Synthetic Circuit-Driven Expression of Heterologous Enzymes for Disease Detection

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ABSTRACT: The integration of nanotechnology and synthetic biology could lay the framework for new classes of engineered biosensors that produce amplified readouts of disease states. As a proof-of-concept demonstration of this vision, here we present an engineered gene circuit that, in response to cancer-associated transcriptional deregulation, expresses heterologous enzyme biomarkers whose activity can be measured by nanoparticle sensors that generate amplified detection readouts. Specifically, we designed an AND-gate gene circuit that integrates the activity of two ovarian cancer-specific synthetic promoters to drive the expression of a heterologous protein output, secreted Tobacco Etch Virus (TEV) protease, exclusively from within tumor cells. Nanoparticle probes were engineered to carry a TEV-specific peptide substrate in order to measure the activity of the circuit-generated enzyme to yield amplified detection signals measurable in the urine or blood. We applied our integrated sense-and-respond system in a mouse model of disseminated ovarian cancer, where we demonstrated measurement of circuit-specific TEV protease activity both in vivo using exogenously administered nanoparticle sensors and ex vivo using quenched fluorescent probes. We envision that this work will lay the foundation for how synthetic biology and nanotechnology can be meaningfully integrated to achieve next-generation engineered biosensors.

KEYWORDS: biomarkers, proteases, activity probes, nanosensors, synthetic biology, nanotechnology, cancer

The computational principle of sense-and-respond is fundamental to engineering efforts that seek to build biological or chemical tools that can detect and report on the presence of disease. In synthetic biology, the forward engineering of biological systems toward this goal relies on the use of biological parts to create novel sense-and-respond functionalities inside living cells, enabling circuit-inspired control at the genetic, transcriptional, or protein levels.1−3 In nanoparticle engineering, material and chemical systems are treated as the programmable entity and can be engineered as multifunctional, cooperative systems that can detect and respond to molecular events at the nanoscale.4,5 In recent years, both these fields have seen the emergence of engineered, exogenous agents that can interrogate biological states in vivo to generate amplified readouts of disease.6 Specifically, several technologies, both in synthetic biology and nanotechnology, have leveraged enzymatic activity to measure or produce biomarkers of disease.7−11 Because they harness the unique substrate specificity and catalytic signal amplification properties of enzymes, these engineered activity-based diagnostics have the potential to overcome some of the sensitivity and specificity limitations of standard diagnostics. Within this sphere, the integration of synthetic biology with nanoparticle sensing tools could yield hybrid sensors that interface directly with the body to produce highly specific, amplified disease readouts.

Recent years have seen independent advances in the design of programmable diagnostics within each of synthetic biology and nanotechnology. Synthetic biology and genetic engineering tools have promoted the design and use of bacteriophages,15 molecular recorders,13 programmable probiotics,16 genetically encoded nanosensors,12, and mammalian cells10,17 for disease detection applications. For example, probiotic bacteria have been programmed to serve as delivery vectors that conditionally produce secreted enzymes to generate a noninvasive, signal amplified readout in urine.9 However, this system lacked any disease-specific sensing circuitry upstream of the activity-based enzymatic readout. To this end, molecular engineering tools have been leveraged to develop DNA vectors carrying a tumor-specific promoter driving expression of a reporter enzyme.13 More recently, macrophages were en-
engineered to secrete a heterologous reporter enzyme in response to adopting a tumor-associated (M2 macrophage) transcriptional profile, and were administered in mice with colorectal cancer to generate a sensitive, signal amplified disease readout that could be queried ex vivo in the blood. In addition, gene therapy delivery techniques, such as engineered lentiviral vectors, have recently been tested both preclinically and clinically for in vivo therapeutic applications in cancer and other diseases, and could enable the targeted delivery of synthetic disease-sensing gene circuits in vivo.

In parallel to these synthetic biology and gene therapy advances, nanoparticle sensors and chemical probes have been deployed to actively query disease activity in vivo to generate diagnostic readouts. Several such approaches convert the activity of enzymes, such as proteases, involved in disease progression into an imaging or biofluid-based detection readout. For example, activity-based nanosensors, an emerging biosensor class, sense dysregulated protease activity at the site of disease and respond by releasing a renal clearable synthetic reporter that enables detection of disease via a simple urine test. However, this and related approaches rely on endogenous protease dysregulation as the measured disease biomarker. Because proteases are expressed and active in healthy cells as well as in a variety of disease etiologies, this places a heavy engineering requirement to try to ensure sensor specificity.

