dCas9 regulator to neutralize competition in CRISPRi circuits

Hsin-Ho Huang1,5, Massimo Bellato2,5, Yili Qian1, Pablo Cárdenas3, Lorenzo Pasotti2, Paolo Magni2 & Domitilla Del Vecchio1,4

CRISPRi-mediated gene regulation allows simultaneous control of many genes. However, highly specific sgRNA-promoter binding is, alone, insufficient to achieve independent transcriptional regulation of multiple targets. Indeed, due to competition for dCas9, the repression ability of one sgRNA changes significantly when another sgRNA becomes expressed. To solve this problem and decouple sgRNA-mediated regulatory paths, we create a dCas9 concentration regulator that implements negative feedback on dCas9 level. This allows any sgRNA to maintain an approximately constant dose-response curve, independent of other sgRNAs. We demonstrate the regulator performance on both single-stage and layered CRISPRi-based genetic circuits, zeroing competition effects of up to 15-fold changes in circuit I/O response encountered without the dCas9 regulator. The dCas9 regulator decouples sgRNA-mediated regulatory paths, enabling concurrent and independent regulation of multiple genes. This allows predictable composition of CRISPRi-based genetic modules, which is essential in the design of larger scale synthetic genetic circuits.
The clustered regularly interspaced short palindromic repeats interference (CRISPRi) allows to create many orthogonal transcriptional repressors, each constituted of catalytically inactive Cas9 (dCas9) bound to a single guide RNA (sgRNA) \( c_i \) in Fig. 1a. This is instrumental to modulate complex transcriptional programs \(^\text{1,2}\) and to build larger scale synthetic genetic circuits \(^\text{3,4}\). Multiplexing is possible because many sgRNAs can be concurrently expressed to recruit dCas9 each to a different promoter with high specificity to regulate transcription \(^\text{5}\). High specificity of sgRNA-promoter binding allows the creation of a large library of transcriptional repressors that, in principle, do not interfere with one another \(^\text{6}\). High specificity combined with lower loading to gene expression resources than with protein-based transcription factors makes the CRISPRi-dCas9 system an optimal solution to create increasingly large and sophisticated transcriptional programs \(^\text{5}\).

Despite the high specificity of sgRNA-promoter binding, multiple sgRNAs can still interfere with one another by competing for binding to dCas9 \(^\text{9,10}\). This binding imparts a load to the dCas9, which affects the level of free dCas9 (apo-dCas9) (Fig. 1a). Specifically, Zhang et al. showed that the fold-repression exerted by any one sgRNA through CRISPRi can decrease by up to five times when additional sgRNAs are expressed \(^\text{9}\). In ref. \(^\text{10}\), the authors observed that the ability of one sgRNA to repress its target was hampered when a second sgRNA without a target was constitutively expressed, consistent with predictions from mathematical models \(^\text{11,12}\). In these works, and more generally in all CRISPRi-dCas9 transcriptional programs to date, dCas9 protein is generated at a constant pre-fixed rate. As a consequence, the concentration of dCas9 bound to any sgRNA species decreases as additional sgRNAs that bind to dCas9 become expressed. This leads to undesirable coupling among theoretically orthogonal sgRNA-mediated regulatory paths, wherein each sgRNA modulates directly the transcription of its targets but also indirectly affects transcription of non-target genes \(^\text{12}\). This interference confounds design since the combined input/output (I/O) response of multiple CRISPRi modules (CM) operating concurrently in the cell is different from that predicted using the I/O responses of each CM characterized in isolation. Although the effects of dCas9 loads may, in principle, be mitigated by increased levels of dCas9 (see simulations in Supplementary Fig. 1), it is difficult to increase dCas9 level in practice because this causes severe growth defects (see refs. \(^\text{6,13,14}\) and Supplementary Note 2).

To address this problem, less toxic mutations of dCas9 protein have appeared. Yet, even at the allowed higher dCas9 levels, the effects of dCas9 loading remain prominent \(^\text{9}\).

