Synthetic Biology-

A Sensitive Switch for Visualizing Natural Gene Silencing in Single Cells

Karmella A. Haynes,*^{,†} Francesca Ceroni,[‡] Daniel Flicker,[§] Andrew Younger,[§] and Pamela A. Silver^{§,||}

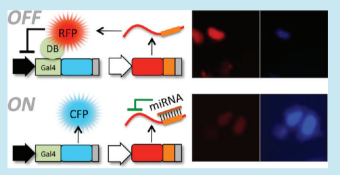
[†]School of Biological and Health Systems Engineering, Arizona State University, Tempe, Arizona 85287, United States [‡]Laboratory of Cellular and Molecular Engineering, University of Bologna, I-47521 Cesena, Italy

[§]Department of Systems Biology, Harvard Medical School, Boston, Massachusetts 02115, United States

^{II}The Wyss Institute for Biologically Inspired Engineering, Harvard Medical School, Boston, Massachusetts 02115, United States

Supporting Information

ABSTRACT: RNA interference is a natural gene expression silencing system that appears throughout the tree of life. As the list of cellular processes linked to RNAi grows, so does the demand for tools to accurately measure RNAi dynamics in living cells. We engineered a synthetic RNAi sensor that converts this negative regulatory signal into a positive output in living mammalian cells, thereby allowing increased sensitivity and activation. Furthermore, the circuit's modular design allows potentially any microRNA of interest to be detected. We demonstrated that the circuit responds to an artificial microRNA and becomes activated when the RNAi target is replaced by a natural microRNA target (miR-34) in



U2OS osteosarcoma cells. Our studies extend the application of rationally designed synthetic switches to RNAi, providing a sensitive way to visualize the dynamics of RNAi activity rather than just the presence of miRNA molecules.

KEYWORDS: genetic switch, synthetic repressor, RNA interference, miR-34

T ranslational application of sophisticated synthetic devices for medicine and basic research is enabled by enhancing the synthetic biology toolkit with parts from mammalian cells,^{1,2} cellular mechanisms that play an important role in normal cell development and disease. Recently, functional synthetic gene switches have been successfully designed for applications in human health.^{3,4} Similar artificial networks could be designed to report important dynamic cellular mechanisms that play a role in normal cell development and disease.

RNA interference (RNAi) is linked to essential processes including cell cycle progression,^{5–7} cellular differentiation,⁸ apoptosis,⁹ and cancer development¹⁰ in myriad organisms. One class of small noncoding RNAs acts on the posttranscriptional level to inhibit translation from target messenger RNAs (mRNAs). MicroRNAs (miRNAs) are transcribed in the cell nucleus as long precursor molecules (pri-miRNA), processed into hairpin RNAs (pre-miRNA) by the protein Drosha11, and exported into the cytoplasm. Pre-miRNAs are cleaved, by the protein Dicer, into a RISC–miRNA complex that can then bind target mRNAs and inhibit translation.

Recent studies are beginning to connect the dynamics of miRNA expression with cellular and tissue phenotypes, advancing our knowledge of RNAi beyond a collection of data that show which miRNAs are present in a tissue at a fixed point in time. So far, time-course miRNA expression profiles have been created for different human developmental processes. The accumulation of a set of miRNAs including miR-1, miR-133a, miR-133b, and miR-206 has been observed during natural human fetal muscle development, as well as during artificial induction of muscle cell differentiation in situ.¹² A key step in neural development is also linked to a dynamic miRNA expression profile, where differentiation-associated miRNAs accumulate and proliferation-regulating miRNAs decrease upon Schwann cell maturation.¹³ Periodic expression of some miRNAs appears to rely upon the circadian clock machinery¹⁴ and in some cases may regulate the clock itself.¹⁵ Rapid reporter systems that track closely with miRNA activity in real time will enable us to discover to what extent miRNA timing has a biological role.