When integrated with responsive nanoparticles or molecular probes, synthetic biology tools could open new diagnostic pathways by expanding the set of accessible input signals specific to a disease of interest, such as cancer. Namely, due to the variety of molecular parts available, such as synthetic promoters with enhanced cell-state specificity (SPECS), synthetic circuits based on transcriptional control are highly tunable and can be engineered to detect the altered transcriptional state present in cancer cells. In turn, the sensed transcriptional signal could be transduced into an easily detectable, biorthogonal output, such as a heterologous reporter enzyme, which could then trigger the activation of a nanoparticle sensor to ultimately produce a highly specific, amplified readout. More generally, despite the foundational principles common to both synthetic biology and nanotechnology-inspired biosensors, they have yet to be meaningfully integrated. Simultaneously harnessing the programmability of genetically controlled synthetic biology systems and the modular targeting, sensing, and reporting capabilities of nanomaterials could open new avenues in the design of engineered diagnostics and therapeutics for cancer and other diseases.

In this work, we explored how synthetic biology and nanoparticle sensors can be integrated to achieve a cooperative sense-and-respond system that produces amplified readouts of disease states. We engineered a synthetic gene circuit that, when integrated into cancer cells following lentiviral delivery, drives specific production of a heterologous enzyme biomarker by the modified tumor cells. The secreted heterologous enzyme in turn catalyzes highly specific cleavage reactions measurable by nanoparticle sensors, enabling noninvasive, amplified readouts in either the blood or urine. As a proof of concept, we deployed this system in a mouse model of disseminated ovarian cancer, a disease with a dismal mortality rate in which endogenous, blood-based biomarkers face significant sensitivity limitations. Our integrated system consisted of an initial synthetic biology-driven sensing phase followed by a nanotechnology-enabled detection phase. For the sensing phase, we engineered an RNA-based AND gate circuit that integrated the activity of two ovarian cancer SPECS via a Boolean AND gate and generated a biomarker output only when both promoters were simultaneously active. The circuit was delivered into ovarian tumors, where it could sense aberrant cancer-associated transcriptional activity and drive the tumor-specific expression of a heterologous biomarker, the Tobacco Etch Virus (TEV) protease. Because of its stringent cleavage preference and orthogonality to endogenous proteases in the body, TEV is particularly well suited for use as a heterologous biomarker. For the detection phase, nanoparticle probes carrying the cognate peptide substrate for TEV were used to measure the activity of this enzyme in vivo or ex vivo to generate amplified signals detectable in the urine or blood, respectively. This work presents a new paradigm for the integration of synthetic gene circuits and nanotechnology toward the development of engineered sense-and-respond diagnostic or therapeutic systems.

### RESULTS

**Sensing Ovarian Cancer-Associated Transcriptional Dysregulation with an Engineered Synthetic Circuit.** Our integrated sense-and-respond platform consists of (1) a synthetic circuit that drives tumor-specific expression of the heterologous biomarker TEV protease during the sensing step, and (2) two readout methods to measure the subsequent protease activity during the detection step (Figure 1). The input component of the circuit integrates the activity of two ovarian cancer SPECS (P1 and P2) to sense cancer-associated transcriptional activity in circuit-expressing cells and to ultimately produce secreted TEV protease (Figure 1a). To implement the synthetic circuit in vivo, we used three separate lentiviruses that encode the input modules (Module 1 and Module 2) and the output module (Module 3) to transduce ovarian cancer cells in tumor-bearing mice (Figure 1b). Sensing and transduction of the transcription factor input signals resulted in expression and secretion of TEV protease (Figure 1a). In turn, the activity of secreted TEV protease was subsequently queried either by responsive nanoparticles in vivo or by a Förster resonance energy transfer (FRET)-based detection assay ex vivo, to ultimately generate an amplified readout in either the urine or blood, respectively (Figure 1c).

To design the input modules for the sensing circuit, we used two SPECS, S(E2FJ)P and S(cMyc)P, which were previously demonstrated to be specific to the OVCAR8 ovarian cancer cell line, to transcribe the endoribonuclease Cys4 gene (Input 1) and a novel miRNA-based self-inhibitory gene module (Input 2), respectively (Figure 2a). The miRNA-based gene module is made of two separate exons for a fusion protein (GAD), consisting of a GAL4 DNA binding domain and the transcriptional transactivator VP16 flanking a synthetic miRNA sequence (miR1), a 28bp Cys4 binding site, and a downstream miR1 binding site sequence (Figure 2a). We defined four different states for the synthetic circuit according to the input status of each module (Figure S1): no inputs ([0,0]), one of Input 1 (S(E2FJ)P/Cys4; [1,0]) or Input 2 (S(cMyc)P/miR1; [0,1]), or both inputs ([1,1]). All four of these circuit states incorporate the output module, which drives expression of the output protein (GFP for optimization experiments; TEV protease for detection experiments) only when triggered by both inputs ([1,1]) (Figure 2a). In the [1,0]
We reasoned that the performance of this synthetic circuit would depend on how efficiently each module was delivered and expressed, and how accurately the input modules could sense the transcriptional landscape of transduced cancer cells. To streamline initial characterization of the circuit’s input sensing performance, we used a membrane GFP as output to simplify the analysis process. All four different cellular states (no input ([0,0]), only one input ([1,0] or [0,1]), or both inputs ([1,1])) were tested by transducing OVCAR8 cells \textit{in vitro} with different combinations of lentiviruses carrying individual circuit components (Figure 2). Cells were then fixed for immunofluorescence imaging and flow cytometry analysis to quantify the fraction of GFP-expressing cells and the mean fluorescent intensity of the population. At a multiplicity of infection (MOI) of approximately 5 for each virus, all the cells in the [1,1] state expressed GFP (data not shown), indicating that the synthetic circuit could be turned on in OVCAR8 cells. To avoid signal saturation caused by over infection, which may complicate the assessment of the synthetic circuit’s performance, we purposely infected the cells with a lower virus titer for each virus. As expected, cells in the [0,0] or [1,0] states displayed no GFP expression (Figure 2b,c). On the other hand, about 6% of cells in the [0,1] state and 22% of cells in the [1,1] state were GFP positive (Figure 2b, Figure S2). The 6% GFP positivity in the [0,1] state points to leaky transcription of the Input 2 cassette. Cells in the [1,1] state also had the highest mean fluorescent intensity, about 3.5 fold higher than that of the [0,1] state (Figure 2c). The nonzero frequency of GFP+ cells observed in the [0,1] state-treated cells further supports the hypothesis that the transcription of the Input 2 gene (GAD) was not entirely inhibited by the miRNA repressor module, and that further optimization is required to eliminate this leakiness. Nevertheless, the percent of GFP+ cells detected in the [0,1] state was significantly lower than that detected in the [1,1] state (Figure 2c, $P = 0.0236$).

