehold repressor sequences using identical secondary structures and design parameters as the first-generation library. The scoring function was applied to rank each of the devices and select the top 96 for a second-generation library (see Supplementary Table 6 for sequence information). Figure 2a presents the fold reduction of GFP fluorescence for the second-generation toehold repressors (see Supplementary Fig. 2c for the on- and off-state GFP expression levels). There is a dramatic increase in GFP fold reduction for the devices in general, with eight switches exhibiting a dynamic range greater than 100 and 81 switches exhibiting a dynamic range greater than 10. The second-generation systems exhibit an average GFP fold reduction of 40 compared to 20 for the first-generation library. Highly performing toehold repressors exhibit fold changes rivaling the dynamic range of protein-based regulators12 without requiring any in vitro evolution or large-scale screening experiments. We quantified the effectiveness of our selection criteria by calculating the percentage of toehold repressors with GFP fold reductions exceeding a given minimal level (Fig. 2b). The yield of high-performance devices is higher for the second-generation devices across all fold reductions.

SHAPE-Seq measurements of 3WJ repressor structure. To better understand the operating mechanism of the synthetic repressors, we performed in-cell SHAPE-Seq16 on devices with varying repression
also showed the formation of the 3WJ structure upon repression (Supplementary Fig. 11). To the best of our knowledge, these results represent the first structural confirmation of the regulatory mechanism of a completely de novo-designed riboregulator.

Evaluation of repressor orthogonality. One of the prerequisites for higher-order logic processing is the orthogonality of regulatory components with respect to one another. We thus measured in vivo the interactions between pairwise combinations of different repressor trigger and switch RNAs. For the second-generation toehold repressors, we performed in silico screening to identify 16 devices that provided more than 10-fold GFP reduction and also displayed low levels of predicted crosstalk with non-cognate triggers. Flow cytometry was used to quantify GFP output in *E. coli* for all 256 trigger–switch interactions (Fig. 4a). Crosstalk was calculated by dividing the GFP fluorescence obtained from a non-cognate trigger–switch pair, as shown in Fig. 4a. However, the crosstalk level was high in many instances, such that the set of orthogonal devices that maintained at least 12-fold dynamic range was reduced to four devices (red boxes, Fig. 4a). For the less stringent orthogonality condition of at least sevenfold dynamic range, the toehold repressor library yielded a set of eight independent riboregulators (blue boxes, Fig. 4a).

Based on the shorter exposed single-stranded regions of the 3WJ repressor trigger RNAs, we anticipated that these repressors would show improved orthogonality compared to the toehold repressors. We measured the pairwise trigger–switch interactions for 16 of the top devices using the same methods (Fig. 4b). The 3WJ repressors showed substantially reduced crosstalk while maintaining strong repression of cognate trigger–switch pairs (see Supplementary Fig. 12 for GFP expression levels). In fact, we found that a set of 15 3WJ repressors provided at least 17-fold reductions in GFP expression in the presence of the cognate trigger compared to any of the other 14 non-cognate triggers. Moreover, we only observed substantial crosstalk in a single pairwise interaction (red outlined box, Fig. 4b).

To quantify device orthogonality, we determined the maximum number of repressors that could be used to provide a given minimum level of overall dynamic range (Fig. 4c; see Supplementary Table 8 for
3 biologically independent samples. GFP fluorescence was measured from the dynamic range increase over the 3–5 h induction time. Crosstalk was determined by dividing the arithmetic mean of the GFP fluorescence from a given trigger–switch combination by the arithmetic mean of the GFP fluorescence for the cognate trigger–switch interaction. GFP fluorescence was measured from the overall library dynamic range and orthogonal library size for the toehold repressors and 3WJ repressors. 3WJ repressor orthogonal library size and crosstalk measured by flow cytometry for 256 trigger–switch combinations 3 h after induction. All but one of the 240 non-cognate combinations provide at least 14-fold higher repression efficiency.

Assessment of toehold and 3WJ repressor orthogonality.

Fig. 3 | In-cell SHAPE-Seq confirmation of the 3WJ repressor mechanism. a, Design schematic for testing 3WJ repressor variants. A 3WJ repressor switch was characterized using in-cell SHAPE-Seq, either expressed alone or co-expressed with a trigger RNA. Several triggers were tested, varying in their designed binding length (ab) to either side of the switch hairpin. Functional characterization of switch plasmid expressed without trigger (green) and with triggers of increasing interaction length (blue). Strong repression is observed upon trigger binding, with longer triggers showing increased repression efficiency. c, In-cell SHAPE-Seq reactivity profile of the switch RNA expressed alone. A trend of high reactivities is observed across the molecule, consistent with the design hypothesis that the switch hairpin can be disrupted by ribosome binding, leading to active translation. d, In-cell SHAPE-Seq reactivity profile of the switch co-expressed with trigger RNAs. Sharp drops in reactivity are observed at the predicted trigger binding sites (a–a* and b–b*) and within the switch hairpin, suggesting the formation of a stable 3WJ structure when the trigger is bound. The RBS and start codon (AUG) positions are indicated. Fluorescence (b) and reactivity (c, d) values are the arithmetic mean of n = 3 biologically independent samples. Error bars represent the s.d. from n = 3 biologically independent samples and individual points are shown for the fluorescence and reactivity of each sample.