In the studies cited above, populations of cells or tissues were collected at specific time points and lysed for RNA analyses. Other recent studies have aimed to generate expression profiles *in vivo* by placing a fluorescence-producing gene (i.e., GFP) under the control of miRNA or short interfering RNA (siRNA) so that the loss of expression indicates RNAi activity.^{16–18} Using loss of signal as a proxy for RNAi has several disadvantages. Photobleaching or cell division in addition to RNAi can contribute to signal depletion, confounding the results of time-course experiments. Moreover, clear loss-of-

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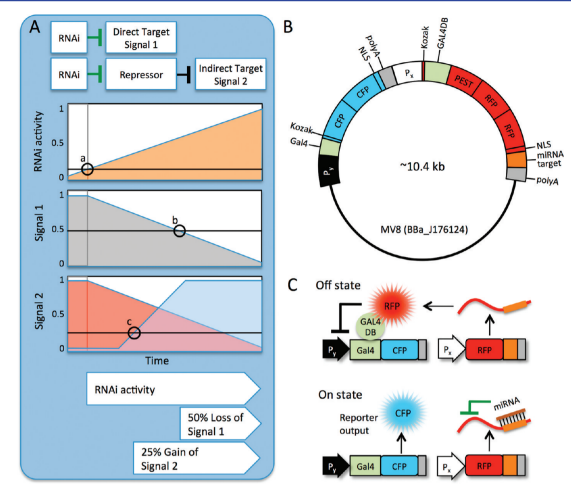


Figure 1. Schematic diagram of the mammalian miRNA reporter circuit. (A) Qualitative comparison of two approaches for detecting RNAi, loss of signal vs accumulation of signal. Signal 1 is directly repressed by RNAi. Signal 2 is repressed by a protein that is deactivated by RNAi. Once RNAi activity reaches an arbitrary threshold "a," target protein degradation exceeds production. When Repressor concentration is insufficient to repress expression, signal 2 accumulates ("c") to a visible level sooner than signal 1 is depleted by 50% ("b"). (B) General construct map. For all experiments, both genes were placed on the same vector. P_y and P_x = constitutive promoters driving expression of the CFP-Reporter and RFP-Repressor genes, respectively; NLS = nuclear localization sequence; CFP = AmCyan; RFP = mCherry; polyA = bovine growth hormone polyadenylation signal; PEST = protein degradation signal. (C) In the absence of miRNA, RFP-Repressor is expressed and inhibits CFP (Off). When a miRNA recognizes its specific target site within the 3'-UTR of the *RFP-Repressor* mRNA transcript, the Repressor protein is degraded, and CFP is expressed (On).

signal can be delayed by the slow degradation of stable fluorescent proteins.¹⁹ Lastly, natural miRNAs with low activity levels may not efficiently knock down highly expressed reporters. Ideally, RNAi reporters should be fast, sensitive, and produce a positive output.

We have engineered a synthetic reporter that converts RNAimediated gene silencing into a positive, visible signal on the order of hours in single living cells. We built a new genetic circuit based in part on the double repression scheme previously used by others.²⁰ Thus far, such circuits have used strong synthetic gene promoters to create cell-type classifiers that detect high levels of miRNA.⁴ Our device extends the range RNAi detection for synthetic circuits. We engineered a constitutive human promoter that can be held in the initial inactivate state without relying on an overwhelmingly strong repressor, giving the device the sensitivity to detect low levels of miRNA near the onset of RNAi. The double repression scheme places an output gene under the control of a repressor protein, which is targeted by a specific miRNA. This double repression approach links a gain of signal to RNAi activity. Thus, the observer can look for an accumulation of signal above a

negative background level, which can be seen more readily than a loss of signal relative to a positive background (Figure 1A). The sooner the RNAi signal is seen, the more closely the reporter tracks with RNAi in real time, and the more useful the reporter is for live cell analysis.

