A primer on emerging field-deployable synthetic biology tools for global water quality monitoring

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Tracking progress towards Target 6.1 of the United Nations Sustainable Development Goals, “achieving universal and equitable access to safe and affordable drinking water for all”, necessitates the development of simple, inexpensive tools to monitor water quality. The rapidly growing field of synthetic biology has the potential to address this need by isolating DNA-encoded sensing elements from nature and reassembling them to create field-deployable “biosensors” that can detect pathogenic or chemical water contaminants. Here, we describe current water quality monitoring strategies enabled by synthetic biology and compare them to previous approaches used to detect three priority water contaminants (i.e., fecal pathogens, arsenic, and fluoride), as well as explain the future potential for engineered biosensors to simplify and decentralize water quality monitoring. We conclude with an outlook on the potential for engineered biosensors to simplify and decentralize water quality monitoring. We conclude with an outlook on the potential for engineered biosensors to simplify and decentralize water quality monitoring.

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INTRODUCTION

Reliable access to clean drinking water is essential for human well-being, economic development, and political stability. Impaired water quality, quantity, and accessibility, however, are projected to increase both in frequency and severity due to population increase, climate change, persistent water infrastructure degradation, and poor water governance. As such, institutions like the World Economic Forum and the US Government have identified the burgeoning water crisis as a top global threat that may undermine progress in protecting human health and serve as a structural driver of poverty and inequity.

The turn of the millennium saw the creation of the United Nations (UN) Millennium Development Goals—eight humanitarian grand challenges to be resolved by 2015. These goals were monitored and refined over the next 15 years, and after an extensive revision process, 2016 saw the launch of the Sustainable Development Goals (SDGs) for 2030, each of which is accompanied by targets and progress indicators. Sustainable Development Goal (SDG) 6 aspires to the availability and sustainable management of water and sanitation for all, with SDG Target 6.1 seeking to “achieve universal and equitable access to safe and affordable drinking water for all.” Progress towards SDG 6.1 is tracked by Indicator 6.1.1, “the proportion of population using safely managed drinking water services,” defined as services that are located on premises, available when needed, and free from contamination. The Joint Monitoring Program (JMP), housed within the United Nations Children’s Fund (UNICEF) and the World Health Organization (WHO), is the official UN mechanism that has been tasked with monitoring progress towards this goal.

Accurate tracking and surveillance of global drinking water sources will require significant advances in water quality monitoring technology. Although location on premises and availability when needed can be relatively easily quantified, objectively determining drinking water safety (i.e., if a source is “safely managed”) necessitates the use of technologies to detect the presence of specific contaminants. There are countless potential contaminants that could pose health risks; JMP focuses on three that are globally prevalent and universally recognized as deleterious to human health: arsenic and fluoride (naturally abundant chemical contaminants), and Escherichia coli (an indicator of fecal contamination). Owing to the ubiquity of these contaminants and resource limitations in most affected areas, ideal technologies for global water quality monitoring should be inexpensive, simple enough for an untrained individual to use, and capable of rapidly (within minutes to hours) providing results onsite. Notably, they do not necessarily need to be quantitative; the ability to determine if a contaminant is above or below a risk threshold can provide sufficient actionable information, though technologies that can provide quantitation would enhance their use and impact. Current gold-standard methods for assessing water quality, however, do not fulfill these criteria. Most technologies require expensive equipment and reagents, reliable electricity sources, technically skilled operators, and transportation infrastructure. For example, the equipment to run qPCR (a DNA amplification technique for pathogen detection) and mass spectrometry (a molecular analysis technique for chemical detection) costs tens of thousands of dollars excluding operational expenses, must be operated by a trained technician, and cannot be brought into the field, thus...
necessitating sample transport for centralized analysis. As such, these methods come at a significant resource burden, which prohibits widespread deployment.