Fig. 4 | Assessment of toehold and 3WJ repressor orthogonality. a, Toehold repressor crosstalk measured by flow cytometry for 256 trigger–switch combinations 3 h after induction. Red outlined boxes designate a subset of four repressors that exhibit sufficiently low crosstalk to provide at least 12-fold GFP reduction. Blue boxes designate a subset of eight repressors that provide at least sevenfold GFP reduction. b, 3WJ repressor crosstalk measured by flow cytometry for 256 trigger–switch combinations 3 h after induction. All but one of the 240 non-cognate combinations provide at least 14-fold higher GFP expression than cognate pairs. The red outlined box marks the trigger–switch combination with the most substantial crosstalk. c, Comparison of overall library dynamic range and orthogonal library size for the toehold repressors and 3WJ repressors. 3WJ repressor orthogonal library size and dynamic range increase over the 3–5 h induction time. Crosstalk was determined by dividing the arithmetic mean of the GFP fluorescence from a given trigger–switch pair by the arithmetic mean of the GFP fluorescence for the cognate trigger–switch interaction. GFP fluorescence was measured from n = 3 biologically independent samples.
the orthogonal repressor sets). This analysis showed large improvements in orthogonal library size and dynamic range for the 3WJ repressors compared to the toehold repressors. For example, the most orthogonal eight-device toehold repressor set provided an overall dynamic range of at least sevenfold, while the corresponding eight-device 3WJ repressor set yielded an overall dynamic range of 29-fold. As the induction time increased, we also observed steady increases in the fold reduction of GFP in the cells, leading to parallel increases in device orthogonality for the 3WJ repressors. For example, a set of six 3WJ repressors provided a remarkable library dynamic range of 118 at the 5th time point. Based on the orthogonality of the repressors, we also investigated their ability to respond to intracellular mRNAs. These mRNA-sensing toehold and 3WJ repressors, which were designed to bind to regions of low secondary structure in the trigger mRNAs, successfully detected multiple antibiotic resistance genes using GFP and mCherry reporter proteins (Supplementary Fig. 13; see Supplementary Table 9 for sequence information).

**Two-input repressor-based logic circuitry.** The modular and programmable nature of the toehold and 3WJ repressors makes them ideal candidates for integration into ribocomputing devices for implementing sophisticated genetic programs. We have previously demonstrated that toehold switch riboregulators can be incorporated into such RNA-based computing systems for multi-input intracellular computation using RNA input signals and protein output signals. We thus applied the ribocomputing strategy to the repressors to enable efficient computation of NAND and NOR logic functions in living cells (see Methods and Supplementary Figs. 14 and 15 for circuit design details and Supplementary Table 10 for circuit sequence information).

We studied two-input NAND gates based on toehold repressors optimized for these logic operations. The trigger RNA sequence was divided into two input RNAs A1 and A2, and complementary bridging domains u and u* were appended to each input (Fig. 5a and Supplementary Fig. 14a). Only when both inputs A1 and A2 are present do they hybridize to one another through the u–u* interaction and bring both trigger halves into close proximity for binding to the gate RNA, which consists of a single switch RNA hairpin upstream of an output gene. Similar associative toehold mechanisms have been demonstrated in vitro and have been used for AND logic in ribocomputing devices. Figure 5b shows the mean GFP fluorescence from this gate RNA showed a substantial reduction in fluorescence compared to the input conditions to compute the fold reductions for the circuit (Fig. 16 for GFP population histograms of all ribocomputing circuits). We found that the dynamic range of the repressor-based ribocomputing devices increased with IPTG induction time up to ~6 h (see Methods and Supplementary Fig. 17). For the logical TRUE input conditions with at least one input missing, GFP output from the gate RNA remained high. When the logical FALSE condition occurred with two input RNAs expressed, the NAND gate provided strongly reduced GFP expression. Mean GFP fluorescence for the null input condition with no cognate input RNAs expressed was divided by the mean GFP fluorescence obtained from each of the input conditions to compute the fold reductions for the circuit (Fig. 5c). GFP was reduced by 40-fold in the logical FALSE state. A noticeable decrease of 2.5-fold GFP reduction was observed upon expression of input A2 alone, potentially due to A2 binding causing a partial disruption of the gate RNA stem or cross-interactions between the u* bridging domain with exposed single-stranded regions of the gate.

