Cell-free biosensors for rapid detection of water contaminants

Jaeyoung K. Jung, Khalid K. Alam, Matthew S. Veroslof, Daiana A. Capdevila, Morgane Desmau, Phillip R. Clauer, Jeong Wook Lee, Peter Q. Nguyen, Pablo A. Pastén, Sandrine J. Matiasek, Jean-François Gaillard, David P. Giedroc, James J. Collins, and Julius B. Lucks

Lack of access to safe drinking water is a global problem, and methods to reliably and easily detect contaminants could be transformative. We report the development of a cell-free in vitro transcription system that uses RNA Output Sensors Activated by Ligand Induction (ROSALIND) to detect contaminants in water. A combination of highly processive RNA polymerases, allosteric protein transcription factors and synthetic DNA transcription templates regulates the synthesis of a fluorescence-activating RNA aptamer. The presence of a target contaminant induces the transcription of the aptamer, and a fluorescent signal is produced. We apply ROSALIND to detect a range of water contaminants, including antibiotics, small molecules and metals. We also show that adding RNA circuitry can invert responses, reduce crosstalk and improve sensitivity without protein engineering. The ROSALIND system can be freeze-dried for easy storage and distribution, and we apply it in the field to test municipal water supplies, demonstrating its potential use for monitoring water quality.

Synthetic biology is creating new capabilities for engineered biological systems to address pressing global challenges through repurposing, evolving and rewiring the functions of biological components for a range of applications. Among these functions, the ability of cells to use molecular biosensors and genetic networks to sense and respond to changing conditions is being harnessed to create new diagnostic technologies. In particular, cell-free nucleic-acid diagnostics are being developed to detect pathogens and human health biomarkers in complex matrices. These systems use RNA genetic switch design or CRISPR-mediated interactions to detect specific target DNA or RNA sequences, and trigger gene expression, fluorophore release or other biochemical reactions to produce a detectable signal. Changing the aTF allows cellular systems to be engineered to sense a range of contaminants including metals, aromatic compounds and antibiotics. Advantages of this approach include signal amplification through gene expression and selectivity afforded by processes that affect compound bioavailability. However, there are challenges associated with WCBs, including maintenance of viable cells, difficulties with sensing toxic contaminants, membrane transport limitations and the need to address biocontainment and regulatory concerns for deploying living organisms in the field.

In the present study, we sought to preserve the modularity and signal amplification advantages of WCBs, while addressing some of their limitations. We developed cell-free diagnostics using ROSALIND (RNA Output Sensors Activated by Ligand Induction)
to detect contaminants in water (Fig. 1). ROSALIND comprises a highly processive phage RNA polymerase (RNAP), aTFs and engineered DNA transcription templates, which together produce visible outputs on exposure to specific ligands. The use of RNA-level outputs allows signals to be observed within minutes and removes the need for complex and resource-intensive protein translation to detect responses. In addition, the use of in vitro transcription (IVT) reactions removes the need for cells and overcomes biocontainment concerns. ROSALIND represents a capability for simple and rapid field monitoring of chemical species to address growing global health crises in water quality.

**Results**

**Fluorescent aptamers are rapid visible transcription reporters.** We reasoned that fluorescent RNA-level outputs generated through IVT would allow biosensing reactions to be simplified, thereby resulting in faster signal generation than protein-level outputs that must be translated in more complex reaction environments. To demonstrate the viability of fluorescent RNA outputs, we chose the three-way junction dimeric Broccoli (3WJdB) aptamer and the fast T7 phage RNAP capable of transcribing 300 nucleotides per second in vitro. DNA transcription templates encoding 3WJdB behind a T7 RNAP promoter were applied to a commercially available transcription kit supplemented with the Broccoli-binding dye DFHBI-1T (Fig. 2a). Fluorescence activation was monitored in real time and standardized to known amounts of fluorescein (fluorescein isothiocyanate (FITC)) (Supplementary Fig. 1). The IVT reactions generated a fluorescence signal comparable to micromolar equivalents of fluorescein (MEF) across a range of input DNA-template concentrations within minutes (Fig. 2b and Supplementary Video 1). Furthermore, the fluorescence activation of 3WJdB was easily visualized at 0.1 MEF (µM FITC) with low-cost light-emitting diodes and a stage-lighting filter (Fig. 2c and Supplementary Data 1). In addition, we found that inclusion of a T7 terminator in the transcription templates resulted in improved fluorescence activation (Supplementary Fig. 2).

We next compared the fluorescence intensity and kinetics of the RNA output with that of superfolder green fluorescent protein (sfGFP) produced using a commercial kit based on the PURE system. 3WJdB-expressing IVT reactions and sfGFP-expressing PURE reactions were run using equimolar amounts of DNA templates. In this setup, the RNA output immediately began to increase in fluorescence intensity, whereas sfGFP required at least 9 min before a detectable signal was observed (Fig. 2d and Supplementary Fig. 3). In addition, over the 4-h time course, 3WJdB transcription produced substantially more fluorescence than sfGFP (Fig. 2d and Supplementary Fig. 3), and the signal generated with a homemade transcription reaction was comparable to that of the commercial kit (Extended Data Fig. 1).

Together, these results show that IVT of fluorescence-activating RNA aptamers can be repressed by aTFs and induced by targets of interest, thereby establishing a proof of concept for the ROSALIND system.

**aTFs enable ligand responsiveness.** We next sought to create a ligand-responsive biosensor by using an aTF to regulate T7 RNAP IVT. We reasoned that the binding of an aTF to its cognate operator sequence would prevent T7 RNAP from promoter escape in the absence of ligand, and ligand-mediated de-repression of an aTF would result in a fluorescence signal (Fig. 1).

We first investigated how aTF–operator interactions can block T7 RNAP transcription in the absence of ligand. We chose TetR, a well-characterized model aTF that binds the 19-base pair (bp) palindromic tet operator sequence (tetO) in the absence of its cognate ligand tetracycline. We designed a series of transcription templates to explore the effect of spacing between the minimal 17-bp T7 promoter and the tetO sequence (Fig. 2e). We used the native sequence that follows the canonical T7 RNAP promoter as a spacer in 2-bp increments from 0 to 14 bp, immediately followed by tetO, 3WJdB and T7 terminator sequences. IVT reactions were set up as before using homemade reactions, with the addition of purified recombinant TetR protein in 100-fold excess of the DNA template. We observed that the absence of any spacing resulted in no fluorescence activation in either the presence (regulated) or the absence (unregulated) of TetR (Fig. 2f). However, robust fluorescence activation was observed without TetR when using a 2–6-bp spacer, which was reduced to baseline levels when regulated by TetR. Spacers...
that were ≥8 bp resulted in fluorescence activation in the regulated reactions. We therefore chose the shorter functional spacers for subsequent promoter–operator designs.

We next sought to determine whether TetR can be de-repressed with a cognate ligand to permit IVT and therefore act as a biosensor (Fig. 2g). We first performed IVT reactions over a range of TetR concentrations using the transcription template with a 4-bp spacer (Extended Data Fig. 2). We found that a 25-fold excess of TetR dimer to transcription template was the minimal amount of repressor needed to ensure tight regulation of transcription, whereas a 50-fold excess of TetR dimer performed similarly to the 100-fold excess (Extended Data Fig. 2). We then set up IVT reactions using these TetR and DNA template concentrations and included a range of anhydrotetracycline (aTc) concentrations. We observed strong repression over 4 h through high nanomolar amounts of aTc, with half-maximal induction between 1 and 2.5 μM aTc (Fig. 2h and Extended Data Fig. 3). Importantly, when induced with 2.5 μM aTc, the kinetics of induction follow those of the unregulated transcription reaction (Extended Data Fig. 2).