In this work, we present a regulated dCas9 generator that adjusts the production rate of dCas9 such that the concentration...
of apo-dCas9, and thus that of dCas9 bound to any sgRNA species, becomes practically independent of the load (Fig. 1b). This, in turn, ensures an approximately constant repression strength of any sgRNA when additional sgRNAs become expressed, thus decoupling regulatory paths of different sgRNAs. We create an experimental model system that recapitulates the dCas9 loading problem, in which dCas9 is produced by either an unregulated (UR) (Fig. 1a) or a regulated (R) (Fig. 1b) dCas9 generator. In the unregulated dCas9 generator, the production rate of dCas9 is constant (Fig. 1a). By contrast, the regulated dCas9 generator produces dCas9 at a rate that is negatively regulated by the level of apo-dCas9 itself through CRISPRi by means of sgRNA g0 (Fig. 1b). The regulator’s performance is evaluated on genetic circuits that show severe alterations of their I/O responses when a competitor sgRNA is expressed.

Results
Decoupling NOT gates’ I/O responses from competitor sgRNA expression. We first evaluated the performance of the regulated dCas9 generator by comparing the extent by which the I/O response of a CM-based logic NOT gate (Fig. 1c) is affected by the expression of a competitor sgRNA. The CM-based NOT gate is constituted of a CM with sgRNA g1, expressed by an inducible promoter and a genetic module expressing red fluorescent protein (RFP) as an output. The sgRNA g1 represses RFP’s transcription through CRISPRi. A second CM (competitor CM) contains sgRNA g2 expressed by a constitutive promoter. This competitor CM plays the role of any additional CM that may become activated in a system, such as from other logic gates15. When competitor sgRNA g2 is present and binds to dCas9, the level of apo-dCas9 decreases. In the regulated dCas9 generator, this drop reduces the concentration of the dCas9-g2 complex c0, thereby de-repressing dCas9 transcription. As a result, the level of apo-dCas9 increases, thus balancing the initial drop due to the load exerted by competitor sgRNA g2. This ultimately allows the concentration of dCas9 bound to sgRNA g1 in the NOT gate to remain approximately constant when competitor g2 becomes expressed.

We created an ordinary differential equation (ODE) model of the regulated dCas9 generator, the CM-based logic NOT gate, and the competitor CM (Supplementary Eqs. (33–35)). Simulations of this model show that the unpressed production rate of dCas9 and the production rate of sgRNA g0 are the two key design parameters. When these production rates are both sufficiently high, expression of competitor sgRNA g2 no longer affects the NOT gate I/O response (simulations in Fig. 1d). In-depth mathematical analysis of the regulated dCas9 generator ODE model further shows that the sensitivity of apo-dCas9 level to the expression rate of the competitor sgRNA g2 can be arbitrarily diminished by picking a sufficiently large expression rate for sgRNA g0 (Supplementary Note 1.3). This sensitivity reduction, in turn, mitigates the effects of competitor sgRNA g2’s expression on the NOT gate I/O response, although potentially affecting the ability of sgRNA g1 to fully repress its target (simulations in Supplementary Fig. 1 and experimental data in Supplementary Fig. 5d, e). By also increasing dCas9 protein production rate, sgRNA g1 can fully repress its target without hampering robustness of the NOT gate I/O response to competitor sgRNA g2’s expression (simulations in Supplementary Fig. 1 and experimental data in Supplementary Fig. 5e, f)). Therefore, our regulated dCas9 generator employs strong promoter for sgRNA g0 and also higher dCas9’s promoter and RBS strengths compared to those of the unregulated generator (compare Fig. 1a (UR) with Fig. 1b (R)).