We also implemented repressor-based gate RNAs integrating multiple repressor hairpin modules upstream of the output gene. Attempts using toehold repressor hairpins proved unsuccessful because their strong hairpin secondary structure and long target RNA binding sites both prevented efficient translation of the output gene by impeding translation from upstream RBS regions. However, the comparatively weak secondary structure of the 3WJ repressors and their short trigger RNA binding sites were ideal for incorporation into gate RNAs. We implemented a two-input NAND gate RNA composed of two orthogonal 3WJ repressor hairpins separated by a 17-nt single-stranded spacer domain (Fig. 5d). With only one input RNA present, translation of the output gene will continue from the unrepressed hairpin module, because the ribosome can translate through weak hairpin secondary structures and duplexes formed by the input and gate RNAs. As a result, only simultaneous binding of both input RNAs to the gate RNA will fully inhibit gene expression. We evaluated the two-input NAND gate and found that GFP expression remained strong except for the logical FALSE case with both inputs expressed (Fig. 5e). Small decreases in GFP expression were observed when only one of the input RNAs was present, probably as a result of inhibition of one of the two translation initiation sites from the gate RNA. The GFP fold reductions of the circuit show a large 88-fold decrease in expression in response to the two input RNAs compared to the null input case (Fig. 5f). This reduction was at least a factor of 33 higher than any of the changes in expression observed for single-input cases.

To implement NOR ribocomputing devices responsive to sequence-independent input RNAs, we developed a gate RNA architecture that exploited co-localized intramolecular interactions (Fig. 5g and Supplementary Fig. 15a). These NOR gate RNAs contain multiple sequestered trigger RNA sequences upstream of a 3WJ repressor module regulating the output gene. The trigger RNA domains x and y, which are complementary to the downstream repressor hairpin, are confined within the loops of strong hairpin secondary structures. These hairpins function as input RNA sensors that provide toehold domains for binding to complementary input RNA sequences. When an input RNA is expressed, binding to the input sensor leads to a branch migration that unwinds the sensor stem. This interaction releases the trigger RNA domain and enables the trigger to repress the downstream 3WJ repressor domain through an efficient gate RNA intramolecular interaction. We constructed the two-input NOR gate RNA using a validated 3WJ repressor and two input sensor hairpins, resulting in a gate RNA regulatory region of 312 nt. Measurements of GFP fluorescence from this gate RNA showed a substantial reduction in fluorescence upon expression of any of the cognate input RNAs (Fig. 5b). Analysis of the GFP fold reductions from the circuit show between an 8- and 12-fold decrease in GFP output in response to one or two input RNAs (Fig. 5i).

**Three- and four-input repressor-based logic circuitry.** NAND gate RNAs based on 3WJ repressors were extended to three- and four-input operation by adding additional repressor modules upstream of the output gene. A three-input gate RNA for NOT (A AND B AND C) computations was constructed using three orthogonal 3WJ repressor hairpins separated by 11-nt single-stranded spacer domains (Fig. 6a). This device showed high GFP expression for all logical TRUE conditions lacking at least one of the input RNAs (Fig. 6b), while providing low expression for the logical FALSE condition was at least a factor of 33 higher than any of the changes in expression observed for single-input cases.

The GFP fold changes for this circuit showed a sixfold reduction in GFP expression in the sole logical FALSE state and provided at least a 3.7-fold reduction in GFP compared to all logical TRUE states (Fig. 6f). We also tested multiple additional NAND gates with two to four inputs using combinations of six orthogonal 3WJ repressor
Overall, we found that 23 out of 25 gates operated successfully in *E. coli* (Supplementary Figs. 18 and 19). Further studies also indicated that the performance of the NAND gates can be affected by the copy number of the plasmids used for the input RNAs, with a higher-copy plasmid in one case causing lower expression when targeting a 3WJ repressor downstream of another translation start site (see Supplementary Fig. 20 and Methods for details on additional NAND gates).