Our genetic circuit has two states, which we qualitatively describe as "Off" and "On." In the Off state, a cyan fluorescent signal is repressed. In the On state, the cyan fluorescent signal becomes activated. The circuit has a modular design (Figure 1B), which in principle allows us to simply choose a corresponding target site for different miRNAs and could be readily switched to On in the presence of a miRNA of interest. The Off state is maintained by a double RFP-tagged protein (RFP-Repressor) that binds downstream of the CFP-Reporter promoter. The RFP-Repressor is placed under the control of miRNA such that miRNA expression leads to repressor silencing and activation of the reporter (Figure 1C). Gene circuit components were optimized by identifying a combination of constitutive promoters for the CFP-Reporter and the RFP-Repressor that established the greatest ratio between the active cyan and repressed cyan states. A synthetic RNAi system

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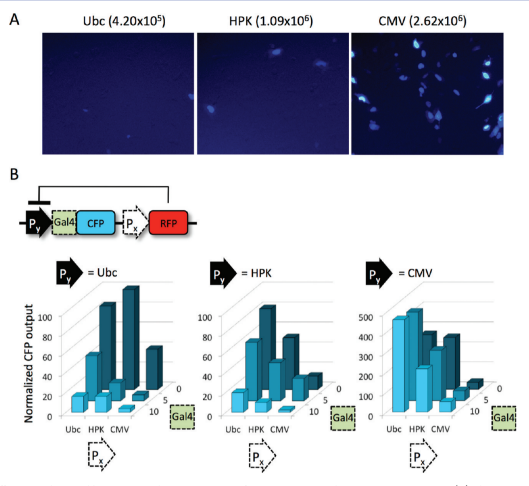


Figure 2. The off state can be tuned by increasing the concentration of RFP-Repressor at the output gene promoter. (A) Fluorescence microscopy of cells carrying CFP driven by a *Ubc, HPK,* or *CMV* promoter. Numbers show total CFP intensity (mean CFP fluorescence in CFP-positive cells, multiplied by the frequency of CFP-positive cells) from flow cytometry. (B) *RFP-Repressor* expression levels were regulated by *Ubc, HPK,* and *CMV* promoters (P_x), respectively. The local concentration of RFP-Repressor at the CFP promoter (P_y) was modulated by inserting 0, 5, or 10 copies of RFP-Repressor binding sites (Gal4) downstream of the P_y transcription start site. "Normalized CFP output" is the CFP intensity (mean CFP fluorescence in CFP/RFP-positive cells, multiplied by the frequency of CFP-positive cells) divided by the corresponding values shown in part A (*i.e.*, P_y = Ubc values/Ubc 4.2 × 105; P_y = HPK values/HPK 1.09 × 106; P_y = CMV values/CMV 2.6 × 106). Each column is the average of triplicate flow cytometry experiments. Standard error is less than 25% of the CFP value with the exception of 44% for P_x = HPK, P_y = Ubc, Gal4 = 5, and 31% for P_x = CMV, P_y = Ubc, Gal4 = 10. The arithmetic mean of the fluorescence was used for all calculations.

(miR-luc) was used to test the performance of the device in cultured cells. We then added natural miRNA target sites to detect miR-34 in U2OS osteosarcoma cells. Our sensor system has produced the first evidence of cell-cycle-arrest associated miR-34 activity in unstressed U2OS cells.

RESULTS

Regulating Transcription by Steric Hindrance from a Fluorescent Protein. The RFP-Repressor is a novel synthetic transcriptional repressor that binds near a promoter to occlude the transcriptional machinery through steric hindrance. Most of its bulk comes from two fluorophores (~54 kD); it does not rely upon a transcriptional repression domain to maintain gene repression. Thus, its promoter-silencing activity is essentially a function of repressor protein production and degradation, not epigenetic silencing effects. For instance, previously reported synthetic genetic switches utilize the Krüppel associated box (KRAB) repression domain.^{20,21} KRAB interacts with heterochromatin components such as HP1, which recruit chromatin modifying enzymes that form mitotically stable gene repression.²² We aimed to engineer fast reporter activation, where repression is reversed on the order of minutes or hours, before cell division occurs. Therefore, we used steric hindrance, reminiscent of the TetR and LacI regulation systems,^{23,24} to compete with PolII at the promoter and block transcriptional activation. Instead of tagging a full-length repressor with a fluorescent protein, we fused two mCherry monomers to the Gal4 DNA-binding domain from yeast to create a visible repressor. We included a PEST sequence motif to decrease repressor protein half-life when RNAi is active.