We implemented the CM-based NOT gate by using an HSL-inducible promoter controlling the expression of g1. The competitor CM was created by expressing a competitor sgRNA g2 through a library of constitutive promoters (Fig. 2a and Supplementary Note 6). The competitor sgRNA g2 was designed to target a DNA sequence not present in the circuit nor in the bacterial genome (as predicted by Benchling’s sgRNA design tool, see Supplementary Note 5) to avoid additional interactions that could confound analysis16. We experimentally evaluated the performance of the regulated dCas9 generator by first building CMs in high copy number plasmids (~84 copies, Supplementary Note 6) in order to create large dCas9 loads. In the unregulated generator configuration, the choice of dCas9’s promoter and RBS ensures production of dCas9 at a level sufficient to enable complete repression of target promoters by sgRNAs without substantially affecting growth rate (see ref. 17 and Supplementary Note 6, Supplementary Fig. 9, Supplementary Note 7). With the unregulated dCas9 generator, expression of the competitor sgRNA causes changes of 16, 17, or 42 fold in the I/O response of the CM-based NOT gate depending on the input level (Fig. 2b). By contrast, with the regulated dCas9 generator no appreciable change is observed (Fig. 2c), indicating that the regulated dCas9 generator can attenuate large dCas9 loads.

We then investigated to what extent dCas9 competition still affects the function of logic gates when the CMs are built on low copy number plasmids. Thus, the NOT gate and competitor CMs were both assembled on pSB4C5 plasmid (giving ~5 copies, Supplementary Note 6). Interestingly, even when the CMs are on a low copy number plasmid, expressing competitor sgRNA g2 still leads to appreciable changes in the I/O response of the NOT gate, leading to a 2-fold change in the NOT gate’s output when the unregulated generator is used (Fig. 2d). These changes disappear when the regulated dCas9 generator is employed (Fig. 2e).

We also tested the load mitigation property of the regulated dCas9 generator in an alternative bacterial strain (Supplementary Note 4) and adopting a different design of the NOT gate, which uses an aTc-inducible promoter to control the expression of sgRNA g1. In this case and with CMs in high copy number plasmids, expressing a competitor sgRNA results in up to 13-fold change in the NOT gate’s I/O response with unregulated dCas9 generator, while it still gives inappreciable change when the regulated generator is used (Supplementary Fig. 8).

Decoupling layered circuits’ I/O responses from competitor sgRNA expression. To demonstrate that the ability of the regulated dCas9 generator to mitigate the effects of dCas9 loading is not specific to a single CM-based NOT gate, we built a layered logic circuit constituted of two NOT gates arranged in a cascade (Fig. 3a, b). NOT gate cascades are a prototypical example of layered logic gates and are ubiquitous in circuits computing sophisticated logic functions13,15,18,19. In the cascade design, the LuxR/HSL input activates expression of sgRNA g1 in CM1, which, in turn, represses the expression of sgRNA g3 in CM4 through CRISPRi. The output of CM1 then represses a promoter expressing the RFP output protein (Fig. 3b). As before, a competitor sgRNA g2 was included or omitted from the system (CM4). The two CMs constituting the cascade and the competitor CM are all on a high copy number plasmid (~84 copies). The I/O response of the cascade was measured with or without the competitor sgRNA g2 and with either the unregulated or regulated dCas9 generators. The I/O response of the cascade shows approximately a 4-fold change at low induction levels when the competitor sgRNA is included and the unregulated dCas9 generator is used (Fig. 3c). By contrast, the cascade’s I/O response shows no appreciable changes when the regulated dCas9 generator is used.
The dCas9 regulator’s ability to decouple the function of a circuit from expression of additional sgRNAs is independent of the specific genetic components of the circuit (i.e., promoters and inducers) and of whether the circuit contains one or multiple connected CMs, such as in layered systems.