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**Fig. 5 | Two-input logic operations using repressor-based devices.**

**a.** Design of a toehold repressor NAND gate where the input RNAs hybridize to form a complete trigger for repression. **b, c.** GFP fluorescence (b) and fold reduction (c) for the toehold repressor NAND device. **d.** Design of a 3WJ repressor NAND gate. In the gate RNA, two switch modules are inserted in-frame and upstream of the reporter gene and both input RNAs must bind to the gate to prevent gene expression. **e, f.** GFP fluorescence (e) and fold reduction (f) for the 3WJ repressor NAND device. **g.** Design of a 3WJ repressor NOR gate. The gate RNA contains a repressor module to regulate the output gene and two trigger modules sequestered within the loops of strong hairpin secondary structures that sense the input RNAs. Binding of either input RNA causes the corresponding hairpin to unwind, which releases the trigger to bind to and inhibit the 3WJ repressor module. **h, i.** GFP fluorescence (h) and fold reduction (i) for the 3WJ repressor NOR device. Devices were measured 6 h after induction. GFP fluorescence values and their error bars are the arithmetic mean and s.d., respectively, of *n* = 3 biologically independent samples. Fold reductions for each device were calculated by dividing the GFP fluorescence value from the gate RNA obtained for the null input case by the GFP fluorescence value for each input combination. Relative errors for the fold reductions were obtained by adding the relative fluorescence errors in quadrature. Individual points show the fluorescence measured from *n* = 3 biologically independent samples (b, e, h) or the fold reduction (c, f, i) from *n* = 3 pairs of biologically independent samples.
An additional four-input logic system was implemented using the toehold repressors. Starting from the A1–A2 NAND gate in Fig. 5a–c, we designed a second pair of bridge domains v and v* and shifted the trigger splitting point 4 nt to generate new NAND inputs B1 and B2 (Fig. 6g and Supplementary Fig. 14b). The resulting ribocomputing device performed the computation NOT ((A1 AND A2) OR (B1 AND B2)) (Fig. 6h–i). As expected, we observed substantial reductions in GFP expression only when A1 and A2 or B1 and B2 were expressed simultaneously. Furthermore, only weak crosstalk was observed when the non-interacting A triggers and B triggers were tested in pairs. The crosstalk observed for the trigger A2 and B1 combination was at least fivefold less than the cognate pair of triggers.

**Discussion**

We have developed two types of high-performance translation-repressing riboregulator using de novo RNA sequence design. Toehold repressors exploit strong RNA secondary structures to provide a very wide dynamic range of gene expression and are best suited for applications requiring tight translational control.
3WJ repressors exhibit very low device crosstalk while using weaker RNA secondary structures and are optimal for multiplexed sensing and multi-input logic. While this work was being performed, an independent study uncovered similar designs for RNA-based transcriptional repressors. These designs exhibited weaker repression efficiency than the systems presented here, but they enabled targeting of endogenous mRNA transcripts.

The synthetic RNA-based repressors provide a wide dynamic range comparable to protein-based transcriptional repressors. A previous 20-component protein repressor library yielded an average 51.3-fold reduction in reporter expression. In comparison, the top 20 second-generation toehold and 3WJ repressors provided average GFP reductions of 122- and 43-fold, respectively. The 3WJ repressors also exhibited good orthogonality, with 15 devices providing 17-fold dynamic range, while the toehold repressors had eight devices with sevenfold GFP reduction.

The toehold and 3WJ repressors were also incorporated into genetically compact ribocomputing devices that effectively computed NOT-related logic expressions with up to four different input RNAs. 3WJ repressor designs, in particular, were amenable to integration into long NAND gate RNAs to simultaneously detect multiple sequence-independent trigger RNAs and displayed excellent modularity, with 92% of the 25 devices tested operating correctly. NOR gates based on 3WJ repressors that exploited intramolecular RNA interactions enabled two-input regulation without requiring translation through downstream hairpins, in contrast to previously reported OR gate systems.

We also successfully applied in-cell SHAPE-Seq to simultaneously characterize RNA structure and function for the 3WJ repressors. Analysis of 3WJ repressors yielded the first direct structural evidence to support the mechanistic model of a de novo-designed riboregulator and also revealed potential pitfalls in our design strategies. These results highlight how SHAPE-Seq can be used to confirm design principles and understand potential failure modes of synthetic riboregulators, which can be used to guide future design improvements.

Overall, the toehold and 3WJ repressors represent versatile new components for the rapidly expanding RNA synthetic biology toolkit. The development of these NOT, NAND, and NOR logic devices coupled with advances in RNA-guided CRISPR/Cas systems and RNA-based transcriptional regulators and systems that merge these capabilities into increasingly sophisticated forms of RNA-enabled genetic circuits that exploit regulation at the transcriptional, translational, and post-transcriptional levels to achieve more dynamic and programmable cellular functions.