We optimized the Off state of the circuit by testing increasingly active constitutive promoters to drive RFP-Repressor expression. The relative strengths of the promoters we used are *Ubiquitin C* (*Ubc*) < *human phosphoglycerate kinase* (*HPK*) < *cytomegalovirus* (*CMV*), based on fluorescence microscopy and flow cytometry of cells transfected with CFP driven by each promoter (Figure 2A). In constructs, where a weak CFP promoter (P_y = *Ubc*) had five Gal4 repressor binding sites (Gal4 = 5), we observed stronger CFP repression (relative to controls with zero binding sites, Gal4 = 0) as RFP-Repressor expression was increased (Figure 2B): 0.8-, 2.5-, and 2.7-fold repression for RFP-Repressor promoters (P_x) *Ubc, HPK*, and

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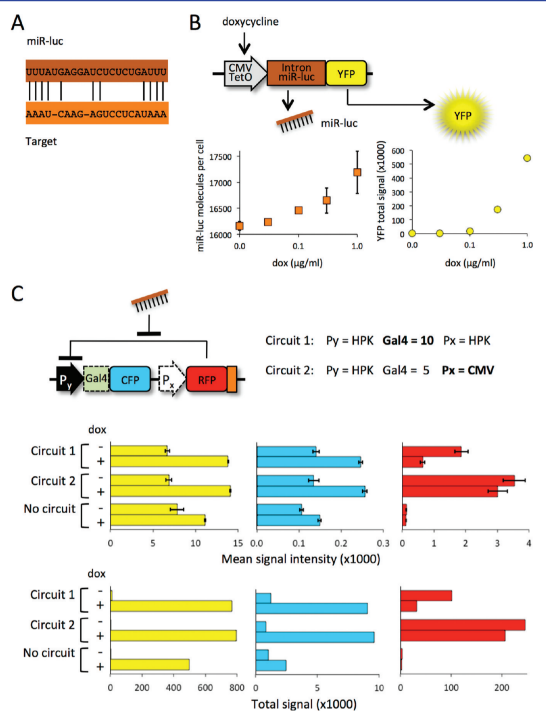


Figure 3. The miRNA sensor responds to an artificial miRNA. (A) YFP signal indicates the level of doxycycline-induced miR-luc in JDS33 cells. MiR-luc molecules per cell was measured by real time quantitiave RT-PCR. YFP signal was detected by flow cytometry analysis. YFP total signal = mean fluorescence \times frequency of fluorescence signal. (B) Comparison of two miRNA sensor designs, one with 10 binding sites for the *RFP*-*Repressor* (circuit 1) vs one with higher *RFP-Repressor* expression (circuit 2). YFP, CFP, and RFP signals (left to right) were measured by flow cytometry. Top graphs: error bars show the standard error of the mean signal intensity. Bottom graphs: total signal (mean signal intensity \times frequency of fluorescence signal). The arithmetic mean of the fluorescence was used for all calculations.

CMV, respectively. Switching the CFP promoter to HPK appeared to compromise the effectiveness of the repressor (0.46- and 0.44-fold repression for RFP-Repressor promoters *Ubc* and *HPK*). The *CMV*-driven CFP did not appear to respond to the repressor, compared to corresponding zero-Gal4 controls. Interestingly, the *CMV*-driven RFP-Repressor appears to decrease CFP levels when there are no repressor binding sites at the promoter of the *CFP*-Reporter (where $P_x = CMV$

and Gal4 = 0). This suggests that CMV on one plasmid may indirectly affect expression from the other plasmid, perhaps through competition for a fixed pool of transcription factors in the cell.