Together, these results show that aTF-regulated IVT of fluorescence-activating RNA aptamers can serve as biosensors, establishing a proof-of-concept basis for the ROSALIND system.

Modularity allows sensing of organic molecules and metals. Next, we sought to expand the range of molecules that can be detected with ROSALIND by leveraging the natural diversity of aTFs. Inspired by earlier work with WCBs, we reasoned that ROSALIND could sense a wide range of important targets by changing the operator sequence within the transcription templates and including the corresponding aTF in the reaction (Extended Data Fig. 2 and Supplementary Data 2).

We began with antibiotics, the overuse of which is resulting in the contamination of freshwater supplies and driving antimicrobial resistance. We first tested our TetR-based sensor to detect tetracycline (Fig. 3a and Extended Data Fig. 4a) and the semisynthetic antibiotic doxycycline (Fig. 3b and Extended Data Fig. 4b), both of which are on the World Health Organization’s (WHO’s) list of essential medicines. Reactions with TetR-regulated transcription templates and purified TetR were run using either laboratory-grade water (−) or similar water with added antibiotic (+). In both cases, we observed generation of fluorescence with the antibiotic present and no fluorescence in the water control.

We then leveraged OtrR, a member of the multiple antibiotic-resistance repressor (MarR) family of aTFs, to serve as a biosensor for oxytetracycline (Fig. 3c and Extended Data Fig. 4c). To expand the range of tetracycline antibiotic sensing further, we identified a putative MarR-family aTF analogous to OtrR, CtcS, and the divergent operator regulated by CtcS, and designed a minimal operator sequence to enable its use in ROSALIND for sensing chlorotetracycline (Fig. 3d and Extended Data Fig. 4d).
We next sought to expand sensing to the macroclide class of antibiotics that are also widely used in human health. We chose a TetR-family aTF, MphR\textsuperscript{23}, to detect erythromycin, a drinking water contaminant of emerging concern\textsuperscript{14}, azithromycin, clarithromycin and roxithromycin (Fig. 3e–h and Extended Data Fig. 4e–h). We note that this observed broad range of MphR specificity is expected based on its known promiscuity, and is probably a common feature of xenobiotic-sensing aTFs\textsuperscript{23}.

After demonstrating the ability of ROSALIND to detect complex antibiotic molecules, we sought to leverage the system for the detection of other organic molecules, focusing on those with public health relevance. In particular, there has been a growing concern that unregulated contaminants, such as those generated from the large use of pharmaceuticals and personal care products, interfere with or are unaffected by water treatment, and are causing widespread health and environmental concerns\textsuperscript{26}. We started with the perfume additive 3-hydroxybenzoic acid as a model pharmaceutical, and personal care product that can de-repress the MarR-family aTF, MobR\textsuperscript{27}. When ROSALIND reactions were configured with MobR and its corresponding DNA template and aTF concentrations used in each reaction can be found in Supplementary Table 2. All data shown are for \( n = 3 \) independent biological replicates as lines with raw fluorescence values standardized to MEF (\( \mu M \) FITC). Shading indicates the average value of three independent biological replicates ± s.d. DNA template and aTF concentrations used in each reaction can be found in Supplementary Table 2.

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**Fig. 3** ROSALIND is modular and can be used to sense complex molecules, small molecules and metal ions. a, b, TetR can be used to sense tetracycline (a), doxycycline (b) and other tetracycline-family antibiotics. c, Oxytetracycline sensing with OtrR. d, Chlorotetracycline sensing with CtcS. e–h, MphR senses the macroclides erythromycin (e), azithromycin (f), clarithromycin (g) and roxithromycin (h). i, 3-Hydroxybenzoic acid sensing with MobR. j, Benzalkonium chloride sensing with QacR. k, Naringenin sensing with TtgR. l, Uric acid sensing with HucR. m, Zn\textsuperscript{2+} sensing from ZnSO\textsubscript{4}, with SmtB. n, Cu\textsuperscript{2+}/Cu\textsuperscript{+} sensing from CuSO\textsubscript{4}, with CsR from Bacillus subtilis. o–p, Pb\textsuperscript{2+} sensing from PbCl\textsubscript{2} (o) and Cd\textsuperscript{2+} sensing from CdCl\textsubscript{2} with CadC (p). Each reaction contains the indicated amount of ligand (±) dissolved in laboratory-grade water, ethanol, Tris-based buffer, dimethylsulfoxide or diluted sodium hydroxide, or a laboratory-grade water control (–) (Supplementary Table 2). All data shown are for \( n = 3 \) independent biological replicates as lines with raw fluorescence values standardized to MEF (\( \mu M \) FITC). Shading indicates the average value of three independent biological replicates ± s.d. DNA template and aTF concentrations used in each reaction can be found in Supplementary Table 2.
disinfectant that can promote antimicrobial resistance\(^4\), by leveraging the \textit{Staphylococcus aureus} TetR-family aTF, QacR\(^29\) (Fig. 3j and Extended Data Fig. 4j).

Some organic molecules such as naringenin\(^30\) and uric acid can serve as important markers for human health and nutrition. Using the \textit{Pseudomonas putida} TetR-family aTF, TigR\(^1\), and an optimized \texttt{tgO} variant sequence\(^2\), we designed ROSALIND reactions to sense naringenin in the tens of micromolar range (Fig. 3k and Extended Data Fig. 4k). In addition, we sought to detect uric acid, an early indicator of gout when present at \(\geq 6\) mg dl\(^{-1}\) within bodily fluids\(^31\). By incorporating HucR, a MarR-family aTF from \textit{Deinococcus radiodurans}\(^4\), and the corresponding DNA template, we were able to detect uric acid within water samples at approximately 50 ppm (\(-0.84 \text{ mg dl}^{-1}\)) or higher (Fig. 3l and Extended Data Fig. 4l).

Among water contaminants, metals have become a major issue due to aging infrastructure, mining operations and industrial waste disposal, and from natural abundance\(^32\)-\(^37\). To enable metal detection using ROSALIND, we first identified the canonical zinc-sensing aTF, SmtB, from the ArsR/SmtB family of regulators\(^8\). Using SmtB and the corresponding DNA template, we were able to sense Zn(II) at \(\geq 2.5\) µM (2163 ppm), which is below the US Environmental Protection Agency’s (EPA’s) secondary drinking guideline of 5 ppm. By incorporating HucR, a MarR-family aTF from \textit{Deinococcus radiodurans}\(^4\), and the corresponding DNA template, we were able to detect uric acid within water samples at approximately 50 ppm (\(-0.84 \text{ mg dl}^{-1}\)) or higher (Fig. 3l and Extended Data Fig. 4l).

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Although many metals are toxic at high concentrations, of particular concern are copper and lead\(^38\)-\(^40\) due to toxic effects such as psychiatric disorders, impaired neural development and organ damage. As such, their levels are regulated in drinking water by the US EPA’s Lead and Copper Rule (LCR). Among the many copper-sensing aTFs is CsoR, a Cu(I)-sensing aTF first identified in \textit{Mycobacterium tuberculosis} that forms a homotetramer to bind four copper ions\(^8\).