Online content
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Methods

Strains and growth conditions. The following E. coli strains were used in this study: BL21 Star DE3 (F⁻ ompT hsdS (rK, mK) gal dcm rm3131 (DE3); Invitrogen), BL21 DE3 (F⁻ ompT hsdS (rK, mK) gal dcm (DE3; Invitrogen), E. coli MG1655/ Marionette-Wild²¹ and MG1655/ Marianette-Wild²¹ (endA1 recA1 proAB (1 glnV44 relA1 hisdR17 (rK, mK))). All strains were grown in LB medium at 37 °C with appropriate antibiotics: ampicillin (50 μg/ml⁻¹), spectinomycin (25 μg/ml⁻¹) and kanamycin (30 μg/ml⁻¹). E. coli MG1655/Marionette-Wild²¹ (sAIM.1506) was a gift from C. Voigt (Addgene bacterial strain no. 108254).

Synthetic repressor computational design. Toehold repressors were designed to provide a 15 nt toehold region for trigger binding and refolding into a repressing hairpin structure identical to that used in toehold switches (Supplementary Fig. 1a). The repressed hairpin had a 12-nt loop and the top 3 bp of the stem was specified (Supplementary Table 1). An additional 12-bp stem domain c* was used to contain only A–U base pairing, which was previously associated with high-performance toehold switches²¹. An additional 12-bp stem domain c* was used to ensure that the repressing hairpin structure would only form upon binding to the trigger RNA. A 4-nt single-stranded region (AAAC) was used upstream of the main RBS sequence (AGAGGAGA) to allow efficient translation of the output gene in the absence of the trigger. These design considerations resulted in a 30-nt-long hairpin stem region for the switch RNA in its active translation state. Three bulges were included in this hairpin structure at 8-nt increments to discourage transcriptional termination through the strong secondary structure. As part of the RNA design, the switch RNA sequence was considered up to the 30 nt following the repressed hairpin structure, which included a 21-nt linker previously used for toehold switch libraries (9 nt of GFP ubiquitous complex design package³⁴, which enabled the trigger, switch and trigger–switch transcription–translation reactions (PURExpress, NEB) at different concentrations. Alternative 3WJ repressors were implemented using the 3OC6-HSL-inducible promoter PluxB to regulate trigger RNA transcription and the anhydrotetracycline (aTc)-inducible promoter PmB for the switch RNA³⁵. Trigger and switch RNAs were expressed using separate plasmids. The trigger was on a high-copy plasmid with a ColE1 origin andkanamyacin resistance. The switch was expressed using the 5′-insulating ribosome binding site (Supplementary Table 1). Trigger RNAs for the toehold repressors were designed with a 5′ hairpin region to maximize RNA stability followed by the c, b and a domains responsible for binding to the switch RNA. The trigger RNA was also designed with the 47-nt T7 terminator sequence (Supplementary Table 1). Three-nucleotide spacers were added between the interaction domains and the outer hairpins as part of the trigger design.

The 3WJ RNAs were designed using the core sequence of first-generation toehold switch number 1¹. This core region is indicated by the grey and black bases within the hairpin structure shown in Supplementary Fig. 1b and has the sequence 5′-UUUGUAAUUGUUGUAAAGAGGAGAGACCAUAGACAAAC-3′ where the RBS and start codon are shown in bold. This hairpin sequence provided very high translational output despite its secondary structure in previous studies¹. The core translational element was integrated into the 3WJ repressor by appending binding domains a* and b* with lengths of 15 nt and 12 nt, respectively, and the last 2 nt of the b domain. The remaining 10 nt of the b domain were via flow cytometry (Supplementary Fig. 8).

Evaluation of repressors with inducible promoters. Inducible expression was implemented using the 3OC6-HSL-inducible promoter PmB to regulate trigger RNA transcription and the anhydrotetracycline (aTc)-inducible promoter PmB for the switch RNA³⁵. Trigger and switch RNAs were expressed using separate plasmids. The trigger was on a high-copy plasmid with a ColE1 origin and kanamycin resistance. The switch was expressed using the 5′-insulating ribosome binding site (Supplementary Table 1). Trigger RNAs for the toehold repressors were designed with a 5′ hairpin region to maximize RNA stability followed by the c, b and a domains responsible for binding to the switch RNA. The trigger RNA was also designed with the 47-nt T7 terminator sequence (Supplementary Table 1). Three-nucleotide spacers were added between the interaction domains and the outer hairpins as part of the trigger design.

