De Novo Design of the ArsR Regulated P_{ars} Promoter Enables a Highly Sensitive Whole-Cell Biosensor for Arsenic Contamination

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ABSTRACT: Whole-cell biosensors for arsenic contamination are typically designed based on natural bacterial sensing systems, which are often limited by their poor performance for precisely tuning the genetic response to environmental stimuli. Promoter design remains one of the most important approaches to address such issues. Here, we use the arsenic-responsive ArsR-P_{ars} regulation system from *Escherichia coli* MG1655 as the sensing element and coupled gfp or lacZ as the reporter gene to construct the genetic circuit for characterizing the refactored promoters. We first analyzed the ArsR binding site and a library of RNA polymerase binding sites to mine potential promoter sequences. A set of tightly regulated P_{ars} promoters by ArsR was designed by placing the ArsR binding sites into the promoter’s core region, and a novel promoter with maximal repression efficiency and optimal fold change was obtained. The fluorescence sensor P_{lacV}-P_{arsOC2} constructed with the optimized P_{arsOC2} promoter showed a fold change of up to 63.80-fold (with green fluorescence visible to the naked eye) at 9.38 ppb arsenic, and the limit of detection was as low as 0.24 ppb. Further, the optimized colorimetric sensor P_{lacV}-P_{arsOC2}-lacZ with a linear response between 0 and 5 ppb was used to perform colorimetric reactions in 24-well plates combined with a smartphone application for the quantification of the arsenic level in groundwater. This study offers a new approach to improve the performance of bacterial sensing promoters and will facilitate the on-site application of arsenic whole-cell biosensors.

Arsenic contamination of the atmosphere, water, and soil has become a worldwide health issue.1,2 It can enter the human body through inhalation, drinking, and even eating, and prolonged exposure to high levels of arsenic can cause serious damage, such as visible skin lesions, peripheral neuropathy, cardiovascular disease, diabetes, and renal system damage, such as visible skin lesions, peripheral neuropathy, and cancer (IARC classification).3–5 Consequently, the International Agency for Research on Cancer (IARC) classifies arsenic compounds as Group I carcinogens. To assess arsenic pollution and forestall further arsenic exposure, there is an urgent need to develop a rapid and reliable method to determine arsenic.