Discussion

In summary, our regulated dCas9 generator effectively decouples the regulatory functions of different sgRNAs from one another, despite that these sgRNAs compete for dCas9. This enables true independent modulation of multiple genes simultaneously, which is not possible with commonly used unregulated dCas9 generators. Independent modulation of multiple genes by different sgRNAs is critical for scalability of CRISPRi-based genetic circuits. Indeed, scalability requires that the characteristics of any circuit stay unchanged when other circuits are added to the cell. With an unregulated dCas9 generator the I/O characteristics of various logic gates are severely affected when adding just one more CM to the cell (Figs. 2b, d and 3c). By contrast, with the regulated dCas9 generator, the gates’ I/O characteristics are unperturbed by addition of the same CM (Figs. 2c, e and 3d). The regulated dCas9 generator is implemented in a dedicated plasmid (Supplementary Note 6 and Supplementary Fig. 9) and can

**Fig. 2 Effect of competitor CM on CM-based NOT gate with UR and R dCas9 generators.** a Genetic diagram of a CM-based NOT gate and competitor CM. The NOT gate is composed of a CM comprising sgRNA g1, giving complex c1 as an output, and of a genetic module that takes c1 as input and gives RFP as an output. Here, g1’s expression is regulated by HSL/LuxR (IN). A competitor CM that expresses competitor sgRNA g2 with variable promoter Pc is either present or absent. The CMs were placed either on a low copy (LC) number plasmid (~5 copies) or on a high copy (HC) number plasmid (~84). The dCas9 generator was placed on a medium (~20) copy number plasmid (Supplementary Notes 6). b CM-based NOT gate I/O response with UR dCas9 generator and CMs on HC plasmid. Turquoise line represents I/O responses in the absence of competitor, while yellow (medium competitor) and orange (higher competitor) lines represent system I/O responses in which the expression of g2 is driven by a weaker (BBa_J23116) or stronger (pTrc) promoter, respectively (Supplementary Note 6 and Supplementary Table 13). Inset shows steady state I/O responses obtained by simulating the ODE model of the unregulated system described in Supplementary Eqs. (30–32) with parameters in Supplementary Table 3. (e) CM-based NOT gate I/O response with R dCas9 generator and CMs on HC plasmid. Inset shows steady state I/O responses obtained by simulating the ODE model of the regulated system described in Supplementary Eqs. (33–35) with parameters in Supplementary Table 3. RFP distributions obtained by flow cytometry are reported in Supplementary Fig. 12. d CM-based NOT gate I/O response with UR dCas9 generator and CMs on LC plasmid. Yellow (medium competitor) and orange (higher competitor) lines represent system I/O responses in which the expression of g2 is driven by a weaker (BBa_J233100) or stronger (pTrc) promoter, respectively (See Supplementary Note 6 and Supplementary Table 13). Inset shows steady state I/O responses obtained by simulating the ODE model of the unregulated system described by Supplementary Eqs. (30–32) with parameters in Supplementary Table 3. e CM-based NOT gate I/O response with R dCas9 generator and CMs on LC plasmid. Inset shows steady state I/O responses obtained by simulating the ODE model of the regulated system described in Supplementary Eqs. (33–35) with parameters in Supplementary Table 3. Data in the line plots represent mean values ± SD of n = 3 or 4 biologically independent experiments. Fold changes are normalized to the no competitor data. Bar graphs with error bars and overlaid dot plots are computed according to Eqs. (1-3) in “Methods” section. Insets’ IN and OUT have the same units as in the data plots. Source data are provided with this paper.
be easily transported across compatible bacterial strains and applications of CRISPRi-dCas9 systems.