When configured into ROSALIND with the appropriate DNA template, our sensor was able to detect copper at \(\geq 5\) µM (\(\geq 318\) ppb) (Fig. 3n and Extended Data Fig. 4n). It is of interest that, although the cognate ligand for CsoR is Cu(I), we observed that the reaction senses Cu(II) from added CuSO\(_4\), possibly due to the presence of 10 mM dithiothreitol (DTT) in IVT reactions that can reduce Cu(II) to Cu(I). Another ArsR/SmtB family aTF, CadC, is found in \textit{Pseudomonas aeruginosa} and is known to bind a number of metals, including lead and cadmium\(^6\). When implemented in ROSALIND with appropriately configured DNA templates, we could robustly detect both lead and cadmium at \(\geq 1.25\) µM (\(\geq 259\) ppb for lead and \(\geq 141\) ppb for cadmium) (Fig. 3o,p and Extended Data Fig. 4p,o).

In total, we employed 11 different aTFs to sense 16 different ligands that are structurally and functionally diverse. An investigation into their limits of detection (Extended Data Fig. 4 and Supplementary Data 3) showed that the majority detect their respective ligands in the micromolar range. In addition, characterization of the inhibitory effects of the ligands and the solvents used to prepare them (Supplementary Fig. 4 and Supplementary Table 2) suggests that ROSALIND can tolerate much higher concentrations of the ligands than their respective WHO/EPA-regulated limits (Supplementary Table 1).

RNA circuitry can invert logic and fix crosstalk. Specificity is a key metric for the performance of biosensors. In some cases, a single aTF is well known to sense multiple ligands, as is the case with TetR sensing tetracyclines, MphR sensing macrolides and CadC sensing both lead and cadmium\(^18\)-\(^24\). In these cases, ROSALIND is best viewed as reporting on the presence of a specified family of stereochemically related ligands. To investigate whether ROSALIND reactions are specific for the indicated compounds outside of these known cases, we tested ROSALIND reactions configured with TetR, MphR, MobR, QacR, TtgR, HucR, SmtB, CsoR or CadC, along with their corresponding DNA templates, with a panel of cognate ligands. Tetracycline, erythromycin and lead were chosen as representative members of each class of compounds for TetR, MphR and CadC, respectively. In general, we found strong signals from cognate ligands with minimal crosstalk, indicating that ROSALIND reactions are generally specific for their intended ligands when DNA template and aTF concentrations are optimized (Fig. 4a and Extended Data Fig. 5).

We did, however, find significant crosstalk from the CsoR-containing copper sensor when induced with zinc (Fig. 4a and Extended Data Fig. 5). Notably there was no corresponding crosstalk across SmtB, the zinc sensor and copper. This created the opportunity to construct a sensor by combining the activity profiles of CsoR and SmtB in a single reaction to fix the crosstalk of CsoR. Specifically, this sensor was configured based on a Cu NIMPLY Zn (Cu AND NOT Zn) circuit that would respond only to copper and not zinc. Implementation of this circuit requires the presence of zinc to inhibit the 3WjdB signal produced by CsoR.

To achieve this zinc-mediated inhibition, we leveraged the recent development of RNA kleptamers—short antisense RNAs designed to inhibit Broccoli by disrupting the aptamer folding that enables fluorescence activation\(^1\). We found that fluorescence activation of 3WjdB could be inhibited through unregulated IVT of the KB2 kleptamer with T7 RNAP (Extended Data Fig. 6a), and that KB2 transcription could be regulated by TetR, by including the \texttt{tetO} sequence in the KB2 template (Extended Data Fig. 6b). Addition of its cognate ligand, aTc, effectively de-repressed TetR from the KB2 template, thereby allowing KB2 transcription and inverting the output of the TetR sensor (Extended Data Fig. 6c). We also demonstrated the generality of the kleptamer inversion strategy using the aTc repressor AdcR from \textit{Streptococcus pneumoniae}\(^1\)-\(^2\), which binds its operator in the presence of Zn(II), and inverting it into a zinc-activated sensor (Extended Data Fig. 6d).

We used the kleptamer strategy within the Cu NIMPLY Zn circuit. In this circuit design, the transcription of 3WjdB is regulated by CsoR, whereas the transcription of KB2 is regulated by SmtB (Fig. 4b). In the presence of copper only, CsoR is de-repressed from the 3WjdB transcription template, generating 3WjdB. In the presence of zinc only, both CsoR and SmtB are de-repressed from their respective transcription templates, generating 3WjdB and KB2, which is then able to prevent 3WjdB from binding its dye, shutting off the fluorescence signal. When implemented in ROSALIND, the Cu NIMPLY Zn circuit performed as expected, correcting the crosstalk of CsoR with zinc (Fig. 4c). Furthermore, the circuit was able to fix the crosstalk across a wide range of zinc concentrations, while keeping the dose-response curves against copper unaffected (Supplementary Fig. 5). Overall, these results show that RNA genetic circuitry can be designed to address issues with aTF specificity without having to use protein engineering strategies.

Sensitivity can be tuned with transcriptional circuitry. A shortcoming of our approach is that it is limited by the natural sensitivity of the aTF employed in the ROSALIND reaction. For example, our TetR sensor was activated by doxycycline at 1.25 µM (556 ppb), whereas the typical concentrations of antibiotic pollution found in environmental water samples are in the range of tens of parts per billion\(^4\). We therefore sought to tune the sensitivity of ROSALIND reactions using a generalizable strategy that did not require engineering of the aTF.

As proof-of-concept demonstrations, we first sought to tune the copper- and lead-sensing reactions containing CsoR and CadC, respectively. The dose–response curves of the copper and lead sensors indicate that the limit of detection for copper (0.3 ppm of Cu(II)) is below the threshold set by the US EPA’s LCR (1.3 ppm), whereas that of lead (259 ppb) is an order of magnitude above the desired threshold (15 ppb) (Extended Data Fig. 7 and Supplementary Table 1). To desensitize the copper sensor, we increased the CsoR concentration to allow any excess to act as a copper chelator and shifted the dose–response curve to activate near the desired LCR threshold.
of 1.3 ppm of copper from added CuSO₄ (Extended Data Fig. 7a). To sensitize the lead sensor, we thus employed the converse of our CsoR strategy. By lowering the CadC dimer concentration to 0.5 µM, we were able to shift the dose–response curve to be more sensitive to detect tens of parts per billion of Pb(II) (Extended Data Fig. 7b). However, this strategy resulted in a significant increase in signal in the absence of lead.

To address this limitation, we sought to implement a different method to improve sensor sensitivity, based on the previous development of a TetR-binding RNA aptamer (anti-TetR) which de-represses TetR by mimicking its operator binding site tetO (ref. 43). Taking advantage of this feature of anti-TetR, we implemented an RNA-feedback loop designed to enhance the expression of 3WJdB at low concentrations of doxycycline by regulating anti-TetR transcription with TetR (Fig. 5a). As configured, when doxycycline is present, the circuit is designed to de-repress TetR and initiate the expression of 3WJdB as well as anti-TetR. Once transcribed, anti-TetR can then further de-repress any TetR that is blocking the transcription of 3WJdB. When implemented in ROSALIND, the feedback circuit improved the sensitivity of the doxycycline sensor by an order of magnitude to the ~0.2 µM (89 ppb) range (Fig. 5b and Supplementary Data 3). These results demonstrate that TetR-based sensors can be sensitized to the ranges of antibiotic concentrations typically found in the environment, although preconcentration strategies would further enhance this. Moreover, this demonstrates that ROSALIND is compatible with RNA-feedback circuits that can improve the sensitivity of aTFs without any protein engineering.