At present, whole-cell biosensors have attracted the attention of scientists because they are self-renewable and tolerant to harsh environments.6–9 Compared with traditional high-end analytical techniques, bacterial biosensors are cost-effective, easy to integrate, portable, and easily applied for high-throughput testing.9,10 The development of sensitive whole-cell sensors requires sophisticated sensing and signal transduction elements. Nevertheless, whole-cell sensors based on natural sensing systems have certain shortcomings, such as high leakage, low induced fold change, and poor sensitivity.11–13 Recent advances in synthetic biology have provided many methods for improving the regulation of the genetic response to achieve highly sensitive sensors, such as genetic circuit configurations,14,15 transcriptional promoter engineering,16,17 translational efficiency tuning,18–21 posttranslational protein degradation control,22–24 and output signal amplifiers.22,23 Promoters are the first gate for target gene expression and fundamental elements of the genetic circuit, and selected promoters with excellent performance remain one of the essential considerations for whole-cell biosensor design. The classical prokaryotic inducible promoter (e.g., P_{tet}, P_{lac}, and P_{ara}) comprises two core elements: the RNA polymerase binding sites (−10 and −35 sites) that determine promoter activity and the transcription factor binding sites (TFBSs) that control gene expression. Therefore, promoter engineering usually targets these two core elements for investigation.16,17,26 Taking the arsenic-regulated P_{ara} promoter as an example, previous studies on promoter engineering have focused on directed evolution (high-throughput screening of optimal promoters),13 mutating RNA polymerase binding sites (to enhance or attenuate promoter activity),12 adding an addi-
The straight lines with dots at both ends indicate that the refactored encoding multiple TFBS (e.g., LacO and TetO) into a single emerging, which is generally done by embedding TFBS of (Figure 1A) has a constitutive promoter $P_{\text{ars}}$ sensitively and specifically recognizes arsenic. The sensor circuit analysis, design, and optimization, which can be developed for promoters controlled by repression. 33 Specifical promoter characteristics still have a strong promoter activity. 34,35 Such work provides theoretical support for the de novo design of tightly regulated promoters controlled by transcription factors with long footprint sites. 32 Shortening the length of TFBS will reduce its affinity for transcription factors and then alter the dynamic behavior of the genetic circuit. 16,26 For transcription-factor-regulated prokaryotic $\sigma^+\text{O}$ promoters, deploying TFBS at different positions in the promoter will alter the dynamic behavior of the genetic circuit. A consensus on obtaining maximum repression efficiency has been developed for promoters controlled by repression transcription factors (such as LacI/$P_{\text{lac}}$ and TetR/$P_{\text{tet}}$). 16,29 The repression efficiency is most robust when the position of the TFBS is located between $−10$ and $−35$ sites (approx. 17 bp spacer). 16,26,29–31 However, transcription factors with long footprint sites are difficult to design (such as $P_{\text{ars}}$ up to 33 bp long). 32 Shortening the length of TFBS will reduce its affinity for transcription factors and then alter the dynamic behavior of the sensor. Consequently, there has been no research on the de novo design of tightly regulated $P_{\text{ars}}$ promoters to date, thus limiting the further development of arsenic whole-cell sensors. Recent studies predict that promoter sequences are diverse, and up to 60% of random sequences require only single-base mutations to be promoters. 33 Specific sequences without prominent promoter characteristics still have a strong promoter activity. 34,35 Such work provides theoretical support for the de novo design of tightly regulated promoters controlled by transcription factors with long footprint sites.

To address these challenges, we selected the *Escherichia coli* MG1655 $arsRBC$ operon as the starting element for the arsenic sensor circuit analysis, design, and optimization, which can sensitively and specifically recognize arsenic. The sensor circuit (Figure 1A) has a constitutive promoter $P_{\text{dtg}}$ that drives constant expression of the arsenic receptor ArsR. ArsR binds to the ABS to repress $P_{\text{ars}}$ promoter transcription in the absence of arsenic, which dissociates from the ABS in the presence of arsenic and triggers the reporter gene expression. We collected a library of RNA polymerase binding sites (i.e., $−10$ site sequences and $−35$ site sequences) and aligned them to natural ABS sequences to explore potential promoter sequences (Figure 1B). Based on the aligned results, we analyzed the potential promoter model and selected the optimal promoter model for the de novo design of promoters based on the principle of maximum overlap between the promoter core region and ABS (Figure 1C). After optimization, two promoters ($P_{\text{arsOB4}}$ and $P_{\text{arsOC2}}$) with low leakage and high fold changes were selected, and their applicability was confirmed with *gfp* and *lacZ* as reporter genes, respectively. Ultimately, the low-leakage and highly sensitive $P_{\text{lac}}$-$P_{\text{arsOB4}}$-*lacZ* sensor was used to perform colorimetric reactions in a 24-well transparent plate and combined with a color recognition application of a smartphone to analyze the arsenic content of groundwater samples. This study provided a promising approach for the design of tight regulation promoters and a platform for the rapid detection of arsenic.

### Materials and Methods

**Strains, Growth Conditions, and Reagents.** Plasmid construction and biosensor characterization were performed in the *E. coli* DH5α strain. Essential primers and detailed sequences of target genes used in this study are summarized in Tables S1 and S2. Bacteria were cultured in a lysogeny broth (LB) medium supplemented with 30 $\mu$g/mL tetracycline antibiotics. Solid media were prepared by supplementation with 15 g/L agar. For bacterial culture, the engineered strains were inoculated from a single colony on freshly streaked plates to 5 mL of LB in sterile 15 mL universal tubes and were incubated overnight at 37 °C with shaking (200 rpm). Unless otherwise noted, data were collected 6 h after the addition of the inducer in all characterization tests. 5-Bromo-4-chloro-3-indolyl $\beta$-D-galactopyranoside (X-gal) was purchased from TransGen Biotech. Sodium arsenic (NaAsO$_2$) and other chemicals used were analytical grade and purchased from Sigma-Aldrich. The different concentrations of NaAsO$_2$ used were incubated overnight at 37 °C with shaking (200 rpm).