While there has been progress in developing more user-friendly field kits capable of rapidly detecting even trace contaminant levels, there is still significant work to be done before they can be widely adopted for global monitoring or individual use. Existing field kits frequently require sample processing steps that are beyond the skill level of an untrained user, along with expensive supplemental equipment or consumables, which are often hazardous chemicals. Collectively, these limitations preclude the scale and frequency of monitoring that is needed to effectively track progress towards SDG 6.1. There is thus an urgent unmet need for low-cost, field-deployable water quality tests, as evidenced by the UN High Level Panel on Water’s call for higher resolution data on water quality to better address the global water crisis.

The growing field of synthetic biology, which centers around the design and construction of biological systems, is poised to address this knowledge gap by engineering and repurposing microbial biosensors. In nature, microbes use biosensors to detect and respond to changes in their environment. For example, a biosensor for detecting toxins may activate the production of proteins that export, neutralize, or metabolize the toxins. By deconstructing and modifying naturally occurring microbial biosensors, we can create synthetic, genetically encoded biosensors (henceforth referred to as “biosensors”) to detect targets of global concern. Biosensors have already been developed to detect a wide range of chemicals, as well as bacterial and viral pathogens. Recently reported biosensors have even been packaged in handheld, easy-to-use formats, facilitating widespread field deployment.

Because of their potential to significantly advance the field of water quality monitoring, we seek to provide a primer on emerging biosensors. We specifically focus on the development of field-deployable biosensors— inexpensive, portable tools that can be used onsite by individuals without technical expertise. While we focus solely on purely genetically encoded biosensors in this review, we note there exist other developing biosensors that are not purely genetically encoded that are covered in other excellent reviews. We begin with a conceptual overview of how the gene expression process can be leveraged for biosensing and discuss the design process for a biosensor. We then discuss their potential applications for detecting Escherichia coli, arsenic, and fluoride, as well as other emerging targets including metals and pharmaceuticals. We conclude with an outlook on the future of synthetic biology for water quality monitoring, identifying needs in the field and necessary steps for widespread implementation.

**BIOSENSOR DESIGN AND CONSTRUCTION**

At the core of synthetic biology is the idea that biological systems can be deconstructed into sets of biological parts, which are individual biomolecules with discrete functions (Box 1). Each part is written into DNA, which serves as a genetic blueprint. Once they are written into DNA, the individual parts can be manufactured and assembled into a larger functional system (Fig. 1). Synthetic biology works to construct new DNA blueprints that repurpose and reengineer existing biological parts to produce technologies for high-value applications, such as manufacturing food and fuels, creating medicines, and developing diagnostics.

Biosensors are molecular systems that detect and respond to specific targets. All biosensors are constructed from two modular parts—a sensor and a reporter. First, the sensor recognizes a target of interest. Once recognized, the sensor changes its shape to initiate production of the reporter. The reporter then creates a detectable output, typically in the form of fluorescence or color change.

Natural biosensors govern a microbe’s interaction with its environment and exist for every molecule that microbes naturally sense and respond to. They also serve as a starting point for building biosensors, which can be designed, evolved, and engineered to detect targets of interest. To create a biosensor, the DNA encoding the sensor and the reporter is placed in a platform supporting biological function—typically a live cell or cell-free solution containing the cellular machinery needed for transcription and translation.

This review focuses on biosensors for their potential as low-cost, rapid, and field-deployable water quality monitoring devices. Here, we discuss each component of a biosensor and the overall design process for building them.

**Sensor parts**

Sensor parts are molecules that detect a target compound. These molecules can be either natural or engineered and are most commonly nucleic acids or proteins that fold into intricate shapes to match the physical and chemical properties of their targets. One challenge with harvesting parts from nature, however, is that their sensitivity (i.e., ability to detect a particular concentration) or specificity (i.e., ability to distinguish a target from other surrounding molecules) may not meet the requirements for a given application. For example, some natural metal sensors interact with multiple metal ions rather than a single-specific target.

Fortunately, these properties can be adjusted through biomolecular engineering approaches that change the underlying molecular shape and chemistry of the sensor to match the desired sensitivity or specificity.