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Toehold and 3WJ repressor library construction. Plasmids were constructed using PCR and Gibson assembly. DNA templates for repressor switch and trigger RNA sequence were assembled from single-stranded DNAs purchased from Integrated DNA Technologies. The synthetic DNA strands were amplified via PCR and then inserted into plasmid backbones using 30-nt homology domains via Gibson assembly⁶. All plasmids were cloned in the E. coli DH5α strain and validated through DNA sequencing. Backbones for the plasmids were taken from the commercial vectors pET15b (ampicillin resistance, ColE1 origin), pCOLADuet (kanamycin resistance, ColA origin) and pCDFDuet (spectinomycin resistance, CDC origin) from EMD Millipore, and the repressor DNA was inserted upstream of the T7 terminator sequence to replace their respective multiple cloning sites. GFPmut3b-ASV, GFPmut3b with an ASV degradation tag¹¶, was used as the reporter for the repressor switch plasmids, except for experiments studying GFPPn with different degradation tags. In addition, the kanR mRNA toehold repressor was used and the E. coli BL21 Star DE3 strain without a degradation tag. The 3WJ RNA sequences commonly used in the plasmids are provided in Supplementary Table 1.

Toehold and 3WJ repressor expression. Toehold and 3WJ repressor switch and trigger RNAs were expressed using T7 RNA polymerase in BL21 Star DE3, an RNase-deficient strain, with the T7 RNA polymerase induced with the addition of IPTG. Selected sets of toehold and 3WJ repressor switch and trigger RNAs were also tested in BL21 DE3 strain with the T7 RNA polymerase induced with the addition of IPTG. For both strains, plasmids were grown overnight in LB medium with antibiotics with shaking at 900 r.p.m. and 37 °C. Overnight cultures were then diluted by 100-fold into fresh LB medium with antibiotics and returned to the shaker (900 r.p.m., 37 °C). After 80 min, both strains were induced with 0.1 mM IPTG and cells were returned to the shaker (900 r.p.m., 37 °C) until the flow cytometry measurements at specified times post-induction.

Flow cytometry measurements and analysis. Flow cytometry measurements of toehold repressor libraries and their ribocomputing devices were performed using a BD LSRII Fortessa cell sorter with a high-throughput sampler. Before loading to the flow cytometer, cells were diluted by a factor of ~65 into phosphate-buffered saline and approximately 50,000 cells were recorded for each measurement. Flow cytometry measurements of 3WJ repressor libraries and their ribocomputing devices were performed using a Stratagene S1300EXI cell sorter equipped with an A600 high-throughput autosampler. Cells with the 3WJ repressor systems were diluted by a factor of ~17 into phosphate-buffered saline and detected as described above, with 40,000 cells recorded for each measurement. Cell populations were gated according to their FSC and side scatter (SSC) distributions as described previously⁶ (Supplementary Fig. 21), and the GFP or mCherry fluorescence levels of these gated cells were used to measure circuit output via the geometric mean from at least three biological replicates. Fold reductions of GFP or mCherry fluorescence levels were then calculated by taking the geometric mean fluorescence output of the toehold or 3WJ repressor switch with a non-cognate trigger and dividing it by the fluorescent output with a cognate trigger. Cellular autofluorescence was not subtracted before determining the fold reduction.

Evaluation of repressors in cell-free systems. Trigger and switch RNAs for toehold and 3WJ repressors were separately transcribed, quantitated and added to transcription–translation reactions (PURExpress, NEB) at different concentrations. Time-course measurements were then conducted on the reactions using a plate reader (Biotek H11MF) to determine GFP expression (Supplementary Fig. 6).

3WJ repressors with stem sequence variations. Alternative 3WJ repressors with the same secondary structure but different RNA sequences were studied to determine the effect of sequence changes on 3WJ repressor performance. The variants all used the trigger RNA of 3WJ repressor index 20. The ‘NN’ variants were generated by allowing any base (that is, N bases) to be present within the stem region of the switch RNA (white bases in Supplementary Fig. 7a). To generate weaker stems, ‘SW’ variants were designed with a combination of Strong (G–C) base pairs and Weak (A–U) base pairs that matched those of the original hairpin sequence. Both types of design were screened to ensure that they did not have any in-frame stop codons following the start codon in the stem.

Two versions of the NN design and three versions of the SW design were tested in E. coli BL21 Star DE3 (see Supplementary Table 5 and Supplementary Fig. 7b for sequence information). Both NN designs were unable to modulate GFP expression and provided near-background GFP levels even in the absence of the trigger (Supplementary Fig. 7c). Two of the three SW designs (design A and C) provided substantial GFP output without the trigger, while expression from the third (device A) was nearly undetectable (Supplementary Fig. 7c,d).