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**Figure 1.** Schematic showing the de novo design of the arsenic-responsive tightly regulated $P_{\text{ars}}$ promoter. (A) Genetic circuit configurations of an arsenic-responsive sensor module coupled to a *gfp* reporter. The dashed arrow indicates the dissociation of ArsR-$\text{As}^{3+}$ from ABS; the three $T$-shape structures indicate the terminators. (B) Alignment of a library containing $−10$ and $−35$ site sequences with ABS sequences; potential $−10$ and $−35$ sites are shown in different colors. (C) Analysis, design, and optimization of tightly regulated $P_{\text{ars}}$ promoters. The $−35$ site sequence of the $P_{\text{arsWT}}$ promoter is shown in bold. Sequences of ABS are underlined. The potential $−10$ or $−35$ sites are shown as yellowish ($P_{\text{arsR}}$), greenish ($P_{\text{arsM}}$), and blueish ($P_{\text{arsL}}$) in the three refactored promoters, respectively. The refactored $−35$ and $−10$ sites and mutated sequences in ABS are marked in red. The straight lines with dots at both ends indicate that the refactored $−35$ sites (TATTACA) are sequentially located at six different positions.
in this study were all converted to the actual arsenic content for graphical purposes (in ppb).

**Plasmid Circuit Construction.** Standard molecular manipulations were used to construct the plasmids by using pPROBE-TT \(^{15}\) (pPROBE-TT was a gift from Steven Lindow, Addgene plasmid #37822) as a skeleton. The plasmid carries the gfp (mut) gene, referred to as gfp, derived from *Aequorea victoria*. Its product is a green fluorescent protein with a long half-life. The arsR gene with the constitutive promoter P\(_{I109}\) and the ribosome binding site (RBS) B0030 was amplified from the J109-ParsD-ABS-2 plasmid and inserted into the HindIII and SacI sites of the sensor detection platform pPROBE-TT plasmid using a recombination reaction to generate the p-TT-P\(_{I109}\)-30arsR plasmid (Figure S1). Primer extension polymerase chain reaction (PCR) was used to obtain the wild-type P\(_{\text{ars}}\) promoter and variant promoters with a refactored ~35 site at varying locations. The PCR product was cloned into the SacI and EcoRI sites of the p-TT-P\(_{I109}\)-30arsR plasmid to generate the p-TT-P\(_{I109}\)-30arsR-P\(_{\text{arsXX}}\) (XX means a different name) plasmids (Figure S2), which were named P\(_{I109}\)-P\(_{\text{arsXX}}\). To change the density of the receptor protein ArsR, the P\(_{\text{lacV}}\) Promoter was used to drive ArsR expression. The P\(_{\text{arsXX}}\) gene with the mutation was amplified from the p-TT-P\(_{I109}\)-30arsR-P\(_{\text{arsXX}}\) plasmid by PCR and then ligated with EcoRI and SacI restriction-enzyme-digested p-TT-lacVarsR \(^{12}\) to generate p-TT-P\(_{\text{lacV}}\)-arsR-P\(_{\text{arsXX}}\) plasmids, which were named P\(_{\text{lacV}}\)-P\(_{\text{arsXX}}\). For the construction of the sensor with lacZ as the reporter, lacZ with the same RBS as gfp was PCR-amplified from the wild-type *E. coli* MG1655 genome and then ligated with EcoRI and Hap1 restriction-enzyme-digested P\(_{\text{lacV}}\)-P\(_{\text{arsXX}}\) to generate P\(_{\text{lacV}}\)-P\(_{\text{arsXX}}\)-lacZ. The plasmids used in this study are summarized in Table S3. Genewiz Inc. (Suzhou, China) conducted the oligonucleotide primer synthesis and plasmid sequencing.