**Reporter parts**

Reporter parts are molecules that produce detectable signals. Like sensor parts, they can be nucleic acids or proteins, but they are more varied in their modes of operation because of their range of potential outputs. Fluorescent reporters, the most commonly used reporter type, are molecules that produce a fluorescent signal when illuminated by specific wavelengths of light. They come in a range of colors and generally require external illumination to visualize their fluorescence. In contrast, colorimetric reporters are enzymes that react with a supplied substrate to produce a visible color. While colorimetric reporters do not require additional equipment to interpret their results, their outputs are more difficult to quantify than fluorescent outputs. Although other reporters that produce luminescent or electrical outputs exist, we focus on fluorescent and colorimetric reporters given their widespread use in the majority of reported biosensors, as well as their relative field-friendliness.

**Biosensor platforms**

Biosensors must be housed in a platform that supports the biological processes needed for them to operate. For many sensors, this requires an environment that can support transcription and translation. There are two main biosensing platforms enabling this: “whole-cell” biosensors, which are live cells genetically modified to express the sensor and reporter, and “cell-free” biosensors, which consist of either cellular extract or purified cellular machinery that can perform the processes of gene expression. While both of these platforms are amenable to field deployment because they can be freeze-dried for transport and long-term storage, they each have individual strengths and weaknesses that must be considered during biosensor design.

Whole-cell biosensors have several important advantages. As living sensors, they can be simply and inexpensively mass-produced by allowing the engineered microbe to multiply. They
Box 1. Gene expression and the central dogma of molecular biology

Gene expression is the process by which the information encoded in DNA is transcribed into RNA, which is then translated into proteins. This flow of information gives rise to the rich diversity of biological function and is known as the central dogma of molecular biology. 

Deoxyribonucleic acid (DNA) serves as the blueprint for guiding how life develops and functions. It is made of individual components called nucleotides, which are linked together to form longer strands called nucleic acids. The specific order of these nucleotides in a DNA strand is called its sequence and determines what information is stored within the DNA. The totality of the information in an organism’s DNA is called its genome. In bacteria, the genome consists of one long, circular piece of DNA with a sequence unique to each particular bacterial strain.

Ribonucleic acid (RNA) is produced from DNA in a process called transcription. While both are nucleic acids that can fold into complicated structures, they differ in their function; DNA is used for information storage while RNA is used for information processing. Broadly speaking, RNA can be divided into two categories—messenger RNA (mRNA) and noncoding RNA (ncRNA). mRNA carries information from DNA that guides protein production, while ncRNA regulates the steps of gene expression and many other cellular processes. RNA-based sensors are examples of ncRNA; while they do not code for proteins, they fold into analyte-binding structures to either control protein production or generate a signal in response to ligand binding.

Proteins are produced from mRNA in a process called translation. They are composed of chains of individual components called amino acids that fold into complicated structures and have a staggering diversity of functions, ranging from carrying information to structurally supporting the cell. In this review, we focus primarily on sensor and reporter proteins.

These advantages are counterbalanced by the fact that it is difficult to use a part’s performance in a live cell to predict its function in a cell-free platform. Indeed, many sensors require assistance from pieces of peripheral cellular machinery to function properly and the exact differences between the composition of a cell-free reaction and a live cell are still unclear. Despite this, significant progress has been made towards optimizing cell-free systems to accept sensor parts.

Interaction between sensors and reporters

When placed in a biosensing platform, a sensor controls the activation of a reporter by suppressing its signal until the sensor recognizes its target. As the reporter is only produced when this recognition occurs, the reporter’s signal indicates that the sensor’s target is present. This interaction between a paired sensor and reporter is guided by the way that they are written into the biosensor’s DNA blueprint. For example, protein-based sensors can bind to specific regions of DNA to physically block production of the reporters they regulate, attaching or releasing based on the presence of their target. In contrast, RNA-based sensors can fold into different shapes based on whether or not a target is present, with different configurations allowing or preventing reporter production. This diversity of sensor and reporter functions, combined with the staggering number of possible sensor-reporter combinations, also better replicate the cellular environment that the sensors evolved to function in.