Next, we sought to demonstrate the ability of the synthetic circuit to distinguish different cancer cell lines. The SPECS in the circuit were designed to bind with ovarian cancer-associated transcription factors enriched in OVCAR8 cells and could putatively be selected for particular input signals relevant to different disease applications. We have previously shown that these SPECS are highly active in OVCAR8 cells, but not in normal ovarian epithelial cells. To test whether these SPECS were specific to OVCAR8 relative to a select subset of cancer cell lines, we infected three additional ovarian cancer lines (ES-2, CaOV3, and OAW-42), a hepatocarcinoma cell line (Huh7), and a cervical cancer cell line (HeLa), with the input and output viruses ([1,1]). There was no detectable GFP expression in OAW-42, Huh7, or HeLa cells, and negligible GFP expression in ES-2 and CaOV3 cells (Figure 3d, $P = 0.0001$ for ES-2, $P = 0.0015$ for CaOV3; Figure S2). Altogether, these data suggest that the synthetic circuit was specific for the OVCAR8 cancer line relative to the cell lines assayed, highlighting the power of synthetic biology to tune circuit behavior based on specific cell phenotypes.

**Protease Reporter Module for Amplified Response and Readout.** Having confirmed that the input modules could specifically regulate the output of a fluorescent reporter protein, we set out to develop and validate the TEV protease-based response module. Because of the signal amplification properties of enzyme catalysis, we reasoned that measurements of TEV protease activity, rather than abundance, could yield
highly sensitive readouts. To that end, two assays were developed to measure TEV protease activity in vivo or ex vivo (Figure 1c). Both detection assays utilized similar designs as our previously developed protease-responsive systems. For the first assay, we engineered an exogenously administered nanoparticle capable of measuring TEV protease activity in vivo to release reporters that can be detected in the urine. A 40 kDa eight-arm poly(ethylene glycol) (PEG-8 40 kDa) nanoparticle was conjugated with a TEV peptide substrate bearing a ligand-encoded exogenous reporter (Figure 1c). The average size of these nanoparticles was about 20 nm, as characterized by transmission electron microscopy imaging (Figure S3), larger than the size limit of renal filtration. Following systemic administration, the nanosensor is disassembled in response to TEV protease activity, releasing ligand-encoded reporter peptides small enough to renally clear.