The general problem of resource competition in genetic circuits has been widely studied in the context of competition for transcriptional resources in mammalian cells\textsuperscript{22,23}. In particular, studies in bacteria have shown that the effects of ribosome competition on the I/O response of a genetic circuit can be very subtle, yet dramatic, as the inner circuit’s modules compete with one another for ribosomes\textsuperscript{20}. Similar phenomena are also observed to be required in more sophisticated CRISPRi-based genetic circuits\textsuperscript{12}. Feedback regulation systems have been designed before to enhance robustness of bacterial genetic circuits to loading of gene expression resources, i.e., the ribosome\textsuperscript{24–26}. However, none of these approaches is directly applicable to regulate apo-dCas9 level as they either use ribosome-specific parts\textsuperscript{28}, require the protein to be regulated (dCas9 in our case) to act as a transcriptional activator\textsuperscript{24} or to sequester a transcriptional activator\textsuperscript{25}. The dCas9 regulator is simple, compact, and exploits directly the ability of dCas9 to act as a transcriptional repressor when recruited to a promoter by sgRNAs. Feedback controllers enabled by CRISPRi have also appeared in other applications, where expression of a target gene is downregulated through a CRISPRi-mediated negative feedback to prevent growth rate defects due to over-expression\textsuperscript{27}. When expressing multiple sgRNAs from the chromosome, i.e., in one copy, reduced loading effects are expected\textsuperscript{28} and a regulated dCas9 generator may not be required in such cases. However, we have shown that for CRISPRi circuits constructed on plasmids, loading effects are prominent even at low plasmid copy number (~5 copies, Fig. 2d). In these cases, it is expected that a regulated dCas9 generator will be required in order to ensure that multiple sgRNAs can concurrently and independently control their targets.

By combining high dCas9 expression rate with a strong feedback repression, our regulated dCas9 generator neutralizes any loading effect, while keeping dCas9 at sufficiently low levels to prevent growth defects (Supplementary Note 7). Nevertheless, transient toxicity is observed immediately after transformation of the regulated dCas9 generator plasmid into bacteria. In fact, immediately after this plasmid is transformed into cells, the initial concentrations of sgRNA g0 and dCas9 are both zero, but dCas9 production rate is high and, due to a zero g0 concentration, is initially unregulated. Thus, dCas9 concentration increases rapidly after the plasmid is transformed and before g0 level is sufficiently high to repress dCas9 transcription. This, in turn, may create an overshoot in dCas9 concentration resulting in toxic effects to cells. To decrease the overshoot in dCas9 concentration following transformation of the plasmid, we prepared host cells with a removable plasmid that produces the holocomplex dCas9-g0. This represses dCas9 transcription from the plasmid encoding the regulated dCas9 generator immediately upon its transformation into cells, thus removing transient toxicity (see “Methods” section and Supplementary Note 2).

The adoption of multiple orthogonal dCas variants\textsuperscript{29} could theoretically help mitigate the competition between two or more sgRNAs by distributing the resource demand among different DNA binding proteins. However, in large circuits composed of many CMs, it is still expected that multiple sgRNAs will need to share the same dCas variant. Indeed, since multiple variants are
expressed, it is also expected that the level of each single variant will be more limited, to prevent toxicity, than when using a single variant. Therefore, the effects of dCas competition can, in principle, be even more prominent than those observed here due to lower levels of the shared resource. In these situations, a dCas regulator will likely be required to neutralize competition. In general, a hybrid approach in which the variants shared among multiple sgRNAs include a regulator, and those that are not shared are not regulated could form an optimal solution. Although we did not investigate this aspect in this paper, the impact of resource competition on the dynamic behavior of a genetic circuit can also result in dramatic outcomes\(^\text{30}\). Further investigations are required to determine the benefit of the regulated dCas9 generator when dynamic behavior is of interest\(^\text{13,2}\).

Overall, we expect that adoption of our regulated dCas9 generator in CRISPRi systems will make CRISPRi-based programs more scalable, reliable, and predictable.

**Methods**

**Strain and growth medium.** Bacterial strain *E. coli* NEB10B (NEB, C30190) was used in genetic circuit construction and characterization. The growth medium used in construction was LB broth Lennox. The growth medium used in characterization was M9 medium supplemented with 0.4% glucose, 0.2% casamino acids, and 1 mM thiamine hydrochloride. Appropriate antibiotics were added according to the selection marker of a genetic circuit. Final concentration of ampicillin, kanamycin, and chloramphenicol are 100, 45, and 12.5 μg mL\(^{-1}\), respectively.