Use of live hosts, however, also presents several challenges. For instance, whole-cell biosensors must be kept alive during use, requiring bacterial growth media and potentially a field-deployable incubator, which increases the amount of supplemental equipment that must be brought into the field. Furthermore, whole-cell biosensors can only detect targets that do not kill the cell. The synthetic DNA engineered into the cell may also mutate or be lost as cells grow and divide, preventing or distorting sensor and reporter production. Furthermore, the use of live cells inherently confers biocontainment concerns, though methods to encapsulate or disable whole-cell sensors are being explored to mitigate this risk.

Cell-free biosensors aim to emulate the cellular environment in a non-living system. Placing biosensor DNA in a cell-free gene expression reaction allows the system to act in much the same way as a whole-cell biosensor, but without the complications of needing to maintain and contain living cells. Cell-free biosensors can also be easily tuned and optimized by changing the concentration of the biosensor DNA or other reaction components, which is more difficult to do in a living cell. Additionally, because some of the physical and biological constraints of live cells are removed, such as the cell’s outer membrane that restricts the import of some targets, they can detect a broader range of contaminants.
Components of a Synthetic Biosensor

**Platforms**
- Engineered Cell
- Cell-free System

**Sensors**
- Protein
- RNA
- Enzyme

**Reporters**
- fluorescent
- colorimetric

**Fig. 1 The three components of a biosensor: platform, sensor, and reporter.** A sensor is a biomolecule that recognizes a specific target chemical or fragment of a pathogen’s genome. This recognition event activates cellular machinery (gray), which uses gene expression to generate an output signal in the form of a reporter RNA or protein to indicate the presence of the contaminant. When engineering a biosensor, the sensor and reporter are combined in either an engineered cell or a cell-free system that supports the biological reactions necessary to generate a signal. Active Sensors
- Cellular Machinery Activation
- Reporter Production
- Enzyme

**Signal Generation**

**PATHOGEN DETECTION**

Waterborne pathogens, including bacteria, protozoa, and viruses, are leading causes of poor water quality globally that pose both immediate and long-term risks to human health. As such, they are currently amongst the highest priority contaminants of global concern. Fortunately, every pathogenic organism has a unique genetic sequence, which serves as a DNA “barcode” that can be used to identify a specific species and strain in a biosensing reaction. The first step of pathogen detection is sample preparation, where pathogens are broken open to expose their DNA barcodes. These unique DNA sequences are then processed in two steps: amplification of a targeted DNA sequence and production of a signal in response to its detection. This is quite different from existing field-deployable methods that detect secondary indicators of pathogen presence such as H₂S production from bacterial metabolism, presence of indicator protein activity, or biomolecule fluorescence. While these methods are powerful tools for pathogen detection that are currently in use, target DNA sequence detection enables specific pathogen identification, which can provide additional information on water quality and guide treatment more accurately.

There are three basic steps for detecting waterborne pathogens with a biosensor (Fig. 2). While these steps are discussed in the context of detecting fecal coliforms and compared to existing field-deployable coliform detection methods (Table 1), they can be reconfigured to detect virtually any pathogen.

**Target amplification**

Pathogenic DNA in contaminated water is typically only present in trace amounts. To maximize sensitivity, pathogen detection techniques require some form of amplification to increase the amount of target DNA in a sample. Most synthetic biology approaches use isothermal amplification strategies, where DNA is amplified while being held at a single temperature. These methods use some of the natural biological machinery used for DNA and RNA replication; by targeting this machinery towards specific sequences in the genome, it is possible to selectively amplify them for detection. These techniques can therefore be made to be highly specific by targeting the unique barcode regions of specific pathogens.