**Characterization and Data Analysis of the Fluorescent Biosensor.** For fluorescent sensor characterization, the overnight cultures were diluted 50-fold into a fresh LB medium. Then the diluted cultures were loaded into 24-well deep-well plates (Canvic, China) and induced with 100 \(\mu\)L of various concentrations of NaAsO\(_2\) to a final volume of 2 mL per well. After incubation for 6 h at 37 °C, culture samples of 1 mL were centrifuged for 1 min at 13,400g, and the supernatant was decanted. The bacterial pellet was suspended in 1 mL of phosphate-buffered saline (1× PBS, with 1 mg/mL kanamycin to stop the synthesis of GFP), and 200 \(\mu\)L of this suspension was transferred into a clear-bottom 96-well black plate (Fluorotrac 200; Greiner, Germany) to measure the cell growth and GFP expression. The OD\(_{600}\) (absorbance at 600 nm) and fluorescence (480 nm for excitation, 510 nm for emission, sensitivity = 60%) were read by a microplate reader (Synergy H1 multimode plate reader, BioTek). The PBS averaged fluorescence backgrounds \((n = 3, \text{OD}_{600}\) and fluorescence) were determined from wells loaded with 1× PBS and were subtracted from the readings of other wells. The fluorescence/OD\(_{600}\) (Fluo/OD\(_{600}\)) for a sample culture was determined after subtracting the averaged \((n \geq 3)\) counterpart of the negative control cultures (GFP-free) at the same time, and the fold change was calculated as follows:

\[
\text{Fold change} = \frac{\text{Fluo/OD}_{600}(+\text{inducer})}{\text{Fluo/OD}_{600}(-\text{inducer})}
\]

For fold change data, the Fluo/OD\(_{600}\) value obtained with inducer NaAsO\(_2\) was divided by the Fluo/OD\(_{600}\) value obtained without inducers. Unless otherwise stated, all fluorescence data were obtained as above. Dose–response curves were fitted using a nonlinear regression model with the Hill slope (log(agonist) vs response — variable slope (four parameters)). All data analyses were performed on GraphPad Prism 7 (GraphPad Software). The limit of detection (LOD) was calculated based on the formula LOD = limit of background (LOB) \(+ 1.645 \times SD(\text{background})\) \(\times \sqrt{n}\), where LOB = mean(Blank) + 1.645 \(\times SD(\text{blank})\). The definition of this equation is based on the fact that the output signal is the concentration of the analyte, so we need to convert the corresponding fluorescence signal to the concentration of arsenic based on the linear equation.

To visualize the expression of GFP, the same induction experiment was performed independently \((n = 1)\). Cell pellets held in 2 mL tubes were obtained as described above and photographed under daylight by a Samsung Galaxy S21 cell phone. Meanwhile, the bacterial precipitates were suspended in 1 mL of 1× PBS with 1 mg/mL kanamycin to stop translation, and 200 \(\mu\)L of the bacterial suspension was loaded into a 96-well black plate for fluorescence imaging \((\lambda\text{ex} = 475 \text{ nm}, \lambda\text{em} = 520 \text{ nm})\) using a small-animal imaging system (Night OWL II LB 983 NCI100, Berthold, Germany).

**Characterization of the Colorimetric Biosensor.** The growth conditions for the engineered sensors are described above. For colorimetric biosensor characterization, the overnight cultures were diluted 50-fold into a fresh LB medium. The chromogenic substrate of 20 mg/L X-gal was supplied to the cell cultures before incubation (unless otherwise indicated, 200 \(\mu\)g/L X-gal was used as the final concentration). Ten microliters of various concentrations of arsenic was added to 96-well clear flat-bottom plates, 190 \(\mu\)L of the culture was added to each well (the induction concentration of arsenic was 0, 1.17, 2.34, 4.69, 9.38, and 18.75 ppb), and the plate was placed in an orbital shaker (200 rpm, 30 °C). Cultures incubated for 6 h were measured for OD\(_{600}\) (absorbance at 650 nm) using a BioTek Synergy H1 microplate reader and photographed using a Samsung Galaxy S21 cell phone.