**Application to municipal water samples.** Next, we examined whether ROSALIND can be formulated for distribution in the field. Many of the advances in synthetic biology diagnostic technologies have been enabled through freeze-drying cell-free systems for convenient storage and distribution before rehydration at the point of use44. To test whether the ROSALIND system can be freeze-dried, we explored a range of known lyoprotectants and determined that nonreducing disaccharides, such as sucrose and trehalose, in combination with a sugar alcohol, mannitol, provided substantial lyoprotection. In addition, we developed a freeze-drying and packaging procedure (Supplementary Video 2) that enabled freeze-dried ROSALIND reactions to be stored at ambient temperature for at least 2.5 months and remain functional, although with some impact on measured fluorescence (Extended Data Fig. 8).

In addition to freeze-drying, field deployment also requires a convenient approach to detecting the fluorescence signals generated by ROSALIND. To this end, we developed a three-dimensional (3D)-printed, handheld device that could visualize fluorescent output using the naked eye (Extended Data Fig. 9 and Supplementary Data 4). The use of off-the-shelf components assembled onto a
Fig. 5 | Sensitivity of ROSALIND reactions can be tuned using an RNA-feedback circuit. a, A TetR-binding RNA, anti-TetR, can be used to improve the limit of detection of the TetR sensor. In this feedback circuit, TetR regulates both the expression of 3WJdB and anti-TetR. In the presence of the cognate ligand, doxycycline, anti-TetR is transcribed and further de-represses TetR to enhance 3WJdB expression. b, When implemented in ROSALIND, the anti-TetR feedback circuit improves the sensitivity of TetR by an order of magnitude. Each reaction contains 25 nM 3WJdB template, 50 nM anti-TetR template and 1.25 μM TetR dimer. A transcription template encoding the reverse sequence of anti-TetR was used as a control to demonstrate that sensitization required the anti-TetR interaction. All data shown are for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (μM FITC). Error bars indicate the average values of three independent biological replicates ± s.d. The ligand concentrations at which the signal was distinguishable from the background were determined using a two-tailed, heteroscedastic Student’s t-test against the no-ligand condition, and their P value ranges are indicated with asterisks (**P < 0.001, ***P = 0.001-0.01, *P = 0.01-0.05). Exact P values along with degrees of freedom can be found in Supplementary Data 3. Data for no-ligand condition were excluded in b because the x axis is on the log scale, and are presented in Supplementary Data 3.

Discussion

In the present study, we report the development of a cell-free biosensor platform, ROSALIND, for detection of a wide range of chemical compounds and elements in water. ROSALIND consists of simple and defined aTF-regulated IVT reactions that produce RNA outputs as a function of ligand concentration. Inspired by previous WCB approaches, this modular architecture allowed us to develop 16 different sensors by changing the aTF–operator pair (Fig. 3). The use of fast phage polymerases generated visible signals to the naked eye in less than an hour. In addition, the reactions and handheld illuminators could be manufactured inexpensively (Supplementary Table 3 and Supplementary Data 4).

A key advantage of transcriptional RNA outputs was our ability to design RNA-level circuits to address well-known shortcomings of aTFs without performing any protein engineering. In particular, we were able to demonstrate that aTF–ligand crosstalk can be eliminated with RNA-level logic gates (Fig. 4), and aTF sensitivity can be improved through an RNA-mediated feedback circuit (Fig. 5). We anticipate that these RNA-level circuit strategies will be applicable to optimizing broad classes of aTFs.

The simple and defined nature of ROSALIND makes it amenable to extensions. For example, multiplexing can be added with additional fluorescent aptamers, and further engineering of the aTF or operator sequences can be used to enhance sensitivity, modify specificity and otherwise tune system performance. In addition, other uses besides water contaminant detection are possible, including the high-throughput screening of poorly characterized aTFs to uncover their cognate ligands and putative operator sequences, as we showed with our validation of the putative aTF, CtcS (Fig. 3d).

A key limitation of ROSALIND is that it is currently limited to aTFs that block RNAP elongation unless bound to a ligand. In addition, although the sensors are easy to operate from a user perspective, purifying aTFs that are functional in vitro can be nontrivial.
Finally, signal detection by eye can be difficult due to variations in lighting and reaction conditions. Although the inclusion of control reactions helps with this, a more sophisticated illuminator design that incorporates quantification would improve this aspect of the system.

A main feature of our study is demonstrating that freeze-dried ROSALIND reactions could be manufactured and shipped across the USA at ambient temperatures and applied to municipal water samples in the field (Fig. 6). We also found that ROSALIND reactions could be applied to other environmental water sources, although with some inhibition of the observed fluorescence levels (Extended Data Fig. 10). This can potentially be due to molecular components in the water samples that can inhibit transcriptional activity or quench fluorescence. Nevertheless, we did find that a practical workflow of testing serial dilutions could be used to detect copper, which we anticipate dilutes the interfering components in the reactions (Extended Data Fig. 10). In addition, different test formats may be used to include calibration curves that control for matrix effects.

ROSA LIND builds on significant earlier work in developing WCBs and cell extract-based biosensors. Compared with WCBs, the defined in vitro environment of ROSALIND enables rapid optimization of sensor performance (Extended Data Fig. 2) and implementation of RNA-level circuitry to address key limitations of aTF-based sensors. However, this flexibility comes at the cost of purifying functional aTFs and optimizing reaction conditions, and one advantage of WCBs is the simplicity of creating engineered strains. WCBs may take advantage of active transport to accumulate and detect only a bioavailable ligand, that is, the fraction that can cross the cell membrane, although more systematic comparisons are needed to precisely define these benefits. Finally, WCBs require additional equipment or procedures to maintain cells in physiologically active states, encapsulation to manage biocontainment and normalization of signals to account for different numbers of cells in each reaction—challenges that are significantly mitigated by cell-free biosensing approaches such as ROSALIND.

Taken together, WCBs and cell-free synthetic biology approaches such as ROSALIND help fill a gap in existing water-quality-monitoring technology, which currently consists mainly of centralized and expensive instrumentation that cannot always be deployed in the field (Supplementary Table 4). We anticipate ROSALIND as part of a synthetic biology-based solution to enable communities and individuals to rapidly and economically monitor their water quality to meet their needs, to inform when and where more accurate measurements are needed and to facilitate the gathering of critical information necessary for achieving the United Nations Global Sustainable Development Goal of safe and clean water.

Online content
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Methods

Strains and growth medium. Escherichia coli strain K12 (NEB Turbo Competent E. coli, New England Biolabs, catalog no. C2984) was used for routine cloning. E. coli strain Rosetta 2(DE3)pLysS (Novagen, catalog no. 71401) was used for recombinant protein expression. Lysogeny broth supplemented with the appropriate antibiotic(s) (100 µg ml⁻¹ carbenicillin, 100 µg ml⁻¹ kanamycin and/or 34 µg ml⁻¹ chloramphenicol) was used as the growth medium.