We also considered the possibility that CMV might produce too much RFP-Repressor, the direct target of RNAi in this circuit, and overwhelm the natural silencing machinery. Therefore, we altered the binding sites to increase the local

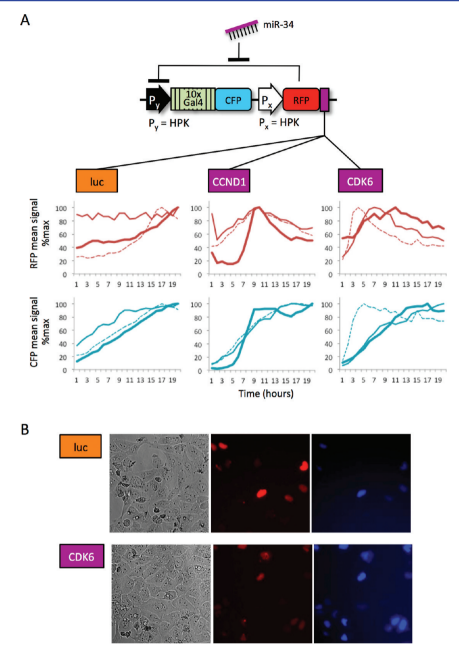


Figure 4. The behavior of natural microRNA sensors indicates miR-34 activity in U2OS cells. (A) Live cells carrying one of three miR sensor circuits with different miR target sites (CCND1, CDK6, or luc) were imaged via fluorescence microscopy over a \sim 30 h period. Each thick, thin, or dashed line corresponds to the same single cell in the RFP (top) and CFP (bottom) channel. RFP intensity and CFP output were measured as the mean signal of an individual (within a single cell cycle). In cells where RFP-Repressor is targeted by the miR-34 silencer (CCND1 and CDK6), a rise and then fall in RFP signal is associated with faster accumulation of CFP output, compared to a negative control where RFP-Repressor is not targeted by miR-34 (luc). (B) The addition of a PEST tag to CFP reduces accumulation of CFP expression from a sensor, whereas the sensor that carries the CDK6 target is activated. Phase contrast, RFP, and CFP channels are shown from left to right.

concentration of RFP-Repressor protein at the *CFP-Reporter* promoter. We found that doubling the number of repressor binding sites (from 5 to 10 copies of Gal4) achieved greater repression (Figure 2B). For instance, in constructs where a weak CFP promoter ($P_y = Ubc$) was suppressed by an *HPK*-driven RFP-Repressor ($P_x = HPK$), we observed 2.5- and 2.7-fold repression with 5 and 10 binding sites, respectively (Figure 2B). Repression of *HPK*-driven CFP was 0.4- and 2.4-fold with 5 and 10 binding sites. We chose the *HPK*-CFP-Reporter construct since expression of CFP from the weak *Ubc* promoter was difficult to visualize via microscopy (Figure 2A). This analysis allowed us to identify an optimal design ($P_x = HPK$, P_y

= HPK, Gal4 = 10), where CFP repression can be achieved without high CMV-driven expression of the repressor.

The Sensor Is Activated upon miRNA Induction. We demonstrated that the miR-sensor produces an increased CFP signal in the presence of miRNA by using a drug-inducible orthogonal RNAi system. The JDS33 cell line²⁵ carries a transgene that expresses luciferase miRNA (Luc-1601,²⁶ referred to hereafter as "miR-luc") when doxycycline (dox) is added to the cell culture medium. miR-luc has imperfect complementarity with the target sequences at the 3' end of the RFP-Repressor gene (Figure 3A), which is characteristic of natural miRNA regulation. Coexpressed visible YFP indicated

the amount of miR-luc present, as confirmed by real time PCR analysis and flow cytometry (Figure 3B). Therefore, the miR sensor should become activated when YFP is expressed.