Figure 2. Design and validation of the synthetic sense-and-respond circuit. (a) Input 1 relies on a SPEC (S(E2F1)P) to drive the specific expression of endoribonuclease Csy4. Input 2 relies on a SPEC (S(cMyc)P) to drive the transcription of a fusion protein, GAD, consisting of the GAL4 DNA binding domain and the transcriptional transactivator VP16, and a miRNA transcript. The 3′ end of GAD transcripts has a hairpin structure for Csy4 binding and a miRNA binding site. When both inputs are present ([1,1] state), Csy4 will bind to the hairpin structure to release the inhibitory effect of the miRNA on GAD translation, driving the expression of GAD. The output expression is driven by a synthetic promoter GAL4BS, which is targeted by transcription activator-binding domain fusion, GAD. Two outputs are used in this study: membrane GFP (hTFR-GFP) or secreted TEV protease (SecTEV). (b) OVCAR8 cells were either transduced with the output virus carrying a membrane GFP expression cassette ([0,0]), or the output virus with one of two input viruses ([1,0], [0,1]), or both input viruses and the output virus ([1,1]). After 2 days, cells were fixed for immunofluorescence imaging or flow cytometry. Scale bar: 100 μm. (c) Quantification of flow cytometry analysis of mean GFP fluorescent intensity with different input and output configurations. Mean ± s.d.; N = 3; ordinary one-way ANOVA with Tukey’s correction for multiple comparisons, ***P = 0.0007 for [1,1] vs [0,0], **P = 0.5151 for [0,1] vs [0,0], ***P = 0.9717 for [1,0] vs [0,0]. (d) Human ovarian cancer cell lines (ES-2, OAW-42, CaOV-3, and OVCAR8), a hepatocarcinoma cell line (Huh7), and a cervical cancer cell line (HeLa) were transduced with both input viruses and the output virus with GFP output ([1,1]). After 2 days, cells were fixed for flow cytometry analysis. The mean GFP fluorescent intensity is shown. Mean ± s.d.; N = 3; unpaired two-tailed t test, ***P = 0.001 for ES-2, **P = 0.0015 for CaOV-3.
and accumulate in urine. Urinary reporter concentrations are then detected via immunoassay. In parallel, we also developed a detection assay to measure TEV protease activity in vivo in the blood. To this end, we designed a fluorogenic peptide reporter carrying a TEV cleavage site and measured TEV activity in the blood by monitoring fluorescence increase over time (Figure 1c).

We first assessed the function of our TEV peptide substrate in vitro by incubating a FRET-based sensor with purified enzymes or with cell-secreted TEV present in conditioned culture media (Figure 3a). First, we measured cleavage of the FRET-based reporter by monitoring fluorescence signal over time following incubation with purified enzymes and found that fluorescence intensity increased over time only in the presence of recombinant TEV protease, but not other proteases (Figure 3b). Next, we tested whether TEV protease maintained its cleavage activity following recombinant expression by mammalian cells. As TEV protease is usually expressed intracellularly, we inserted a N-terminal secretory signal peptide to produce a secreted version of the protein. The expressed intracellularly, we inserted a N-terminal secretory signal peptide to produce a secreted version of the protein. The expressed protease were screened against a FRET-paired TEV substrate, and fluorescence activation was monitored over time. (b) Kinetic fluorescence curves are shown for 1 μM FRET-paired reporter incubated with TEV (20 nM or 40 nM) as well as MMP2, MMP9, MMP13, thrombin, and tPA (all 20 nM), or buffer without protease (Buffer). Mean ± s.d.; N = 3. (c, d) Activity of cellular wild-type TEV, secreted wild-type TEV (WT SecTEV), and secreted, unglycosylated TEV (SecTEV QSG), present in cell culture media, against FRET-paired TEV substrate, thrombin, and uPA (all 20 nM), or buffer without protease (Buffer). Mean ± s.d.; N = 3. (c, d) Activity of cellular wild-type TEV, secreted wild-type TEV (WT SecTEV), and secreted, unglycosylated TEV (SecTEV QSG), present in cell culture media, against FRET-paired TEV substrate, showing cleavage kinetics (c) and fluorescence fold changes at 30 min (d). Mean ± s.d.; N = 2; unpaired two-tailed t test, *P = 0.0123 SecTEV QSG vs Cellular TEV, *P = 0.0106 for SecTEV QSG vs WT SecTEV.

**In Vitro Validation of the Integrated Sense-and-Respond Circuit.** Having validated the input (sensing) and output (response) portions of the synthetic circuit individually, we next tested whether the integrated circuit system could detect OVCAR8-associated transcriptional changes in vitro. Cultured OVCAR8 cells were transduced with various combinations of input viruses as in Figure 2, as well as with the output virus carrying the SecTEV QSG expression cassette. In this case, the SecTEV QSG cassette was further modified to express TEV protease tagged with the epitope V5 to facilitate immunodetection of the produced TEV protease. Two days post viral transduction, cells were fixed and permeabilized for immunofluorescence imaging or flow cytometry analysis via detection with anti-V5 antibody. Like our characterizations using membrane GFP as the output, V5-labeled SecTEV QSG was not detected in conditions transduced with state [0,0] or [1,0] lentiviruses (Figure 4a–c), and low levels were measured for state [0,1]. In contrast, state [1,1]-transduced cells were positive for SecTEV QSG via immunostaining for V5, as measured by both immunofluorescence imaging (Figure 4a) and flow cytometry (Figure 4b; Figure S4). Indeed, the [1,1]-transduced population exhibited a significantly greater percentage of TEV+ cells relative to all other state configurations (Figure 4c). Specifically, 55% of cells transduced with state [1,1] viruses expressed the TEV protease, based on V5 antigen expression, as compared to approximately
7% of cells exposed to state [0,1] inputs (Figure 4c). We defined the ON-OFF ratio for the input AND gate as the relative fold change in output mean intensity in the [1,1] state relative to [0,1]. The ON-OFF ratio for the SecTEV QSG output was 6 (Figure S4), with the output performance of this miRNA-based self-inhibitory AND gate on par with that of other previously reported circuits.29,34

Next, we analyzed the system’s sense-and-respond performance by testing whether the activity of SecTEV QSG produced by circuit-transduced OVCAR8 cells could be monitored in vivo. Activity was monitored as increase in fluorescence signal over time (d) as measured by the FRET-based reporter and quantified using fluorescence fold change at 120 min (e). Mean ± s.d.; N = 3; ordinary one-way ANOVA with Tukey’s correction for multiple comparisons, ****P < 0.0001. For comparison, [1,1] vs [0,0], **P = 0.0022; [1,1] vs [1,0], *P = 0.0015; [1,1] vs [0,1], *P = 0.0284.