Plasmids and genetic assembly. DNA oligonucleotides for cloning and sequencing were synthesized by Integrated DNA Technologies. Genes encoding aTFs were synthesized either as gBlocks (Integrated DNA Technologies) or gene fragments (Twist Bioscience). Expression plasmids were cloned using Gibson Assembly (NEB Gibson Assembly Master Mix, New England Biolabs, catalog no. E2611) into a PET-28c plasmid backbone and were designed to overexpress recombinant proteins as either C terminus or N terminus, His-tagged fusions. Certain constructs additionally incorporated a recognition sequence for cleavage and removal of the His-tag using tobacco etch virus protease (Supplementary Table 5). Gibson-assembled constructs were transformed into NEB TURBO cells, and isolated colonies were purified for plasmid DNA (QIAprep Spin Miniprep Kit, Qiagen, catalog no. 27106). Plasmid sequences were verified with Sanger DNA sequencing (Quintara Biosciences) using the primers listed in Supplementary Data 5.

Plasmids encoding transcription templates were constructed using pUC19-T7-3WJdB-T (Addgene, catalog no. 87308) as a backbone. These plasmids included a T7 RNApolymerase promoter, an aTF promoter site, the 3WJdB coding sequence and a T7 terminator. In the present study, we define the T7 RNApolymerase promoter as a minimal promoter region ending with either 3WJdB or the T7 promoter region ending with either 3WJdB or the T7 terminator. In the primer set designed for the reaction mix, an overexpression construct contains the T7 promoter and a 3′-region ending with either 3WJdB or the T7 terminator, with the primers listed in Supplementary Data 5. Amplified templates were then purified (QIAquick PCR purification kit, Qiagen, catalog no. 28106) and verified for the presence of a single DNA band of expected size on a 2% Tris–acetate–EDTA (TAE)–agarose gel, and concentrations were determined using the Qubit dsDNA BR Assay Kit (Invitrogen, catalog no. Q32653).

The KB2 DNA transcription templates were generated by denaturing two complementary oligonucleotides (Supplementary Data 5) at 95 °C for 5 min and slow cooling (~0.1 °C/min) to room temperature in 1X annealing buffer (100 mM potassium acetate and 30 mM HEPES, pH 8.0). Annealed oligonucleotides were then purified by resolving them on a 20% Tris–borate–EDTA (TBE)–PAGE gel, isolating the band of expected size and eluting at room temperature overnight in 1X annealing buffer. The eluted DNA template was then ethanol precipitated, resuspended in MilliQ H₂O and quantified using the Qubit dsDNA BR Assay Kit. The KB2 DNA template for the Cu NIMPLY Zn circuit was constructed by placing the tetO region immediately upstream of the T7 promoter and a 3′-region ending with either 3WJdB or the T7 terminator, with the primers listed in Supplementary Data 5. Amplified templates were then purified (QIAquick PCR purification kit, Qiagen, catalog no. 28106) and verified for the presence of a single DNA band of expected size on a 2% Tris–acetate–EDTA (TAE)–agarose gel, and concentrations were determined using the Qubit dsDNA BR Assay Kit and verified for the presence of a single band of expected size on a 1% Tris–acetate–EDTA–agarose gel.

Expression and purification of aTFs. Supplementary Table 5 provides a listing of the exact expression and purification process for each aTF. In general, with the exception of CadC and AdcR, each protein was expressed and purified as follows: sequence-verified petET-28c plasmids were transformed into the Rosetta 2(DE3) pLysS E. coli strain for protein expression. A single transformed colony from each expression plasmid transformation was then used to seed a small overnight culture grown at 37 °C. Overnight cultures were then seeded into larger expression cultures (10 ml of LB starter culture) and expression cultures (100 ml of LB starter culture). Cultures were induced for overexpression at an optical density (600 nm) of ~0.5 with 100–500 µM isopropyl-β-D-thiogalactopyranoside and grown for 4 h more at 37 °C or overnight at 30 °C. Cultures were then pelleted by centrifuging at 4,000 rpm for 20 min at 4 °C in a Thermo Scientific Sorvall Lynx 4000, and then were either immediately stored at ~80 °C or resuspended in lysis buffer (10 mM Tris–HCl, pH 7.5–8.5, 500 mM NaCl, 1 mM tris(2-carboxyethyl)phosphine (TCEP) and protease inhibitor (complete EDTA-free Protease Inhibitor Cocktail, Roche)). Resuspended cells were lysed on ice through sonication (Bertan sonicator) for 1 min, with 2 min of rest on ice in between rounds. Lysates were then centrifuged at 13,000 g for 30 min to remove insoluble material. Clarified supernatants were purified using His-tag affinity chromatography with an Ni-NTA column (HisTrap FF 5 ml column, GE Healthcare Life Sciences), followed by size exclusion chromatography (Superdex HiLoad 26/600 2-00 pg column, GE Healthcare Life Sciences) using an AKTAxpress fast protein liquid chromatography system. The collected fractions from the fast protein liquid chromatography were concentrated and buffer exchanged (25 mM Tris–HCl, 100 mM NaCl, 1 mM TCEP, 50% glycerol, v/v) using centrifugal filtration (Amicon Ultra-0.5, Millipore Sigma). Protein concentrations were determined using the Qubit Protein Assay Kit (Invitrogen, catalog no. Q32850) for the data shown in Fig. 2d and Supplementary Fig. 3, and the purifying the protein previously described 5. For CadC purification, the protocol was modified to have a higher concentration of TCEP (5 mM) in its buffers.

IVT reactions. NEB HiScript T7 Quick High Yield RNA Synthesis Kit (New England Biolabs, catalog no. E2050S) was used for the data shown in Fig. 2d, Supplementary Figs. 2 and 3, and Extended Data Fig. 1, according to the manufacturer’s protocol, with the addition of 2.23 mM dDFHBI-1T (Tocris, catalog no. 2901) to the reaction mix. Briefly, IVT reactions were purified by adding the following components listed at their final concentration: IVT buffer (40 mM Tris–HCl, pH 8, 8 mM MgCl₂, 10 mM DTT, 20 mM NaCl and 2 mM spermidine), 2.25 mM dDFHBI-1T, 1.14 mM Tris-buffered nucleotide triphosphates, pH 7.5, 0.3 U thermostable inorganic pyrophosphatase (New England Biolabs, catalog no. M0296S) was used for the data shown in Fig. 2d and Supplementary Fig. 3, and the purifying the protein previously described 51. For CadC purification, the protocol was modified into pET-3d and pET-3a plasmid backbones, respectively, were purified with the protocol previously described 5. CadC and AdcR, the genes of which were cloned into PET-3d and PET-3a plasmid backbones, respectively, were purified with the protocol previously described 5. CadC purification, the protocol was modified to have a higher concentration of TCEP (5 mM) in its buffers.