**Genetic circuit construction.** The genetic circuit construction was based on Gibson assembly method\(^\text{13}\). DNA fragments to be assembled were amplified by PCR using Phusion High-Fidelity PCR Master Mix with GC Buffer (NEB, M0552S), purified with gel electrophoresis and Zymoclean Gel DNA Recovery Kit (Zymo Research, D4002), quantified with the nanophotometer (Implen, P330), and assembled with Gibson assembly protocol using NEBuilder HiFi DNA Assembly Master Mix (NEB, E2621S). Assembled DNA was transformed into competent cells prepared by the CCMB80 buffer (TekaNova, C3132). Plasmid DNA was prepared by the plasmid miniprep-classic kit (Zymo Research, D4015). DNA sequencing was used Qunatarbio DNA basic sequencing service. The lists of plasmids and primers used for 13 h. Overnight culture was prepared by inoculating a ~80 °C glycerol stock in 800 μL growth medium per well in a 24-well plate (Falcon, 351147) and grew at 30 °C, 220 rpm in a horizontal orbiting shaker for 13 h. Overnight culture was first diluted to an initial optical density at 600 nm (OD\(_{600nm}\)) of 0.001 in 200 μL growth medium per well in a 96-well plate (Falcon, 351172), and grew for 2 h to ensure exponential growth before induction. The 96-well plate was incubated at 30 °C in a Synergy MX (Biotek, Winooski, VT) microplate reader in static condition, and was shaken at a fast speed for 3 s right before OD and fluorescence measurements. Sampling interval was 5 min. Excitation and emission wavelengths to monitor RFP fluorescence were 584 and 619 nm, respectively. To ensure enough time to reach a steady state RFP/OD signal while the cells are in exponential growth, the cell culture was diluted with fresh growth medium to OD\(_{600nm}\) of 0.01 when OD\(_{600nm}\) approached 0.12 at the end of each batch. Multiple batches were conducted for a total experiment time of up to 25 h until RFP/OD reaches steady state (see Supplementary Note 9). The steady state I/O characteristics reported in the main text and the corresponding growth rates in Supplementary Note 7 were both computed from the last batch of each experiment.

**Quantification of competition effects.** To quantify competition effects, fold-change of a system’s output at a given induced condition i was calculated by considering the ratio between the RFP/OD value of the system with competitor sgRNA (g\(_j\)) and that of the corresponding system bearing no competitor sgRNA (e.g., pOP94 and pCL112 for Fig. 2b, c). In particular, the height of the bars represents mean fold-changes computed as follows:

\[
\text{Mean fold -- change (inducer = i)} = \frac{\sum_{j=1}^{N} [\text{RFP}/\text{OD}(\text{with } g_j, \text{ inducer = i})]}{\sum_{j=1}^{N} [\text{RFP}/\text{OD}(\text{without } g_j, \text{ inducer = i})]} / M
\]

(1)

where subscript j represents steady state RFP and OD measurements for the j-th biologically independent replicate, and N and M are the total number of replicates in the experiments with and without competitor g\(_j\), respectively.

The size of the error bars represents one standard deviation (SD) of the respective fold-change. Specifically, for each induction level i, we account for propagation of uncertainty according to

\[
\text{SD of fold -- change} = \frac{\text{mean fold -- change}}{\sqrt{\text{CV}^2(\text{RFP/OD with } g_j) + \text{CV}^2(\text{RFP/OD without } g_j)}},
\]

where \(\text{CV}(\text{RFP/OD with } g_j)\) and \(\text{CV}(\text{RFP/OD without } g_j)\) are the coefficients of variation from biological replicates for steady state RFP/OD with and without competitor g\(_j\), respectively.

The dot plots overlaid with bar graphs represent RFP/OD fold-changes for individual biological replicate with g\(_j\) with respect to the mean of RFP/OD without g\(_j\). Specifically, for the jth replicate at induction level i, we draw a dot according to:

\[
\text{Replicate fold -- change (inducer = i)} = \frac{\sum_{j=1}^{N} [\text{RFP}/\text{OD}(\text{with } g_j, \text{ inducer = i})]}{M}
\]

(2)

Note that by Eq. (2), the size of the error bars does not quantify the uncertainty in the dots defined according to Eq. (3), as the latter does not account for uncertainties in RFP/OD without g\(_j\).