**Colorimetric Analysis of Real Water Samples.** A 0.22 \(\mu\)m filter was used to filter the groundwater samples (taken from Mosuwan, Xinjiang) to remove impurities and bacteria. Subsequently, the arsenic content was detected by atomic fluorescence spectrometer (the highest arsenic content in all samples was 20.7 ppb and is referred to as G-Sample 1; all other contents were below 20 ppb). To test the recovery of the sensor for different arsenic levels, G-Sample 2 and G-Sample 3 with arsenic concentrations of 50.7 and 80.7 ppb were prepared by titrating NaAsO\(_2\) standard samples into G-Sample 1. G-Sample 1, G-Sample 2, and G-Sample 3 were diluted at dilution factors of 1, 2, 3, 4, 5, and 10, and 350 \(\mu\)L of diluted samples and different concentrations of NaAsO\(_2\) reference samples (with 0, 2, 4, 6, 8, and 10 ppb As) were pre-spiked to clear flat-bottom 24-well plates. Subsequently, 350 \(\mu\)L aliquots of the P\(_{\text{lacV}}\)-P\(_{\text{arsOC}}\)-lacZ biosensor cultures (with 400 \(\mu\)g/mL X-gal and 60 \(\mu\)g/mL tetracycline antibiotics) in the logarithmic phase (OD \(\approx 0.4\)) were transferred to each well, and the plate was placed on an orbital shaker (200 rpm, 30 °C). Thus, the final concentrations of X-gal and tetracycline were 200 and 30 \(\mu\)g/mL, respectively, and the arsenic concentrations of the actual and reference samples were further diluted twofold. After 5 h of incubation, colorimetric results of bacterial...
cultures were acquired according to the method described in the previous section.

Data Acquisition and Processing. The colorimetric photographs of the 24-well plates were loaded into a mobile app (Color Recognizer) and analyzed for the color intensity of each well. The mobile app can analyze the different parameters (such as RGB, CMYK, and LAB) that represented the color intensity. RGB values vary from the range of pure white (255, 255, 255) to pure black (0, 0, 0). As the blue intensity increases, the value of the R channel gradually decreases, so we use 255 minus R (255 − R) to represent the intensity of blue and establish a linear relationship with the arsenic concentration. The final concentrations of groundwater samples were calculated based on the equation and its dilution factor.

RESULTS AND DISCUSSION

Refactoring the \(P_{ars}\) Promoter. To obtain the maximum repression efficiency of ArsR for the \(P_{ars}\) promoter, the possibility of the maximum overlap between the core region (between the −10 and −35 sites) of the promoter and the ABS was explored. First, the effective length of the ABS was determined by using ABS with variable lengths to reduce the leakage of the arsenic whole-cell biosensors. The results showed that when the ABS sequence was between 24 and 33 bp, the repression efficiency increased with length (Figure S3). Then, we aligned the ABS sequences with the RNA polymerase binding site library (comprising the eighteen −35 sites and thirty-six −10 sites) (Table S4) using a SnapGene software. The alignment results showed that in addition to the wild-type −35 site, there were potential −10 and −35 sites in ABS, which could be generated by one base mutation at most (Figure 1B and Figure S4). Based on these results, we designed three promoter models, \(P_{ars1}\), \(P_{ars2}\), and \(P_{ars3}\) (Figure 1C: Analysis), which provide different options for the de novo design of promoters. \(P_{ars1}\) with the maximal repression efficiency in theory, was selected for further study because its operator site ABS is located in the core region of the promoter and completely overlaps with the −10 site.