Freeze-drying. Before lyophilization, PCR tube caps were punctured with a pin to create three holes. Lyophilization of ROSALIND reactions was then performed by assembling the components of regulated IVT (see IVT reactions) with the addition of 30 mM sucrose and 250 mM β-mannitol. Assembled reaction tubes were immediately transferred into a prechilled aluminum block and placed in a −80 °C freezer for 10 min to allow slow freezing. After the slow freezing, reaction tubes were wrapped in Kimwipes and aluminum foil, submerged in liquid nitrogen and then transferred to a FreeZone 2.5-1 Bench Top Freeze Dry System (Labconco) for overnight freeze-drying with a condenser temperature of −85 °C and 0.04 mbar of pressure. Unless rehydrated immediately, freeze-dried reactions were packaged as follows: the reactions were placed in a light-protective bag (Mylar open-ended food bags, Uline, catalog no. S-11661) with a desiccant (Drier-Card Desiccants, Uline, catalog no. S-19582), purged with argon using an argon canister (Art Wine Preserver, Amazon, Amazon Standard Identification Number (ASIN) B01MHICQ2Q) and immediately impulse heat sealed (Metrocin 8 inch Impulse Bag Sealer, Amazon, ASIN B06XC7C6ZV) (Supplementary Video 2).

Plate reader quantification and MEF standardization. A National Institute of Standards and Technology (NIST) traceable standard (Invitrogen, catalog no. F36915) was used to convert arbitrary fluorescence measurements to MEF. Serial dilutions from a 50-µM stock were prepared in 100 mM sodium borate buffer at pH 9.5, including a 100-µM sodium borate buffer blank (total of 12 samples). The samples were prepared in technical and experimental triplicates (12 samples × 9 replicates = 108 samples total), and fluorescence values were read at an excitation wavelength of 472 nm and emission wavelength of 507 nm for 3WJdB-activated fluorescence, or 485-nm excitation and 515-nm emission for dDFHBI-1T expression a plate reader (Systech H1, BioTek). Fluorescence values for a concentration in which a single replicate saturated the plate reader were excluded from analysis. The remaining replicates (nine per sample) were then averaged at each FITC concentration, and the average fluorescence value of the blank was subtracted from all values. Linear regression was then performed for NATURE BIOTECHNOLOGY | www.nature.com/naturebiotechnology
concentrations within the linear range of fluorescence (0–6.25 μM FITC) between the measured fluorescence values in arbitrary units and the concentration of FITC for identifying the conversion factor. For each plate reader, excitation, emission and gain settings were followed for conversion factor. Calibration curves were used to correlate arbitrary fluorescence values to MEF (μM FITC) (Supplementary Data 3).

To characterize ROSALIND reactions, 19-μl ROSALIND reactions were loaded onto a 384-well, black, opaque clear, flat-bottomed plate using a multichannel pipette and measured on a plate reader (Synergy H1, BioTek). Kinetic analysis was performed by reading the plate at 3-min intervals with excitation and emission wavelengths of 472 and 507 nm, respectively, for 4 h at 37°C. PureXpress reactions were quantified and standardized to MEF (μM FITC) as described for ROSALIND reactions, but with excitation and emission wavelengths set to 485 and 515 nm, respectively.

We note that no background subtraction was performed when analyzing outputs from any ROSALIND reaction. An example of this standardization procedure is shown in Supplementary Fig. 1.

Municipal and surface water sampling. Municipal water was collected from water fountains in public parks of Paradise, CA, in October of 2019, as the town recovered from the most destructive wildfire in California’s history (Camp Fire, November 8, 2018). This urban fire severely damaged the municipal water system in Paradise, resulting in widespread contamination of drinking water pipes. After the fire, a Water Quality Advisory was issued, asking the public not to drink municipal water. With the Water Quality Advisory still in effect at the time of sampling, public water fountains had probably remained unflushed for ~11 months. Water samples were collected from self-standing fountains (brass or stainless steel) using metal-free plastic tubes and filtered (0.45-μm nylon membranes) in the field using acid-washed plastic syringes. Then, 50-ml filtered subsamples were immediately acidified to pH ~2 with HNO₃, for metal analysis (FAAS) and kept cold (4°C) with 50-ml filtered, unacidified subsamples until their return to California State University, Chico, where ROSALIND tests were initiated within 4 h of collection using filtered, unacidified sub-samples.

Tests were initiated by applying 20 μl of filtered unacidified samples to reaction tubes, or laboratory-grade, metal-free water (ultrapure type 1 water, resistivity >18 MΩ cm), as indicated. Reactions were incubated in a portable incubator at 37°C and periodically monitored. Photographs were taken of reaction tubes after 2 h and overnight using the handheld incubator.

Reaction strips and water samples were shipped back to Northwestern University for fluorescence measurements by the plate reader (see Plate reader quantification and MEF standardization) and metal analysis (see FAAS measurement).

For Chilean environmental samples, river water samples were taken from the Yerba Loca Creek in the upper Mapocho–Maipo watershed, and the Toro River in the upper Elqui River watershed. Both streams are metal-polluted tributaries to rivers that provide ecosystem services (for example, drinking water sources, habitat for ecosystems and recreation) to major Chilean cities. Acid-washed high-density polypropylene bottles were used to collect 200 ml of filtered (0.45-μm nylon membranes) samples and were shipped in a cooler. Given the low pH (<pH 4) and the objective of the experiments, the pH was not adjusted.

For copper/zinc-spiked municipal water from Evanston, IL, two bottles of approximately 250 ml of the water samples were collected from a drinking fountain. One of the bottles was then filtered at 0.45 μm using 47 mm cellulose acetate filter membranes (Thermo Fisher Scientific). The other was spiked with a known amount of CuSO₄/ZNSO₄ and then filtered using the same method. Then, the copper/zinc-spiked fountain water was diluted with the filtered nonspiked fountain water to create 40-ml solutions in the range of 0.125–4 ppm for copper and 0.05–3 ppm for zinc. Of these water samples, 10 ml was then acidified with HNO₃, and measured using FAAS for concentration validation (see FAAS measurement).

Of the water samples, 20 μl was used to rehydrate freeze-dried CsSO₄, SbIIb and Cu NIMPY Zn fibers. Upon rehydration, fluorescence measurements of the reactions were performed by a plate reader (see “Plate reader quantification and MEF standardization”). For the copper-spiked Michigan water sample, the same method was used as for the Lake Michigan water sample.

FAAS measurement. An FAAS instrument (PerkinElmer PinAAcle 900) equipped with PerkinElmer Lumina hollow-cathode lamps was used to determine copper and zinc concentrations in the drinking water spiked with copper and zinc (Supplementary Fig. 6). The lake water spiked with copper, the Chilean surface water samples (Extended Data Fig. 10) and the Paradise municipal water samples (Fig. 6 and Supplementary Fig. 7). The spectral lines were 324.8 nm for copper and 213.9 nm for zinc. FAAS was calibrated for CuII using CuN, reference solution as a standard (Thermo Fisher Scientific, catalog no. CAS10031–43–3) and for Zn(II) using certified International Organization for Standardization Guide 17034 standard solution (Fluka Analytical, catalog no. 18562), with calibration curves comprising between 0.05 and 2 ppm for copper and 0.05 and 1 ppm for zinc. Samples were diluted with 18 MΩ cm deionized water (Milli-Q, Milipore). When necessary, and acidified with ultrapure HNO₃ (Fisher Chemical, catalog no. CAS6797–37–2). Three replicates were performed and averaged, and error propagations were taken into account to estimate uncertainties.