Each isothermal amplification method differs in temperature and time, although most can bring their targets to detectable levels within two hours. These methods also require minimal training and infrastructure: a freeze-dried reaction containing the biological parts needed for isothermal amplification can be taken to the sample site then deployed by rehydrating with a minimally processed (e.g., syringe-filtered) water sample and incubating, in some cases with body heat or at room temperature. This simplicity eliminates the need for expensive equipment and training, easing deployment in remote and resource limited areas. Collectively, these methods enable detection of even the most dilute pathogens, with methods reporting up to attomolar sensitivity—<10 molecules of DNA in a 10 µL test sample.

**Signal production**

The simplest method for pathogenic DNA detection uses modified DNA molecules that produce a fluorescent output in the presence of the target sequence. As DNA is double-stranded, two interacting strands can be attached to a single modified DNA molecule; a fluorescent molecule called a fluorophore is attached to the first strand, while a quencher that inhibits its fluorescent signal is attached to the second. As isothermal amplification creates more target DNA, the amplified DNA displaces the quenching strand to generate a fluorescent output (Fig. 2). This method has recently been used to detect as few as 10 contaminating E. coli cells in a 50 mL water sample, with a total assay time of 80 min. Recent approaches have further built on this strategy to design RNA-based biosensors that undergo similar structural changes, but activate the expression of a reporter gene in the presence of specific bacterial or viral DNA barcodes.

Beyond its widely known uses for gene editing, Clustered Regularly Interspaced Short Palindromic Repeats, or CRISPR, provides a powerful new method for pathogen detection. CRISPR systems are hybrid protein-RNA biosensors. In CRISPR, portions of a special “guide” RNA target DNA barcode sequences of interest, which are then destroyed by an associated CRISPR protein. In a biosensor, CRISPR systems can be used alongside DNA or RNA sequences labeled by a fluorophore-quencher pair to produce a detectable signal upon target recognition (Fig. 2). Using this strategy, recently developed CRISPR-based sensors have reached the maximum possible specificity by discriminating between pathogen DNA sequences that differ by only a single base pair.

**Outstanding challenges for pathogen detection**

Biosensors target barcode sequences in genomic DNA, which is protected by the cell’s outer wall and therefore inaccessible in an unprepared sample. Because of this, they require some means to break open cells and access their DNA. This can be difficult to do in the field, although some technologies are beginning to address
this limitation$^{46}$. Another limitation is that these technologies cannot distinguish between live and dead pathogens; DNA from dead pathogens remains detectable in water for days before it degrades$^{76}$, which means that a positive result is not a perfect indicator of water quality. Lastly, while the use of isothermal amplification allows for detection of trace amounts of pathogen DNA, the process of amplification obscures the pathogen’s original concentration and hampers precise quantification. Several existing strategies use mathematical models to infer pathogen concentration from final fluorescent signal strength$^{37,75}$, but precise quantification will likely require sophisticated peripheral equipment or an array of tests with built-in thresholds.

CHEMICAL CONTAMINANT DETECTION

Some of the most significant threats to our water supply are chemical contaminants—molecules that are deleterious to human health when consumed at dangerous levels. Many of these compounds occur naturally in soil, while others enter the water supply from industrial pollution, agricultural runoff, or deficient utility infrastructure. Sensors for chemical contaminants do not require an amplification step and therefore work more similarly to natural biosensors than pathogen biosensors do. There are two steps to detecting a chemical contaminant: the biosensor first recognizes its target chemical, which in turn initiates production of a reporter that generates a detectable signal. Currently, significant progress has been made in developing biosensors to detect arsenic$^{79}$ and fluoride$^{28}$, two of the WHO’s highest-priority chemical water contaminants$^{16}$. Here, we discuss recent progress towards using biosensors for chemical sensing with a focus on arsenic and fluoride, comparing these tools to existing field-deployable methods (Table 1).

Arsenic

Arsenic contamination of groundwater is typically caused by the leaching of naturally occurring arsenous compounds from the surrounding soil$^{77}$. Consumption of arsenic-contaminated water is associated with lesions, cardiovascular and pulmonary disease, and cancer in humans$^{78}$. Current field-deployable methods for arsenic detection utilize a colorimetric chemical test strip to semiquantitatively detect as low as 5 ppb arsenic within a few minutes$^{20}$. However, these tests require significant technical skill from the user and produce toxic byproducts, such as arsine gas.