We observed that the circuit becomes activated in the presence of miR-luc. Two different circuit configurations were compared. Circuit 1 had 10 Gal4 sites at the CFP-Reporter and an HPK (medium strength) promoter driving the RFP-Repressor. Circuit 2 had five Gal4 sites at the CFP-Reporter and a CMV (strong) promoter driving the RFP-Repressor (Figure 3C). We transfected JDS33 cells with each circuit and treated the cells with 1 μ g/mL dox for 48 h. About 1.7 × 10⁴ molecules of miR-luc per cell are present under these conditions (Figure 3B). Shifts from a low to a higher CFP signal showed that both sensors became activated after doxinduced miR-luc expression (Figure 3C). Activation of circuit 1 (*HPK*-driven RFP-Repressor); thus a switch from Off to On does not require an overexpressed repressor.

A Sensor Designed to Detect miR-34 Is Activated in Human Osteosarcoma Cells over a Short Time Scale. miRNA sensors carrying target sequences for the natural miRNA miR-34 become activated in the human bone cancerderived cell line U2OS. miR-34 plays an important role in cell physiology. miR-34 silences cyclinD1 (CCDN1), cyclin dependent kinase 6 (CDK6), and B-cell lymphoma 2 (BCL2),⁹ which promote progression into mitosis and prevent apoptosis. Thus, miR-34 activity has a negative impact on cell proliferation and is expected to be diminished or absent in dividing cancer cells. miR-34 has been detected in U2OS at low or high levels, depending upon a proliferating or arrested growth status, respectively.²⁷ We used our system to track miR-34 activity over a time interval of approximately one cell division (20 h) in proliferating U2OS cells with potentially low levels of miR-34.

Sensors built with miR-34 target sites from CCND1 and CDK6, but not BCL2, showed switch-like activation in U2OS cells. All three sensors were constructed on the basis of the configuration for circuit 1 (Figure 3C), transfected into U2OS cells, and analyzed 6 h later via live cell time-lapse microscopy. Circuit 1, without miR-34 target sites, was used as a negative control since it is not expected to become activated in U2OS cells that do not express artificial miR-luc. The negative control showed a gradual linear accumulation of both signals over the time frame analyzed. The BCL2 sensor behaved the same as the negative control construct (data not shown). In contrast, the CCDN1 and CDK6 sensors showed a rise and fall in RFP signal, accompanied by an increase in CFP production (Figure 4A). The first derivatives of regression curves for CFP expression show higher peaks at earlier time points for the miR-34-regulated sensors (Figure S1). These results indicate that for time-course single cell analysis performed in under 20 h, activated sensors can be distinguished by more rapid accumulation of CFP signal.

Computational analysis of RNA hybridization suggests that the affinity of miR-34 for each of the targets may determine the differences in the behavior of the sensors we tested. We calculated the free energy of binding between CCND1, CDK6, and BCL2 target sites and each member of the miR-34 family (miR-34a, b, and c). Unresponsiveness of the BCL2 sensor might be a consequence of this target's low affinity for miR-34, as indicated by relatively high free energies of the predicted miR-34-target hybrids (Figure S2).

We also observed activation of a redesigned miR-34 sensor in which CFP was destabilized to reduce the signal detected in the Off state. The accumulation of both RFP and CFP expression from the construct that was designed to be constitutively off (lacking miR-34 target sites) was unexpected, since our previous analyses indicated an inverse relationship between RFP and CFP (Figure 4A). The time-course results may be due to some instability of the Off state before proteins reach steady state levels. We reduced accumulation of the CFP signal in the inactive state by adding a PEST protein degradation tag to CFP (in circuit configuration $P_x = HPK$, $P_y = HPK$, Gal4 = 10). CFP destabilization does not appear to improve the signal to the background ratio (0.4-fold, Figure 4B) compared to the circuit with stable CFP (2.4-fold, Figure 2B). However, cells carrying the miR-34 sensor with destabilized CFP did show a greater CFP signal than a sensor that lacked the miR-34 target sites (Figure 4B).

Conclusion. Our work presents two important advances for synthetic device engineering and applications. First, the RFP-Repressor may serve as a generally useful model for designing synthetic transcriptional regulators *de novo*. Other commonly used repressors (i.e., LacI and Tet) are derived from naturally occurring gene repression systems. To our knowledge, the RFP-Repressor is the first reported functional synthetic repressor that has been built entirely from protein modules that are not typically associated with transcriptional repressor may be built from a protein designed simply to occupy sites immediately downstream of a constitutive promoter and that its function depends upon the local concentration of repressors at the promoter.