Photographs of reactions. Photog rafts of reaction tubes in Figs. 2 and 6 were taken on an Apple iPhone X and a Google Pixel 3a using their default camera applications, respectively. Photographs in Extended Data Fig. 8 were taken on an Apple iPhone 15, carried out using the default Apple iOS Camera application. The resulting images were only cropped. Source image files are available as Supplementary Data 1.

Statistics and reproducibility. The number of replicates and types of replicates performed are described in the legend to each figure. Individual data points are shown and, with enough replicates present, the mean ± s.d.; this information is provided for each figure. The type of statistical analysis performed in Fig. 5 and Extended Data Fig. 4 is described in the legend to each figure. Exact P values, along with degrees of freedom computed from the statistical analysis, can be found in Supplementary Data 3.

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

Data availability All data presented in this manuscript are available as supplementary data files. All plasmids used in this manuscript are available in Addgene with the identifiers 140371–140399. Source data are provided with this paper.

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Author contributions J.K.J., K.K.A., J.J.C. and J.B.L. conceived the project. J.K.J., K.K.A., M.S.V. and J.B.L. curated the data. J.K.J., K.K.A. and J.B.L. carried out the formal analysis. J.B.L., J.-F.G., S.J.M., D.P.G. and P.A.P. acquired the funds. J.K.J., K.K.A., M.S.V., M.D., P.R.C., J.W.L. and J.-F.G. carried out the investigations. J.K.J., K.K.A., M.S.V., J.W.L., P.Q.N., D.A.C., M.D., S.J.M., J.-F.G. and J.B.L. devised the methodology. J.K.J., K.K.A. and J.B.L. administered the project. D.A.C., D.P.G., P.A.P. and S.J.M. acquired the resources. J.K.J., K.K.A., J.J.C. and J.B.L. supervised the project. J.K.J., K.K.A., M.S.V., M.D., P.R.C. and J.-F.G. validated the project. J.K.J., K.K.A. and J.B.L. visualized the project. J.K.J., K.K.A. and J.B.L. wrote the original draft. J.K.J., K.K.A., M.S.V., M.D., P.R.C., J.-F.G. and S.J.M. wrote the final manuscript. J.K.J. and J.B.L. edited the manuscript.

Competing interests K.K.A., J.K.J. and J.B.L. have submitted a US provisional patent application (no. 62/758,242) relating to regulated IVT reactions. K.K.A., J.K.J., M.S.V., P.C., J.W.L., J.J.C. and J.B.L. have submitted a US provisional patent application (no. 62/838,852) relating to regulated IVT reactions. K.K.A., M.S.V., J.W.L., P.Q.N., D.A.C., S.J.M., P.A.P., J.-F.G. and J.B.L. have a financial interest in Stemloop, Inc. The latter interests are reviewed and managed by Northwestern University in accordance with their conflict-of-interest policies.