Published biosensors for arsenic are generally whole-cell sensors that are controlled by an arsenic-responsive protein$^{79}$ (Fig. 3a). In the absence of arsenic, the protein binds to the biosensor DNA, stopping the reporter from being produced. Arsenic binds to the protein and causes it to change shape, releasing the biosensor DNA and allowing production of the reporter. Previous arsenic biosensors have used fluorescent, colorimetric, and luminescent outputs, with reported detection as low as single-digit parts per billion$^{79}$. This offers presence/absence results for arsenic concentrations below the 10 ppb WHO guideline for arsenic in drinking water$^{16}$, though these sensors have yet to be extensively validated in field conditions$^{79}$.

Fluoride

Fluoride leaches into groundwater from naturally occurring soil minerals and can also be introduced by agricultural runoff or the precipitation of fluoride-containing industrial ash in rainwater$^{80}$. Chronic consumption of fluoride-contaminated water causes dental and skeletal fluorosis, which manifests as discolored teeth, weakened bones, seizures, and stunted growth$^{80}$. Current field-deployable methods for fluoride detection utilize either
photometric analytical equipment to semi-quantitatively measure a colorimetric reaction, or a quantitative fluoride-sensing electrode. While both of these methods can safely detect down to 0.1 ppm fluoride, they require expensive supplementary equipment for their use, precluding widespread deployment.

A recently developed biosensor for fluoride uses a naturally occurring RNA regulator, called a riboswitch, in a cell-free system (Fig. 3b). In the absence of fluoride, the riboswitch folds into a structure that stops the reporter from being produced. When present, fluoride ions bind to the riboswitch, causing it to fold into an alternate structure that permits production of the reporter. This fluoride biosensor can be paired to both fluorescent and colorimetric outputs, with reported detection as low as one part per million. During preliminary field testing, this sensor provided presence/absence results in environmental samples below the 1.5 ppm WHO guideline for fluoride in drinking water.

### Emerging contaminants

Biosensors also have the potential to detect emerging contaminants beyond arsenic and fluoride, including metals, agricultural products, and pharmaceutical and personal care products (PPCPs), such as antibiotics and cosmetics. Both whole-cell and cell-free biosensors have previously been used to detect metals by utilizing natural or engineered proteins; sensors have been reported for cadmium, lead, mercury, arsenic, copper, zinc, nickel, and cobalt, with sensitivities ranging from low parts per million to parts per billion. Sensors for atrazine, a toxic herbicide, have also been developed by encoding a natural metabolic pathway for atrazine’s conversion to cyanuric acid, which can be detected with a known protein sensor. Furthermore, new cell-free approaches can detect a range of PPCPs, including multiple families of antibiotics and benzalkonium chloride. The ability to detect such a wide range of targets underscores the potential of biosensors as modular chemical sensing platforms, paving the way for rapid sensor development and deployment to detect new and emerging contaminants of concern.

### Outstanding challenges for chemical contaminant detection

While many whole-cell sensors report WHO-relevant limits of detection, they are limited by the deployment and operational concerns that are characteristic of live cells. Additionally, many of these sensors are susceptible to false positives due to interference by other chemical contaminants and unintended reporter production. To overcome these challenges, substantial progress must be made in developing robust biocontainment strategies and methods to tune biosensor sensitivity and specificity. While cell-free biosensors partially resolve some of these problems by virtue of being non-living, easily tunable systems, it is still difficult to completely predict how tuning certain parameters influences a sensor’s function. Further development of cell-free sensors must therefore focus on identifying the factors that contribute to maximal sensor and reporter function in cell-free systems and optimizing them for biosensing.