Second, we have extended the application of the double repression circuit design to sense miRNAs previously detected at low levels in the proliferating cell state,²⁷ over a short time scale. miR sensor activation may be sensitive to low levels of miR-34, or activated sensors might signify the onset of miR-34 activity prior to subsequent cell cycle arrest. For medical applications in cancer therapy, our circuit design could be engineered to activate a cancer-killing gene near the onset of metastasis-associated miRNA activity. So far, we have expressed the switches from extra-chromosomal plasmids. Future applications that stably integrate the circuit into the genome will be useful for observing the stochasticity of miR-34 activity at the individual cell level, which is important for determining whether single cancer cells alter states to become more aggressive. Our work demonstrates the power and potential of modular rational design in extending the utility synthetic devices for research and medical applications.

METHODS

Constructs. DNA fragments with universal cloning sites (*EcoRI*, *NotI*, *XbaI*, *SpeI*, and *PstI*) were generated by PCR and assembled according to a modified BioBrick DNA assembly method,²⁸ using *E. coli* DH5-alpha maintained under standard conditions. All DNA fragment and annotated full-length construct sequences are freely available in the MIT Registry of Standard Biological Parts²⁹ (See Table S1). The RFP-Repressor was constructed from a Gal4 DNA-binding domain followed by two copies of mCherry. The constitutive promoters *Ubc*, *HPK*, and *CMV* were cloned upstream of five (5xGal4) or 10 (10xGal4) copies of the *Gal4* DNA binding sites to generate Gal4-regulated promoters. Complete constructs were excised from the V0120 high-copy cloning vector and inserted into

modified pcDNA 3.1+ vectors. All plasmids were verified by restriction digests and DNA sequencing (Genewiz, Inc., Cambridge, MA) from the vector into the 5' and 3' ends of the inserts prior to transfection.

Cell Culture and Transfections. Cell line JDS33 is described by Shih et al.²⁵ U2OS and JDS33 cells were grown in McCoy's 5A medium supplemented with 10% tetracyclinefree fetal bovine serum and 1% penicillin and streptomycin. Cells were grown at 37 °C in a humidified CO₂ incubator. For transfections, plasmid DNA was extracted and purified from *E. coli* using a QIAGEN miniprep kit. A total of 1 μ g of plasmid DNA, 3 μ L Lipofectamine LTX (Invitrogen), and 3 μ L PLUS reagent in 190 μ L Opti-MEM was added to ~2.5 × 10⁵ cells per well (in a 12-well plate) in antibiotic-free growth medium.

Flow Cytometry. Cells were grown in a 12-well plate (~5 \times 10⁵ per sample/well), trypsinized, collected in growth medium, and washed and resuspended in 1× Dulbecco's PBS. Flow cytometry was performed 18 h after transfection using a BD Biosciences LSRII HTS-3 laser high throughput sampler platform. For flow cytometry analyses to detect mCherry, YFP (Venus), and AmCyan, the following excitation lasers and filters were used: 594 nm Yellow/DsRed (620/22), 488 nm Blue/FITC (520/50), and 405 nm Violet/AmCyan (525/50). For every experiment, nonfluorescent U2OS cells were used to adjust the voltage to minimize the autofluorescence signal in all three channels and to create a 1:1 forward and side scatter ratio. Background was subtracted from the signal by setting the threshold gate above the signal from nonfluorescent U2OS cells. Data were visualized using FACS Diva software and statistically analyzed using FlowJo 9.3.3 software.