Additional information Extended data is available for this paper at https://doi.org/10.1038/s41587-020-0571-7. Supplementary information is available for this paper at https://doi.org/10.1038/s41587-020-0571-7. Correspondence and requests for materials should be addressed to J.B.L. Reprints and permissions information is available at www.nature.com/reprints.
Extended Data Fig. 1 | Homemade transcription reactions can be optimized to perform as well as commercial kits. Homemade in vitro transcription reactions were compared to a commercially available high yield transcription kit (NEB HiScribe™ T7 Quick High Yield RNA Synthesis Kit). 25 nM DNA encoding T7-3WjdB-T was added to each reaction in a total reaction volume of 20 µL. Over the course of 1 hour, the data show similar fluorescence activation. However, the homemade reaction begins to saturate after 1 hour, likely due to the exhaustion of NTPs in the homemade reaction (11.4 mM) when compared to the commercial kit (40 mM). All data shown for n = 3 independent biological replicates as lines with raw fluorescence values standardized to MEF (µM FITC). Shading indicates the average value of 3 independent biological replicates ± s.d.
Extended Data Fig. 2 | Design and optimization of a ROSALIND reaction. Constructing a ROSALIND reaction begins with identifying a ligand of interest, an aTF capable of binding or unbinding a DNA operator sequence as a function of the ligand and the aTF’s cognate operator sequence. Once identified, two DNA constructs are designed, one to separately express and purify the aTF and the other to generate a linear transcription template encoding the promoter, spacer, operator, fluorescent RNA aptamer and optional terminator. The first step in optimizing the sensor is performing a titration of the purified aTF at a fixed DNA template concentration to determine the amount of aTF needed to fully repress the expression of 3WjdB. Then, the target ligand is titrated (at the concentration of aTF previously determined) to test for induction. If the sensor needs improvement, components of ROSALIND can be redesigned and retested. The data shown here are for the Tetr-based ROSALIND reaction described in Fig. 2f,h. The bar graphs shown are data taken at 4 h after initiating reactions with T7 RNAP and 25 nM DNA template, and the kinetics data are shown for 0 µM (green lines) and 1.25 µM Tetr dimer (purple lines). Fluorescence-activation is substantially repressed at 25-fold excess or greater of Tetr dimer over DNA template. The Tetr-aTc dose response curve shown are the data presented in Fig. 2h, and the kinetics data are shown for 0 µM (purple lines) and 2.5 µM aTc (green lines). All data shown for n = 3 independent biological replicates as points or lines with raw fluorescence values standardized to MEF (µM FITC), and bars representing averages of the replicates. Shading and error bars indicate the average value of 3 independent biological replicates ± s.d.
Extended Data Fig. 3 | Characterization of a ROSALIND dose response curve. A dose response curve of a TetR-regulated ROSALIND reaction is plotted from the measured and calibrated kinetics traces, using the 4-h end point fluorescence values at different ligand concentrations. a, kinetic traces of TetR–aTc induction at different aTc concentrations, and b, the corresponding 4-h fluorescence values plotted against the μM of aTc added. Reactions are generated using 25 nM of DNA and 1.25 μM of TetR dimer. All data shown for n = 3 independent biological replicates as lines (a) or points (b) with raw fluorescence values standardized to MEF (μM FITC). Shading (a) and error bars (b) indicate the average value of 3 independent biological replicates ± s.d.
Extended Data Fig. 4 | Dose response curves of ROSALIND reactions. The dose response curves of ROSALIND reactions shown in Fig. 3 are presented. The amount of DNA templates and aTF used in each panel are configured as described in Supplementary Table 2. All data shown for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (µM FITC). Error bars indicate the average value of 3 independent biological replicates ± s.d. The ligand concentrations at which the signal is distinguishable from the background were determined using two-tailed heteroscedastic Student’s t-test against the no ligand condition, and their p-value ranges are indicated with black asterisks (*** < 0.001, ** = 0.001–0.01, * = 0.01–0.05). The asterisks indicated in red have p-values in the range between 0.05–0.15 due to variability between replicates, although the average signals were clearly above background. Exact p-values along with degrees of freedom can be found in Supplementary Data 3. Data for no ligand conditions were excluded in the plots since the x-axis is on the log scale, but they can be found in Supplementary Data 3.
Extended Data Fig. 5 | Orthogonality of ROSALIND reactions. The corresponding bar graph data of the orthogonality matrix shown in Fig. 4a are presented. All data shown for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (µM FITC), and bars representing averages of the replicates. Error bars indicate the average value of 3 independent biological replicates ± s.d. Tet, Tetracycline; Ery, Erythromycin; 3-OH BA, 3-Hydroxybenzoic Acid; BAC, Benzalkonium chloride.
Extended Data Fig. 6 | RNA kleptamers are used to invert transcription factor responses. a, A kleptamer RNA (KB2) antisense to the dye-binding region of the broccoli aptamer can disrupt folding of 3WJdB and lead to the loss of fluorescence. Addition of a KB2-expressing template in a 4:1 ratio with the 3WJdB template inhibits signal. b, KB2 transcription can be regulated by TetR by placing the tetO site in between the T7 promoter and KB2 coding sequence. In the presence of 1.25 μM TetR dimer, the KB2 transcription is fully repressed, and the signal from 3WJdB is restored. c, Kleptamers can be used to invert the response of transcription factors when KB2 is regulated by TetR (1.25 μM dimer) and induced by aTc (2.5 μM shown). d, This scheme was used to create a ROSALIND zinc sensor with the aporepressor AdcR. When bound to Zn²⁺ (30 μM), AdcR (1.5 μM dimer) binds to its cognate operator sequence, adcO, placed upstream of the KB2 coding sequence, preventing KB2 expression and thereby activating fluorescence from 3WJdB. Arrows inside of the plots represent direction of regulation when indicated species are added. All data shown for n = 3 independent biological replicates as lines with raw fluorescence values standardized to MEF (μM FITC). Shading indicates the average value of 3 independent biological replicates ± s.d. 3WJdB template concentrations used are: 25 nM for a-c and 7.5 nM for d. KB2 template concentrations used are: 100 nM for a, and 150 nM for b-d.
Extended Data Fig. 7 | ROSALIND sensitivity can be tuned by changing the concentration of transcription factor. a, Increasing the concentration of CsoR desensitizes the copper sensor (dose response shift to higher concentrations), while b, decreasing the concentrations of CadC sensitizes the lead sensor (dose response shift to lower concentrations). The red arrows in the plots indicate the direction of dose response shift. All data shown for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (µM FITC). Error bars indicate the average value of 3 independent biological replicates ± s.d. 3WJdB DNA template concentrations used are: 25 nM for a and 1.5 µM CadC condition in b, and 10 nM for 0.5 µM CadC condition in b.
Extended Data Fig. 8 | Stability of ROSALIND. **a**, The fluorescence signal generated from unregulated ROSALIND reactions are stable over weeks. The increase in fluorescence from Day 1 is likely due to a concentration increase caused by evaporation when the plate was taken out of the incubator after the first measurements. **b**, The shelf-stability of freeze-dried ROSALIND reactions (unregulated, TetR-regulated and αTc-induced) decay over the course of a month without proper packaging. **c**, Packaging of freeze-dried ROSALIND: 1) reactions are lyophilized overnight, 2) the overnight lyophilized reactions are purged with inert gas such as argon and 3) the reactions are placed into a light-protective bag with a desiccant and immediately impulse heat sealed (Supplementary Video 2). **d**, When this packaging method is implemented, lyophilized reactions are functional out to 2.5 months. Though we observed signal decay, the signal from rehydrated reactions after 2.5 months is clearly visible. Images are shown for one replicate with other replicate images included in Supplementary Data 1. Unregulated reactions were lyophilized with 25 nM of the 3WJdB template, and TetR-regulated reactions with additional 1.25 µM TetR dimer along with the components of IVT specified in the *in vitro transcription reactions* method section. Unregulated and TetR-regulated reactions were then rehydrated with laboratory-grade water, and αTc-induced reactions were rehydrated with 10 µM of αTc. All data shown for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (µM FITC), and center values representing averages of the replicates. Error bars indicate the average value of 3 independent biological replicates ± s.d. The original, uncropped images shown in **c** and **d** can be found in Supplementary Data 1.
**Extended Data Fig. 9 | Handheld fluorescence illuminator design.** a, A low-cost handheld fluorescence illuminator enables point-of-use functionality for ROSALIND. 3D printed components include a, b, front case, c, back case and d, battery cover. A printed circuit board (e, front view facing back case, f, rear view facing front case) mounts in the front case and connects LEDs, resistors, a trimmer potentiometer, a power switch and a battery holder. Full CAD files, 3D print files (.STL), PCB design files and assembly instructions are provided in Supplementary Data 4. The estimated cost of building one device is $8.69 USD.
Extended Data Fig. 10 | ROSALIND reactions can be freeze-dried and rehydrated with an environmental surface water source. a, The ROSALIND reactions shown in Fig. 6 were characterized for response to CuSO₄ as in Supplementary Fig. 6, but using water from Lake Michigan (collected in Evanston, IL). We observed lower signals from these spiked environmental samples than with the laboratory-grade water samples due to uncharacterized matrix effects. b, To test ROSALIND functionality on real-world environmental samples, surface freshwater samples were obtained from two streams known to contain elevated levels of copper: (1) the Yerba Loca Creek, upstream from Santiago, in the Maipo watershed of Central Chile, and (2) the Toro River, upstream from La Serena, in the Elqui watershed of Northern Chile. c, d, Copper concentrations of the samples from the Yerba Loca Creek (c) and the Toro River (d) were determined using Flame Atomic Absorption Spectroscopy (FAAS) and found to be in the range of 6.9–8.5 ppm. FAAS was calibrated with a Cu(NO₃)₂ standard solution as shown in e. Serial dilution was performed on each sample to create three separate diluted samples that are in the operating ppm range of FAAS, and their ppm values were measured and reported in Table f as the average value of 3 independent replicates ± s.d. Using these measurements, linear regression on the averages of each dilution was performed to calculate the ppm value of the undiluted sample. g, h, In order to generate a series of tests that cover our detectable copper range, we serially diluted each sample with laboratory-grade, metal-free water (undiluted, 1:2, 1:4, 1:8, 1:16 and 1:32 dilutions) and measured their concentrations by FAAS before using this series to rehydrate freeze-dried copper sensors. The copper concentrations indicated were taken from c, d, or calculated from the extrapolated ppm value of the undiluted sample in Table f. When rehydrated with the 1:8 dilution, we observed clear visible signals, corresponding to copper concentrations of 0.88 ppm and 1.05 ppm from the Yerba Loca Creek and Toro River samples, respectively. Four-hour end-point data are shown in a, g, h for n = 3 independent biological replicates as points with raw fluorescence values standardized to MEF (µM FITC), and bars representing averages of the replicates. Error bars indicate the average value of 3 independent biological replicates ± s.d. Geographical data in b © OpenStreetMap contributors 46.
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| Sample size | No sample size calculation was used. Sample size of n=3 was chosen for quantitative fluorescence measurements based on similar cell-free / in vitro studies. All individual data points are shown in figures, and total sample size spanning all independent experiments is reported in the figure captions. |
| Data exclusions | No data on presented systems were excluded from the analyses. When field testing we did test another preliminary circuit idea we did not discuss in the manuscript. Photographs of these tests are included and annotated in Supplementary Data File 1 (because they were taken alongside the other included circuit(s), but not discussed in the manuscript). |
| Replication | All experiments were performed three times (independent biological triplicates) by at least one co-author. All attempts at replication were successful. |
| Randomization | While randomization is not relevant to this study since our biochemical reactions are handled uniformly, the same data analysis procedure was applied to all samples of the same type. |
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