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

**Data availability**

Simulation and fluorescence data generated or analyzed during this study are included in the paper and its Supplementary Information files. A reporting summary for this Article is available as a Supplementary Information file. The source data underlying Figs. 2 and 3 are provided as a Source Data file. Any other relevant data are available from the authors upon reasonable request.

**Code availability**

Custom MATLAB (The MathWorks, Inc., Natick, MA, USA) codes are used to perform numerical simulations. A Supplementary Software file is provided, which includes codes to produce simulation results in Figs. 2 and 3.

Received: 3 November 2020; Accepted: 29 January 2021;
Published online: 16 March 2021

**References**

1. Gao, Y. et al. Complex transcriptional modulation with orthogonal and inducible dCas9 regulators. *Nat. Methods* 13, 1043–1049 (2016).
2. Kiani, S. et al. Cas9 gRNA engineering for genome editing, activation and repression. *Nat. Methods* 12, 1051–1054 (2015).
3. Kiani, S. et al. CRISPR transcriptional repression devices and layered circuits in mammalian cells. *Nat. Methods* 11, 723–726 (2014).
4. Liu, Y. et al. Synthesizing AND gate genetic circuits based on CRISPR-Cas9 for identification of bladder cancer cells. *Nat. Commun.* 5, 5393 (2014).
5. Qi, L. S. et al. Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. *Cell* 152, 1173–1183 (2013).
6. Nielsen, A. A. K. & Voigt, C. A. Multi-input CRISPR/Cas genetic circuits that interface host regulatory networks. *Mol. Syst. Biol.* 10, 763 (2014).
7. Gyorgy, A. et al. Tocost lines describe the cellular economy of genetic circuits. *Biophys. J.* 109, 639–646 (2015).
8. Xu, X. & Qi, L. S. A CRISPR–dCas9 toolbox for genetic engineering and synthetic biology. *J. Mol. Biol.* 431, 34–47 (2019).
9. Zhang, S. & Voigt, C. A. Engineered dCas9 with reduced toxicity in bacteria: implications for genetic circuit design. *Nucleic Acids Res.* 46, 11115–11125 (2018).
10. Fontana, J., Dong, C., Ham, J. Y., Zalatan, J. G. & Carothers, J. M. Regulated expression of sgRNAs tunes CRISPRi in E. coli. *Biotechnol. J.* 13, 1800069 (2018).
11. Clamons, S. & Murray, R. Modeling predicts that CRISPR-based activators, unlike CRISPR-based repressors, scale well with increasing gene competition and dCas9 bottlenecking. bioRxiv https://doi.org/10.1101/719278 (2019).

12. Chen, P.-Y., Qian, Y. & Del Vecchio, D. A model for resource competition in CRISPR-mediated gene repression. In 2018 IEEE Conference on Decision and Control (CDC) (IEEE, 2018).

13. Depardieu, F. & Bikard, D. Gene silencing with CRISPRi in bacteria and optimization of dCas9 expression levels. Methods 172, 61–75 (2020).

14. Cho, S. & et al. High-level dCas9 expression induces abnormal cell morphology in Escherichia coli. ACS Synth. Biol. 7, 1085–1094 (2018).

15. Gander, M. W., Vrana, J. D., Voje, W. E., Carothers, J. M. & Klavins, E. Digital logic circuits in yeast with CRISPR-dCas9 NOR gates. Nat. Commun. 8, 1–11 (2017).

16. Kim, B., Kim, H. J. & Lee, S. J. Regulation of microbial metabolic rates using CRISPR interference with expanded PAM sequences. Front. Microbiol. 11, 282 (2020).

17. Bellato, M. et al. CRISPR interference as low burden logic inverters in synthetic circuits: characterization and tuning. bioRxiv https://doi.org/10.1101/2020.08.33.248096 (2020).