Quantitative Real-Time Reverse Transcription PCR of miR-luc and YFP Detection. JDS33 cells were cultured in 6well plates ($\sim 1 \times 10^6$ per sample/well) and treated with 0, 0.01, 0.03, 0.1, 0.3, or 1 μ g/mL doxycycline to induce YFP and miR-luc expression for 48 h. One set of samples was harvested for YFP detection via flow cytometry. A duplicate set was processed for RNA isolation. RNA was extracted with TRIzol Reagent (Invitrogen 15596-018) and purified according to the Invitrogen protocol. cDNA synthesis and PCR were designed as described in Raymond et al.³⁰ cDNA synthesis (SuperScript III, Invitrogen 18080-051) was performed according to the manufacturer's protocol with either 5 μ g of template RNA from the dox-treated JDS33 cells (Experimental) or 2 pmol of synthesized "Input Control" RNA oligos (IDT DNA), plus 50 pmol of miR-luc "tailed" primers (5'-catgatcagctgggccaagaaaatcagagag) instead of oligo dT in a final reaction volume of 20 μ L. To normalize for Experimental RNA loading, oligo dT primers were used in separate reactions to generate Total cDNA for measuring GAPDH transcripts. Completed reactions were treated with 1 μ L of RNaseH.

qRT-PCR was performed on Experimental cDNA, Input Control cDNA, or Total cDNA. Experimental and Input Control cDNA reactions contained 0.8 μ L of cDNA, 36.0 pmol of primers (miR-luc forward 5'-tttatgaggatctctct; miR-luc reverse 5'-catgatcagctgggcca), and 7.5 μ L of SYBR Green Master Mix (Applied Biosystems) in a final volume of 15 μ L. GAPDH reactions contained 0.8 μ L of Total cDNA and 2.25 × 10⁻³ pmol of primers (GAPDH forward 5'ccgcatcttcttttgcgtcgcc; GAPDH reverse 5'-accaggcgcccaatacgacc). For standardization, we used a synthesized DNA oligo template Standard that carries the full miR-luc PCR template sequence. qRT-PCR was performed as described above using 0.8 uL of 1.0 × 10⁻⁵, × 10⁻⁴, × 10⁻³, or × 10⁻² uM synthesized DNA solution and 36.0 pmol of primers (miR-luc forward and miR-luc reverse).

Fold change relative to the least abundant standard template was caluclated as $2^{\wedge}(Ct_{Standard 1} - Ct_{sample})$ for all samples. DNA oligo molecules per reaction for standards 1-4 ($4.816 \times 10^{6}, \times 10^{7}, \times 10^{8}$, and $\times 10^{9}$) was plotted against fold change to generate a standard curve. The line of best fit equation was used to calculate miR-luc molecules per reaction for the Experimental cDNA samples; these values were divided by Total cDNA GAPDH normalization values ($2^{\wedge}(Ct_{GAPDHsample}/Ct_{GAPDHodox})$) and adjusted to take into account 73% cDNA synthesis efficiency (as indicated by the Input Control values). "miR-luc molecules per reaction divided as normalized Experimental miR-luc molecules per reaction divided by 8000 cells per reaction.

Single Cell Fluorescence Assays. U2OS cells were cultured in 12-well glass-bottom plates and transiently transfected with miRNA sensor circuit constructs for 6 h. Phase contrast, red fluorescent, and cyan fluorescent images were collected at 1 h intervals using a Nikon inverted microscope, controlled by Metamorph acquisition software. Mean fluorescence intensities of individual interphase nuclei were manually measured using ImageJ.

ASSOCIATED CONTENT

S Supporting Information

A detailed list of constructs as entries into the MIT Registry of Standard Biological Parts (Table S1), Figure S1, Figure S2, supplemental methods, and references. This information is available free of charge via the Internet at http://pubs.acs.org/.

AUTHOR INFORMATION

Corresponding Author

*E-mail: karmella.haynes@asu.edu.

Author Contributions

K.A.H. and P.A.S. conceptualized the project. K.A.H., F.C., D.F., and A.Y. built and transfected constructs and performed flow cytometry. K.A.H. and D.F. performed time-course microscopy. F.C. and A.Y. designed and performed miR-luc RT-PCR. D.F. performed the RNAhybrid analysis. All authors drafted the manuscript.

Notes

The authors declare no competing financial interest.

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