To test this hypothesis, we refactored a −35 site (TTTACA) by site-directed mutagenesis in different positions upstream of the potential −10 site and eliminated the prototype −10 site of the wild-type promoter (Figure 1C: Design). For \(P_{arsO1}\) to \(P_{arsO9}\), different capital letters A to H, implying that the refactored −35 sites (TTTACA) are sequentially located at different positions. Although the typical promoter has a spacing of 17 bp between the −10 and −35 regions, we assessed the impact of spacer variations 16 to 23 bp (corresponding to \(P_{arsO1}\) to \(P_{arsO10}\)) on the promoter activity and fold change in the presence of 300 ppb arsenic. Among the nine variants in the first round, the promoter \(P_{arsO1}\) with 17 bp spacers was highly active (in the absence and presence of 300 ppb arsenic) with strong fluorescence compared to the unmodified \(P_{arsWT}\) promoter. Due to the high leakage of \(P_{arsO1}\), it exhibited a lower fold change. However, in the presence of 300 ppb arsenic, the signal output of the other promoters was lower than that of the wild-type promoter; in the absence of arsenic, their fluorescence was comparable to that of the negative control (Figure 2A). To avoid generating infinite fold changes, fold changes were not calculated for promoters whose leakage levels were close to the negative control. Nonetheless, \(P_{arsOE1}\) and \(P_{arsO1}\) possess a longer operator site, these may be more suitable for ArsR with a longer footprint site. We selected \(P_{arsOB1}\), \(P_{arsOC1}\), and \(P_{arsOE1}\) to generate \(P_{arsOB2}\), \(P_{arsOC2}\), and \(P_{arsOE2}\) by refactoring different −35 sites at the same position to further regulate the promoter activity; different numbers mean that different −35 sites are refactored at the same position (Table S3). In the second round of optimization, \(P_{arsOB2}\) obtained the highest fold change, and \(P_{arsOC2}\) obtained a simultaneous improvement in leakage and signal output, thus increasing the fold change to 51X (Figure 2B). These results suggest that improvements in repression efficiency can be achieved by mining the potential promoter sequences to refactor novel promoters. Meanwhile, the performance of modified promoters can continue to be optimized by engineering promoter designs that are not shown here. The current sequences with promoter features were gradually expanded, and a series of new promoters were redesigned. Besides, AI (artificial intelligence)-based promoter redesign and prediction are emerging, thus providing powerful tools for the mining of tightly regulated promoters. This study gives a new perspective on promoter engineering that could reconstruct artificial promoters with maximal repression efficiency targeted at the specific TFBS.

Performance of the Fluorescent Biosensor. To examine the performance of the refactored promoter variants, we selected \(P_{J109}P_{arsWT}\), \(P_{J109}P_{arsOB4}\), and \(P_{J109}P_{arsOC2}\) for
further analysis. The expression of arsenic receptor ArsR was driven by the P_{J109} promoter in these sensors. We investigated the dose−response curves, fold change, and cell phone images after the induction of arsenic at various concentrations. Compared with the wild-type promoter, both P_{arsOB4} and P_{arsOC2} showed a lower leaky expression in the absence of arsenic, especially P_{arsOB4}, which was reduced 4.5-fold, whereas P_{arsOC2} showed a 2-fold fluorescence increase in the presence of 300 ppb arsenic (Figure 3A). This finding was likely associated with the configuration in which the ABS was located in the core region of the promoter, which reduced the probability of RNA polymerase binding to the promoter in the absence of arsenic and therefore effectively reduced the basal expression. In the presence of 300 ppb of arsenic, the fold change of P_{J109}−P_{arsOB4} and P_{J109}−P_{arsOC2} sensors was 25.27, 73.21, and 50.54, respectively (Figure 3B); moreover, fold changes of 1.12, 1.20, and 1.25 occurred at the lowest induction concentration of 0.29 ppb (Table S5), respectively, indicating that the refactored promoter improved the sensor sensitivity to some extent. Besides, we found that the LOD was somewhat increased for the refactored promoter (Table S5). We speculate that it is due to the increased repression efficiency of ArsR to the refactored promoter. Nonetheless, the difference in LOD between the refactored promoter and the wild-type promoter was less than 0.1 ppb, which is negligible for the sensor development.