Collectively, these results demonstrated that our integrated sense-and-respond system could generate OVCAR8-specific output expression, and that output activity levels could be easily detected in vitro.

In Vivo Tumor Detection through Activity-Based Readouts in the Urine or Blood. Following successful in vitro validation, we sought to apply the integrated sense-and-respond system for in vivo ovarian cancer detection using circuit readout assays in either urine or blood. A mouse model of disseminated ovarian cancer was generated by seeding OVCAR8 cells through intraperitoneal (i.p.) injection. Tumor-bearing mice were i.p. injected either with only the V5-tagged SecTEV QSG output virus ([0,0]) or with both input viruses and the output virus ([1,1]) (Figure 5a). To test whether the heterologous enzyme biomarker SecTEV QSG was being expressed in tumor cells, tumor nodules were resected and stained for V5. Tumor sections from mice infected with both input viruses and the output virus ([1,1]) revealed detectable TEV protein, while there was no positive staining in the control, output virus-only ([0,0]) group (Figure 5b). Approximately 10% of all resected tumor nodules from the
[1,1] group were positive for at least some TEV protease staining, and furthermore most cells in a representative TEV-positive nodule exhibited detectable TEV levels (Figure 5b). Furthermore, minimal TEV protease staining was observed in tissues surrounding the tumor, such as in the liver, intestine, or other stromal tissues (Figure S5). Having verified that outputted TEV protease could be detected within tumor nodules, we next assessed whether this heterologous reporter enzyme was also secreted into circulation. In mice with tumors that stained positively for TEV, TEV protein was detected in transduced cells and entered circulation (Figure S6). These results show that i.p. lentiviral circuit administration was effective OVCAR8 transduction in vivo, driving expression and secretion of TEV protease by integrating input signals from two OVCAR8-SPCES via a synthetic Boolean AND gate.

Upon confirmation of the output protein expression both in tumor cells and in circulation, we next evaluated whether TEV protease activity could be detected in tumor-bearing, circuit-transduced mice. To this end, we tested the performance of our two nanotechnology-based detection assays in live animals. For in vivo tumor detection experiments, OVCAR8 tumor-bearing mice transduced with the complete circuit ([1,1]) or control cassette ([0,0]) were intravenously injected with TEV-responsive nanosensors 1 week postcircus transfusion. Urine was collected from mice 1.5 h post injection, and ELISA-based detection assay was run to detect the presence of liberated reporters. In parallel, blood was collected, and serum samples were reacted with the fluorogenic reporter for the ex vivo readout. Urine samples from tumor-bearing mice transduced with the complete circuit ([1,1]) exhibited significantly increased urinary reporter signal relative to tumor-bearing mice transduced with the output virus alone ([0,0]) (Figure 5c, \( P = 0.0078 \)). Similarly, using the fluorogenic reporter in the ex vivo detection assay, TEV protease activity was observed in blood samples from mice in the [1,1] group (Figure 5d), indicating significantly increased circulating TEV activity relative to the [0,0] control group (Figure 5e, \( P = 0.0027 \)). Taken together, these results provide a proof-of-concept demonstration that a synthetic gene circuit can be coupled with nanoparticle-based readout tools to sense ovarian cancer-associated signals in vivo in mice and to generate signal-amplified readouts traceable in blood and urine.

**DISCUSSION**

In this work, we designed an integrated sense-and-respond system, comprised of a genetically encoded sensing circuit and nanotechnology-based detection tools, and demonstrated its proof-of-principle application for tumor-specific expression of heterologous biomarkers. The AND-gate synthetic gene circuit was engineered to express the heterologous reporter TEV protease in response to transcriptional dysregulation associated with ovarian cancer cells. Intraperitoneal lentiviral circuit delivery effectively triggered TEV protease expression from within tumor cells in a disseminated ovarian cancer mouse model. We then deployed two detection assays for measuring the activity of circuit-produced TEV protease against engineered nanomolecular probes in vivo or ex vivo. Specifically, we engineered peptide-coated nanoparticles that,
following systemic administration, interacted with TEV protease in vivo to release reporter molecules detectable in the urine. TEV protease activity was also measured in the blood using fluorogenic molecular probes. As a proof-of-concept demonstration, we validated the behavior of this engineered detection system in vivo in an ovarian cancer mouse model, where we showed that circuit-transduced tumor cells produced TEV protease and that the activity of this heterologous biomarker could be measured using either of our detection platforms.