18. Didovyk, A., Borek, B., Hasty, J. & Tsimring, L. Orthogonal modular gene repression in Escherichia coli using engineered CRISPR/cas9. ACS Synth. Biol. 5, 81–88 (2016).

19. Weinberg, B. H. et al. Large-scale design of robust genetic circuits with multiple inputs and outputs for mammalian cells. Nat. Biotechnol. 35, 453–462 (2017).

20. Qian, Y., Huang, H.-H., Jiménez, J. I. & Del Vecchio, D. Resource competition shapes the response of genetic circuits. ACS Synth. Biol. 6, 1263–1272 (2017).

21. Pasotti, L. et al. Re-using biological devices: a model-aided analysis of interconnected transcriptional cascades designed from the bottom-up. J. Biol. Eng. 11, 50 (2017).

22. Jones, R. D. et al. An endoribonuclease-based feedforward controller for decoupling resource-limited genetic modules in mammalian cells. Nat. Commun. 11, 5690 (2020).

23. Frei, T. et al. Characterization and mitigation of gene expression burden in mammalian cells. Nat. Commun. 11, 4641 (2020).

24. Huang, H.-H., Qian, Y. & Del Vecchio, D. A quasi-integral controller for adaptation of genetic modules to variable ribosome demand. Nat. Commun. 9, 5415 (2018).

25. Shopera, T., He, L., Oyetunde, T., Tang, Y. J. & Moon, T. S. Decoupling resource-coupled gene expression in living cells. ACS Synth. Biol. 6, 1596–1604 (2017).

26. Darlington, A. P. S., Kim, J., Jiménez, J. I. & Bates, D. G. Dynamic allocation of orthogonal ribosomes facilitates uncoupling of co-expressed genes. Nat. Commun. 9, 695 (2018).

27. Ceroni, F. et al. Burden-driven feedback control of gene expression. Nat. Methods 15, 387–393 (2018).

28. Reis, A. C. et al. Simultaneous repression of multiple bacterial genes using nonrepetitive extra-long sgRNA arrays. Nat. Biotechnol. 37, 1294–1301 (2019).

29. Pickar-Oliver, A. & Gersbach, C. A. The next generation of CRISPR-Cas technologies and applications. Nat. Rev. Mol. Cell Biol. 20, 490–507 (2019).

30. McBride, C. & Del Vecchio, D. Trade-offs in robustness to perturbations of bacterial population controllers. In 2020 American Control Conference (ACC). (IEEE, 2020).

31. Santos-Moreno, J., Tassioli, E., Stelling, J. & Schauerli, Y. Multistable and dynamic CRISPRi-based synthetic circuits. Nat. Commun. 11, 2746 (2020).

32. Zhang, Q. et al. CRISPRi-based dynamic control of carbon flow for efficient N-acetyl glucosamine production and its metabolomic effects in Escherichia coli. J. Agric. Food Chem. 68, 3203–3213 (2020).

33. Gibson, D. G. et al. Enzymatic assembly of DNA molecules up to several hundred kilobases. Nat. Methods 6, 343–345 (2009).

Acknowledgements
The authors want to thank Pin-Yi Chen (Massachusetts Institute of Technology, USA) for the modeling insights throughout the initial phase of the study and Michela Casanova (University of Pavia, Italy) for the assistance with preliminary experiments and cloning technology. This work was funded by NSF-CMMI Award #1727189 and NSF-CCF Award #2007674.

Author contributions
H.H., M.B., and Y.Q. designed constructs and performed experiments. P.C. and H.H. devised the auxiliary pAUX_OL plasmid. Y.Q. and M.B. performed simulations and mathematical analyses. D.D.V., M.B., Y.Q., and H.H. wrote the paper and analyzed the data. P.M. and L.P. supervised the activities of M.B.; D.D.V. designed the research.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-21772-6.

Correspondence and requests for materials should be addressed to D.D.V.

Peer review information Nature Communications thanks the anonymous reviewer(s) for their contribution to the peer review of this work. Peer reviewer reports are available.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.