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**Table 1.** Comparison between commonly used point-of-use methods for contaminant detection (top) and biosensors (bottom).

| Target          | Method                        | Result type | Time to results | Test cost | Equipment cost | Skills required | Number of steps | Limit of detection | Output type |
|-----------------|-------------------------------|-------------|-----------------|-----------|----------------|-----------------|------------------|-------------------|-------------|
| **Coliforms**   | **Coliform Growth Test**      | Semi-quantitative | Hours           | $         | N/A            | Filtration 4   | 1 cfu/100 mL     | Colorimetric      |
| (Minimal)       |                               |             |                 |           |                |                 |                  |                   |
| Coliforms       | **Coliform Growth Test**      | Quantitative | Days            | $         | $$$            | Filtration 6   | 1 cfu/100 mL     | Colorimetric      |
| (Complex)       |                               |             |                 |           |                |                 |                  |                   |
| Coliforms       | **Tryptophan-Like Fluorescence** | Presence/absence | Minutes        | $         | $$$            | Filtration 3   | 10 cfu/100 mL    | Fluorescent       |
| Coliforms       | **Growth Assay**              | Presence/absence | Hours           | $         | N/A            | Filtration 3   | 1 cfu/100 mL     | Cell growth       |
| Coliforms       | **Hydrogen Sulfide**          | Presence/absence | Days           | $         | N/A            | Filtration 3   | 1 cfu/100 mL     | Colorimetric      |
| Arsenic         | **Chemical Test Strip**       | Semi-quantitative | Minutes       | $         | N/A            | Dilution 10   | 5–200 ppb        | Colorimetric b    |
| Fluoride        | **Fluoride-Sensing Electrode**| Quantitative | Minutes         | $         | $$$            | Dilution 5   | 0.1 ppm          | Numerical         |
| Fluoride        | **Complexone Method**         | Semi-quantitative | Minutes       | $         | $$$            | Dilution 3   | 0.1 ppm          | Colorimetric      |
| Coliforms       | **Loop-LAMP**                 | Semi-quantitative | Hours        | $         | $             | Filtration 4   | 20 cfu/100 mL    | Fluorescent       |
| Coliforms       | **SHERLOCK**                  | Semi-quantitative | Hours        | $         | N/A            | Filtration 4   | Attomolar c      | Colorimetric      |
| Coliforms       | **DETECTR**                   | Semi-quantitative | Hours        | $         | $             | Filtration 4   | Attomolar c      | Fluorescent       |
| Coliforms       | **RNA-Based Sensor**          | Semi-quantitative | Hours         | $         | N/A            | Filtration 4   | Nanomolar c      | Colorimetric      |
| Arsenic         | **Whole-Cell Protein Biosensor** | Presence/absence | Hours    | $         | N/A            | N/A           | 1 ppb            | Colorimetric, fluorescent |
| Fluoride        | **Cell-Free RNA Biosensor**   | Presence/absence | Hours        | $         | N/A            | N/A           | 1 ppb            | Colorimetric, fluorescent |

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* a $ < 1 USD, $$ < 10 USD, $$$ > 10 USD.
* b Generates toxic arsenic gas during operation.
* c As these methods detect DNA in the sample, their limits of detection are measured in DNA concentration rather than cfu.
DISCUSSION

Infrequent monitoring of a narrow range of contaminants has created significant gaps in our current understanding of water quality\(^1\)\(^-\)\(^3\) and therefore water insecurity.\(^87\)\(^-\)\(^89\) Synthetic biology has the potential to fill these knowledge gaps by offering simple, field-deployable tools to report on individual water supplies or serve as pre-screening tools to be used with existing gold-standard methods to provide the large-scale, high-resolution data needed to track progress towards development goals. While there are existing field-deployable tools, they are limited by the need for the technical expertise, supplemental equipment, or dangerous chemical reagents required for their use (Table 1). The potential for biosensors to decrease cost and improve ease-of-use for such diagnostics relative to current methods would enable more frequent measurements across wider and more diverse regions, producing water quality data that are more comprehensive and specific than currently available. There is significant promise for this to become a reality—current biosensor formats are accessible to an untrained user, and recent cost estimates suggest that their production can be scaled for global use. Freeze-dried, cell-free reactions can currently be manufactured for a few cents per sensor, with even lower costs possible for whole-cell biosensors.\(^4\) Moving forward, these costs could decrease by as much as one order of magnitude, further facilitating mass deployment.