Integrating synthetic biology approaches with nanotechnology tools presents a new framework for developing engineered sensors that can monitor, record, and respond to the levels of biological and environmental cues. Each of these approaches presents complementary advantages. Synthetic biology provides a tunable toolkit that could significantly expand the repertoire of accessible disease biomarkers beyond traditional blood-based biomarkers.\(^{30}\) For example, synthetic gene circuits could be engineered to measure multiple transcriptional, protein, or metabolic biomarkers of disease,\(^7\) and to integrate these measurements via logic gating\(^{36–38}\) or more complex computations.\(^9,40\) Such circuits could be designed to in turn generate cell-state specific expression of heterologous biomarkers, such as reporter enzymes or effector proteins. By exploiting the unique properties that emerge at the nanoscale, multifunctional nanoparticles could be engineered to target and penetrate disease sites, either to detect the presence or activity of circuit-generated biomarkers or to sense orthogonal signals to multiplex the inputs to the synthetic circuit. Nanosystems have been extensively used to produce both in vivo imaging signals\(^{22,41}\) and ex vivo readouts in accessible biofluids or tissues.\(^{14}\) Tuning similar nanoparticle sensors toward circuit-produced outputs could provide multiple levels of disease specificity and sensitivity that could ultimately yield highly accurate engineered detection systems.

Here, we demonstrated successful circuit-driven transcriptional sensing as well as output generation in vivo, showing both that TEV protease could be expressed in tumor cells and that its activity could be measured by two downstream detection platforms. However, the complete AND-gate synthetic circuit was only validated against a control circuit carrying just the output TEV expression cassette, a key component\(^7\). For example, synthetic gene circuits could be validated using mouse ovarian cancer SPECS as inputs, to integrate synthetic genetic circuits with nanotechnology tools for tumor detection. Our platform is highly tunable and can generate amplified signals for easy readouts in either the blood or urine following in vivo deployment. Moreover, with simple design modifications, such as replacing the SPECS inputs from ovarian-cancer SPECS to SPECS for other cancers, our system could be adapted to detect additional tumor types. Finally, expression of outputs that activate cytotoxic loads linked to nanoparticles could be potentially implemented as a safe and effective cancer therapy. We envision that the integration of synthetic biology and nanotechnology proposed by this work may enable new classes of engineered sense-and-respond systems.

## MATERIALS AND METHODS

All experimental procedures were performed in accordance with the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Animal Care and Use Committee (IACUC) of the Massachusetts Institute of Technology.

### Antibodies and Recombinant Proteins

The following primary antibodies were used in this study: mouse anti-V5 antibody (ThermoFisher, R960-25), rabbit anti-V5 antibody (Sigma, V8137), Alexa Fluor 555 donkey-antimouse (ThermoFisher, A-21200), goat antimouse serum albumin (Abcam, ab19194), mouse antirabbit HRP (Santa Cruz, SC-2357), mouse antigoat HRP (Santa Cruz, SC-2354), goat antihuman serum albumin (Abcam, ab19194), mouse antirabbit HRP (Santa Cruz, SC-2357), mouse antigoat HRP (Santa Cruz, SC-2354), goat antihuman HRP (Santa Cruz, SC-2005), rabbit anti-FITC (GeneTec, GTX26644), streptavidin-conjugated HRP (Abcam, ab7403).

The following recombinant proteins were used in this study: TEV protease (Anaspec, AS-72227), MMP2 (RD Systems, 902-MP-010), MMP9 (RD Systems, 911-MP-010), MMP13 (RD Systems, S11-MM-010), uPA (RD Systems, 1310-SE-010), thrombin (Haematologic Technologies, HCT-0020).

### Plasmids

All synthetic circuit plasmid maps are provided in Figures S7–S10. The sequences for relevant biological parts are provided in Table S1. The synthetic circuit plasmids were constructed by conventional restriction enzyme cloning and Gibson assembly (Gibson Assembly Master Mix, NEB, E2611) with the backbone of lentiviral vector FUGW (Addgene, 14883). The three synthetic promoters S(E2F1)P, S(cMyc)P, and GAL4BS were designed as previously described.\(^{20}\) Specifically, for the inputs, the synthetic promoter S(E2F1)P...
was placed upstream of the S′ end of Cys4, while S(cMyc)P was placed upstream of the S′ end of the RNA-based self-inhibitory gene cassette. The RNA-based self-inhibitory gene component consisted of two exons of GAD, a miRNA synthetic promoter GAL4BS was placed upstream of transmembrane GFP (hTFR-GFP, Addgene plasmid 45060) or TEV protease (pcDNA3.1-V5-hTEV, Addgene plasmid 65800). The TEV protease gene was modified by inserting an IgK leader sequence at the N-terminus for secretion (cloned from vector pSecTag2 B, ThermoFisher, V90020), a V5 tag sequence at the S′ end, and a Myc tag sequence at the 3′ end. To make active secreted TEV protease, we created a N23Q, C130S, T173G triple TEV mutant by site-directed mutagenesis (QuickChange Site-Directed Mutagenesis Kit, Agilent, 200519), using the following primers:

- N23Q, 5′-ctgacactccacgaggtctgccag-3′
- C130S, 5′-tgaggctttcgctagtacattttggagga-3′
- T173G, 5′-ggtgtcgataacgcagctacatct-3′

**Cell Culture and Viral Production.** All cell lines in this study were maintained in Dulbecco’s modified Eagle medium (DMEM; Invitrogen, 11965092) containing 10% fetal bovine serum (ThermoFisher, 10437028) and antibiotics (25 U/mL (DMEM; Invitrogen, 11965092) containing 10% fetal bovine serum and 1% penicillin and 25 μg/mL streptomycin), and cultured in humidified, 5% CO2, environment at 37 °C.

Lentiviruses were produced by cotransfecting HEK293T cells with 6 μg lentiviral expression vector, 4.5 μg delta 8.9 vector, and 3 μg VSVG packaging vector in a 10 cm dish using a calcium phosphate transfection kit (CalPhos Mammalian Transfection kit, Clontech, 631312). Two days post-transfection, the supernatant was harvested, centrifuged at 3000 rpm for 10 min, and snap frozen in liquid nitrogen.

**In Vitro Viral Transduction and Synthetic Circuit Characterization.** To validate the behavior of the synthetic circuit, OVCAR8 ovarian cancer cells were seeded in 96 or 6-well plates, and cells were transduced with the output virus ([0,1]), output virus with one of two input viruses ([1,0], [0,1]), or both input viruses and the output virus ([1,1]) in the presence of 4 μg/mL Polybrene (Santa Cruz, SC-134220). Eight hours post transduction, cells were replenished with fresh DMEM medium containing 10% fetal bovine serum and 1% penicillin/streptomycin, and cultured for an additional 2 days. Cells in 96-well plates were directly fixed with 4% paraformaldehyde (PFA, EMS, 50–980–495) for immunofluorescence imaging, while cells in 6-well plates were trypsinized and then fixed with 4% PFA for flow cytometry analysis. For the GFP output cassette, fixed cells were directly processed for immunofluorescence imaging or flow cytometry. However, for the V5-tagged SecTEV QSG output cassette, cells were first permeabilized with 0.1% Triton X-100 (Sigma-Aldrich) in PBS for 5 min, blocked in 3% bovine serum albumin in PBS (PBSA) for 30 min, and incubated with mouse anti-V5 antibody (1:500, ThermoFisher, R960–25) diluted in 3% PBSA for 1.5 h. Cells were washed three times in PBS every 5 min, and then incubated with Alexa Fluor 555 donkey antimouse secondary antibody (1:1000 dilution, ThermoFisher, A-31570) or Alexa Fluor 488 chicken-antimouse secondary antibody (1:1000 dilution, ThermoFisher, A-21200) for 1 h. Following staining, cells were imaged on a Nikon microscope for immunofluorescence imaging, or analyzed on a LSRII Fortessa cytometer (BD Biosciences) for flow cytometry. Flow cytometry data analysis was performed in Flowjo (TreeStar Inc., Ashland, OR).

**TEV Activity Sensors and In Vitro Measurement Assays.** For the fluorogenic TEV reporter, the TEV specific peptide substrate Glu-Asn-Leu-Tyr-Phe-Gln-Gly was labeled with S-FAM and QXL520 quencher (AnaSpec, AS-72227). The fluorogenic TEV substrate was incubated with recombinant enzyme according to manufacturers’ protocols. Proteolytic cleavage of this quenched substrate was quantified by increases in fluorescence signal intensity (Ex/Em = 490/520 nm) over time as measured by fluorimeter (Tecan Infinite M200 Pro).

For synthesis of TEV-responsive nanoparticles, 40 kDa, eight-arm multivalent (poly)ethylene glycol (PEG) molecules with maleimide reactive handles (JenKem Technology) were dissolved in PBS and filtered through a 0.2 μm membrane filter (Acrodisc). Filtered PEG nanoparticles were reacted with at least 20-fold excess of cysteine-terminated TEV-specific peptide for at least 1 h at room temperature. The TEV-specific peptide (Sequence: Biotin-eGvndneeGdGGENLYFQGGGC) consists of a glutamate fibrinopeptide B (GluFib) urinary reporter, encoded with an N-terminal biotin and S-FAM on the opposite terminus,28 and a TEV-specific cleavage site, and was synthesized by the Koch Institute Polymers and Peptides Core at MIT. Following the maleimide–cysteine coupling, unconjugated peptides were removed from the nanoparticle complexes through centrifugal filtration with 30kD spin column filters (Millipore, UFC503096). The size of synthesized PEG nanoparticles was measured by transmission electron microscopy (TEM) in with 2% uranyl acetate on an FEI Tecnai Spirit Transmission Electron Microscope at the Koch Institute facility.