Repressors using GFP with different degradation tags. To examine the effect of the degradation tag on riboregulator performance, 3WJ repressor systems were tested in E. coli BL21 Star DE3, using three different types of output GFPmut3b protein: ASV-tagged (~110 min half-life)⁶, LVA-tagged (~40 min half-life)⁶ and untagged. The resulting repressor systems were tested using the same conditions and plasmid combinations as the repressor libraries and characterized using flow cytometry (Supplementary Fig. 8).
In-cell SHAPE-Seq measurements and analysis. In-cell SHAPE-Seq measurements were carried out as described previously19. Briefly, 3WJ repressor variants were transferred into BL21 Star DE3 as in the functional characterization experiments. Overnight cultures were diluted 100-fold into 1.2 ml of fresh LB medium with antibiotics. Following IPTG induction and 5 h additional subculture, 100 µl of culture was removed and diluted by ~100-fold for functional characterization using a BD Accuri cell analyser with a high-throughput sampler. A 500 µl volume of the remaining culture was then added to 13.3 µl of 250 mM IM7 or 13.3 µl of DMSO (control solvent). Cells were returned to shaking for 3 min to allow IM7 to react, then cellular RNAs were Trizol extracted and reverse transcribed using a custom reverse transcription primer specific for GFPmut3b (5′-CAACAAGATCTGCAACACTCCAGTG-3′). Additional 5′ and 3′ sequencing adapters were then added. Following 2× 35 pM-paired-end Illumina sequencing, β reactivities were calculated as described in ref. 19. Error bars represent the standard deviation of three samples, each probed from a separate transformation on a separate day. Duplicate samples were only processed in parallel during final sequencing.

**Experimental testing of ribocomputing devices.** For device testing, the RNA inputs and the gate RNA were separated from separate plasmids through the T7 promoter in BL21 Star DE3 cells. In cases where an input RNA was not present, a non-cognate input RNA was expressed in its place. Culturing and induction of the cells were performed in the same way as the repressor libraries.

**Study of 3WJ repressor NAND gate modularity.** Multiple versions of the two-, three- and four-input 3WJ NAND gates were tested to assess their modularity (see Supplementary Table 10 for sequence information). NAND gates were designed from a parent library of six 3WJ repressor modules selected based on three low reduction levels of 3WJ repressor hairpins. The NAND gates were selected for testing based on their relatively low expected ensemble defect in NUPACK. GFP fluorescence and GFP fold reduction were obtained in E. coli BL21 Star DE3 for 16 different two-input NAND systems: eight with 11-nt spacers (Supplementary Fig. 18a) and eight with 17-nt spacers (Supplementary Fig. 18b). Six different three-input NAND gates and three different four-input NAND gates were generated from a smaller subset of four 3WJ repressor modules after elimination of index 10 and index 21 hairpins based on their lower performance in two-input NAND computations (Supplementary Fig. 19).

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability.** The datasets generated during and/or analysed during the current study are available from the corresponding authors on reasonable request. Supplementary Tables are available from A.A.G. in the spreadsheet format upon request. The following plasmids from the study are available from Addgene: pYZ_3Wrep_N10_switch 132722; pYZ_3Wrep_N19_switch 132723; pYZ_3Wrep_N10_switch 132724; pYZ_3Wrep_N14_switch 132725; pYZ_3Wrep_N22_switch 132726; pYZ_3Wrep_N20_trigger 132727; pYZ_3Wrep_N19_trigger 132727; pYZ_3Wrep_N10_trigger 132728; pYZ_3Wrep_N24_trigger 132729; pYZ_NAND2_L17_S19_S11_32730; pYZ_NAND3_L11_S11_S13_32731; pYZ_NAND4_L17_S14_S11_S13_32732; pYZ_3Wrep_N11_trigger 132727; pYZ_3Wrep_N12_trigger 132734; pYZ_GateB_3W_N13_trigger 132735; pAG_PluxB_ToRep_N01_trigger 132736; pAG_Pet_“ToRep_N01_switch 132737; pYZ_PluxB_3W_N19_N10_trigger 132738; pYZ_Pet_“ToRep_N19_switch 132739; pAG_ToRep_N09_trigger 132740; pAG_ToRep_N09_switch 132741; pIK_ToRepG2_N02_switch 132742; pIK_ToRepG2_N64_switch 132743; pIK_ToRepG2_N19_switch 132744; pIK_ToRepG2_N02_trigger 132745; pIK_ToRepG2_N64_trigger 132746; pIK_ToRepG2_N19_trigger 132747.