This potential is counterbalanced by several existing barriers to rapid biosensor design and deployment. For example, we are currently limited to harvesting sensor parts from nature, rather than designing them from scratch. With the rise of unnatural contaminants such as synthetic antibiotics, pesticides, and other harmful industrial compounds, we may lack the tools to detect some emerging targets. While we are currently on the cusp of engineering entirely synthetic proteins\(^91\) and RNAs\(^92\) to address this need, the technologies to do so are still in their infancy, and it will be some time before they can be applied to targeted contaminant detection.

There is also significant work to be done in developing validated field deployment strategies. Although many biosensors can be freeze-dried for transport and long-term storage,\(^25\)\(^,\)\(^28\)\(^,\)\(^40\)\(^,\)\(^43\)\(^,\)\(^44\) this has not yet been explored in the context of tools to enable their use in real-world settings. Of particular concern is the potential for other compounds present in environmental samples to interfere with biosensor components, or for organic matter to chelate contaminants and mask their presence. Thus, a major next step for biosensor development is to characterize these potential inhibitory effects and devise strategies to make biosensors robust against them.\(^83\)

Fortunately, there have already been several successes using these biosensors in complex samples. For example, cell-free biosensors have been used in the laboratory to detect fecal contamination of unprocessed water samples, including a test for robustness with raw sewage.\(^90\) Additionally, a cell-free fluoride biosensor was capable of detecting environmental fluoride in unprocessed water samples onsite.\(^88\) While these preliminary findings suggest that cell-free biosensors can be robust to a wide range of potential contaminants in complex water samples, more exhaustive testing that considers the effect that common contaminants and other compounds found at target sites is needed. We must also develop comprehensive packaging and usage guidelines that accommodate both the needs and abilities of diverse users.

From a logistical perspective, the lengthy validation and regulatory approval processes for certifying biosensors may delay their potential impact. Furthermore, meeting the manufacturing demands for global deployment will require funding and production capabilities beyond the reach of academic labs. Enabling individuals to more easily monitor their own water quality could also reveal unwelcome information, and may raise some potential societal, ethical, data protection, and regulatory concerns. These issues will require the careful consideration and cooperation of diverse stakeholders to ensure that these technologies are used for the maximal public good. Because of
this, widespread implementation of these technologies will require interdisciplinary collaboration across the synthetic biology and water, sanitation, and hygiene (WASH) communities to foster the use of biological design to advance large-scale humanitarian goals.

As our ability to build biological systems improves, we can begin constructing more sophisticated systems from a wider array of biological parts. Recent work has demonstrated that biosensors can do more than merely produce a single reporter output in response to a target. Indeed, networks of interacting genes can be coupled to form “molecular computers” that take input signals from a biosensor and calculate an appropriate response. For example, a genetic system could be engineered to simultaneously detect multiple targets and produce an output that reports the identity and concentration of each target. Furthermore, new sample calibration strategies are being developed to circumvent biosensors’ intrinsic limitations and enable field-deployable sample quantification. As we continue harvesting parts from nature and clarifying biological design principles, we expect to see an improvement in the sensitivity and specificity of biosensors for an expanding list of detectable targets.

Using biosensors to generate spatiotemporal water quality data will enable more efficient resource allocation by showing exactly when and where interventions are necessary. Not only will such diagnostics provide important population-level information, but they have the potential to usher in the ability to simply and inexpensively assess water quality so that even untrained individuals can personally test the safety of their water. As such, advances in synthetic biology could facilitate global water quality monitoring by producing actionable contaminant data, guiding the development of efficacious policies and programs, and informing choices about the water we consume.

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