To measure the activity of cell-secreted TEV in vitro, OVCAR8 cells were seeded and transduced as previously described. Cells were cultured in OptiMEM (Gibco, 11058021) overnight, and supernatant was collected. 100 μL of collected supernatant was incubated with fluorogenic TEV substrate (1 μM), and proteolytic cleavage was monitored by increase in fluorescence intensity over time as measured by fluorimeter.

**ELISA for Ligand-Encoded Reporters.** 96-well plates (Nunc) were coated with capture antibodies (Rabbit anti-FITC, GeneTex, 1:1000 dilution) for 1 h in coating buffer (0.05 M carbonate bicarbonate, pH 9.6), and blocked with 3% (w/v) bovine serum albumin (BSA) in PBS for 30 min. After incubating with 100 μL sample solution for 1 h at room temperature or overnight at 4 °C, the wells were washed with PBST (PBS with 0.2% Tween) five times, and incubated with detection antibody (Streptavidin-HRP, Abcam, 1:3000 dilution) in PBST for 1 h before adding the chromogenic substrate TMB (ThermoFisher, 34028). Oxidation of TMB for 5–10 min allowed quantification of reporter concentrations. For urine experiments, samples were diluted at 1:300 in 3% BSA in PBS.

**Ovarian Cancer Mouse Model and In Vivo TEV Activity Measurements.** All animal studies were approved by the Massachusetts Institute of Technology (MIT) committee on animal care (MIT protocol 0420–023–23) and conducted in compliance with ARRIVE guidelines. To generate the disseminated ovarian cancer mouse model, three to four-week-old female NCr nude mice (Taconic) were injected with 5 million OVCAR8 cells intraperitoneally (i.p.).
OVCAR8 cells were modified to stably express luciferase, allowing us to monitor tumor growth through intravital IVIS imaging (PerkinElmer, Koch Institute Animal Imaging and Preclinical Testing Core, MIT). For imaging, 100 μL of 15 mg/mL d-luciferin potassium salt (GoldBio, LUCK-100) dissolved in PBS was injected into each mouse subcutaneously, and luciferase activity was measured after 5 min.

Four weeks after seeding of OVCAR8 cells, mice were intraperitoneally injected with different combinations of lentivirus mix (corresponding to the [0,0] or [1,1] state) diluted in 100 μL sterile PBS. After 1 week, 30 μL blood was collected from each mouse through tail vein bleeding and mixed with 30 μL PBS containing 1 μM EDTA (Invitrogen, 15575020). The collected blood was centrifuged at 13 000 rpm for 2 min, and the supernatant plasma was collected. For the fluorogenic detection assay, plasma samples were incubated with fluorogenic TEV peptide substrate (1 μM), and substrate cleavage was monitored by fluorescence increase over time as measured by a Tecan plate reader.

To detect cancer through urinary readouts, tumor-bearing mice were transduced with lentivirus combinations as described above. One week post-transduction, mice were injected intravenously with TEV-responsive nanosensors (1 μM in 200 μL sterile PBS) and subcutaneously with 500 μL of sterile PBS to facilitate urinary production. One hour post nanosensor injection, mice were placed in custom housing with a 96-well plate base for urine collection. After 30 min, bladders of the mice were voided to collect urine. Collected urine samples were diluted 300-fold for ELISA measurement.

**Histology.** Immunohistochemistry was performed to determine whether transduced tumor cells expressed TEV protease. OVCAR8 tumor-bearing mice were transduced with lentivirus combinations as previously described, and tumor nodules were extracted and fixed in 4% PFA in PBS overnight. Fixed tissue was paraffin embedded and sectioned for immunostaining (Histology Core, Koch Institute, MIT). Sections were stained with mouse anti-VS antibody (1:1000 dilution, ThermoFisher, R960–25) and appropriate secondary antibody, and images were acquired by a Nikon microscope.

**Statistics and Data Analysis.** All statistical analyses were performed in GraphPad Prism 9.0. Statistical significance and individual tests are described in figure legends.

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**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssynbio.1c00133.

Additional results and figures including the following: inputs and states of the AND gate; output and specificity characterization of the synthetic circuit; characterization of the PEG-based nanosensor; in vitro characterization of circuit ON-OFF ratio; circuit-produced TEV abundance in healthy tissues; detection of circuit-produced TEV protease in the blood; plasmid maps for circuit components; sequences of biological parts (PDF)

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and S.N.B. wrote the manuscript. All authors reviewed and approved the final manuscript.

Notes
The authors declare the following competing financial interest(s): L.N. holds equity in and is the scientific founder of Circuit-Bio and receives sponsored research funding from FutuRx. S.N.B. holds equity in Glympse Bio, Satellite Bio, Cend Therapeutics, and Catalio Capital; is a director at Vertex; consults for Moderna; and receives sponsored research funding from Johnson & Johnson.

The data that support the findings of this study are available upon request to the corresponding author.

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