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Author contributions

J.K. and A.A.G. designed the toehold repressors and ribocomputing devices. J.K., M.T. and A.A.G. performed experiments for the toehold repressors and their ribocomputing devices. Y.Z., S.C. and A.A.G. designed the 3WJ repressors and ribocomputing devices. Y.Z. and S.C. performed experiments for the 3WJ repressors and their ribocomputing devices. P.D.C. performed SHAPE-Seq measurements. J.K., Y.Z., A.A.G. and P.D.C. analysed the data. J.K., Y.Z., P.D.C. and A.A.G. wrote the manuscript. J.K., A.A.G., P.D.C., J.B.L. and P.Y. edited the manuscript. A.A.G., P.Y., J.B.L., J.J.C., P.A.S. and F.C.S. supervised the research.

Competing interests

US provisional patents have been filed by J.K., A.A.G., J.J.C. and P.Y. and by Y.Z. and A.A.G., based on this work. P.Y. is the co-founder of Ultivue Inc. and NuProbe Global.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41589-019-0388-1.

Correspondence and requests for materials should be addressed to P.Y. or A.A.G.

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Software and code

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Data collection
BD FACSDiva (version 8) and CellCapTure (version 4) were used for collecting flow cytometry data.

Data analysis
Matlab R2017b was used for data analysis.

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The datasets generated during and/or analysed during the current study are available from the corresponding authors on reasonable request. Supplementary Tables are available from A.A.G. in spreadsheet format upon request. The following plasmids from the study are available from Addgene: pAG_ToeRep_N09_trigger, pAG_ToeRep_N09_switch, pYZ_3WJrep_N10_trigger, pYZ_3WJrep_N10_switch, pYZ_3Wirep_N19_trigger, pYZ_3Wirep_N19_switch, pYZ_3Wirep_N20_trigger, pYZ_3Wirep_N20_switch, pYZ_3Wirep_N24_trigger, pYZ_3Wirep_N24_switch, pAG_PluxB_ToeRep_N01_trigger, pAG_PluxB_ToeRep_N01_switch, pYZ_PluxB_3WJrep_N19_trigger, pYZ_PluxB_3WJrep_N19_switch, pAG_Ptet*_ToeRep_N01_trigger, pAG_Ptet*_ToeRep_N01_switch, pYZ_Ptet*_3WJrep_N19_trigger, pYZ_Ptet*_3WJrep_N19_switch, pJK_ToeRepG2_N02_trigger, pJK_ToeRepG2_N02_switch, pJK_ToeRepG2_N19_trigger, pJK_ToeRepG2_N19_switch, pJK_ToeRepG2_N64_trigger, pJK_ToeRepG2_N64_switch, pYZ_NAND2_L17_S19_S11, pYZ_NAND3_L11_S11_S13_S19, pYZ_NAND4_L17_S24_S11_S19_S13, pYZ_3Wirep_N11_trigger, pYZ_3Wirep_N12_trigger, and pYZ_3Wirep_N13_trigger.
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**Life sciences study design**

All studies must disclose on these points even when the disclosure is negative.

**Sample size**  
Flow cytometry was performed on populations of at least 30,000 cells. Three biological replicates were analyzed for each condition using flow cytometry. Sample replicate size was chosen based on the limited degree of variability in reporter protein expression observed in earlier experiments.

**Data exclusions**  
Data were not excluded from the analyses.

**Replication**  
At least three biological replicates were measured for each experiment. All attempts at replication have been successful and confirm device function.

**Randomization**  
Organisms were transformed with different DNA and subject to different inducers as necessary as described in the paper. Any covariates were controlled by processing experimental groups to be compared in identical conditions.

**Blinding**  
Investigators were not blinded to group allocation. Blinding was not possible since the investigators generated the different experimental groups studied.

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**Methodology**

**Sample preparation**

Prior to loading to flow cytometer, cells were diluted by a factor of ~65 into phosphate-buffered saline. Cells were detected using a forward scatter (FSC) trigger and at least 30,000 cells were recorded for each measurement. Flow cytometry measurements of 3WJ repressor libraries and their ribocomputing devices were performed using a Stratedigm S1300EXi cell analyzer equipped with a A600 high-throughput autosampler. Cells with the 3WJ repressor systems were diluted by a factor of ~17 into phosphate-buffered saline and detected as described above with 40,000 cells recorded for each measurement.

**Instrument**

BD LSRFortessa, Stratedigm S1300EXi, BD Accuri
| **Software** | Flow cytometry data was collected using proprietary software packages from BD (BD FACSDiva) and Stratedigm (CellCapTure). The flow cytometry data was processed using Matlab. |
|-------------|---------------------------------------------------------------------------------------------------------|
| **Cell population abundance** | The cells were not sorted. The cells presented a single population based on FSC and SSC measurements. |
| **Gating strategy** | A two-dimensional histogram of cell counts versus FSC and SSC values was generated. The gate was defined to include all cells having FSC and SSC values that provided at least 10% of the cell count obtained from the peak of the two-dimensional histogram. |

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