Previous studies have shown that the density of arsenic receptor ArsR can modulate the sensitivity and fold change of the sensor.42 To show that this improvement in the basal expression, signal output, and sensitivity is robust, we replaced P_{J109} with a more potent P_{lacV} promoter to express the arsenic receptor ArsR. These results showed that the dose−response curves of the different sensors using the P_{lacV} promoter to drive ArsR expression were consistent with those using the P_{J109} promoter (Figure 3A,C). Sensors with the P_{lacV} promoter possessed a lower leaky expression and hence more remarkable fold changes at the same arsenic level than those using the P_{J109} promoter (Figure 3B,D). However, the fold-change of the P_{lacV}−P_{arsOB4} and P_{lacV}−P_{arsOC2} sensors was diminished at low concentrations (Table S5). Interestingly, P_{lacV}−P_{arsOC2} effectively reduced the sensor’s basal expression while maintaining a higher signal output, thus achieving a bidirectional improve-

Figure 3. Characterization of the various arsenic-responsive promoters (P_{arsWT}, circles; P_{arsOB4}, squares; P_{arsOC2}, triangle) within a fluorescent whole-cell biosensor. (A−D) Dose−response curves and fold change of different promoters when ArsR expression was driven by P_{J109} (A and B) or P_{lacV} (C and D). Error bars show the standard deviation (n = 3). (E) Images of cell cultures show pellets under daylight (left) or liquids in a fluorescent imaging system (right).
ment in background noise and signal output. Ultimately, a fold change of 183.52 was produced over the control when induced with 300 ppb arsenic. These fluorescent biosensors were semilog linear ($R^2$ from 0.9868 to 0.9934) at the range of 2.34 to 150 ppb arsenic (Figure S5). The LOD of the optimal sensor $P_{\text{lacV}}-P_{\text{arsOC2}}$ was as low as 0.24 ppb, which is 41.67-fold lower than the safety level of 10 ppb for drinking water defined by the World Health Organization. Therefore, it would fully meet the requirements for real applications.

To visualize the difference in fluorescence of different sensors, we photographed the cell cultures after arsenic induction at different concentrations with and without the fluorescence imaging system (Figure 3E). As shown in Figure 3E (left), the sensor constructed from the $P_{\text{arsOC2}}$ promoter had green fluorescence visible to the naked eye when the arsenic concentration was higher than 9.38 ppb, while the sensors constructed from the $P_{\text{arsOWT}}$ and $P_{\text{arsOB4}}$ promoters required higher concentrations of arsenic to have visible fluorescence. With the increase of arsenic concentration, sensors whose ArsR expression was driven by the $P_{\text{lacV}}$ promoter exhibited a sensitive color transition (from blue to green, to yellow, and finally to red) than $P_{\text{J109}}$ (Figure 3E, right). This phenomenon may be due to their lower leaky expression and higher fold changes at the same arsenic concentration.

Since the overexpression of ArsR may affect the specificity of the arsenic sensor, we performed a specificity assay for $P_{\text{J109}}$, $P_{\text{arsOC2}}$ and $P_{\text{lacV}}$, with a wide range of metals that could be potential water contaminants. Remarkably, the expression of the arsenic receptor ArsR using two different promoters showed high specificity for arsenic and no reactivity to the other 10 metal species, even antimony (a homolog of arsenic) (Figure S6A,B). In addition, we tested the performance of the $P_{\text{lacV}}-P_{\text{arsOC2}}$ sensor at different temperatures. There were a lower signal output and background at 30 °C compared to at 37 °C (Figure S6C), while the fold change was higher at 30 °C (Figure S6D).

**Arsenic Colorimetric Analysis with the $\beta$-Galactosidase Biosensor.** While it is essential to quantify the level of arsenic, it is more important to quickly diagnose if the arsenic level is over the safe limit. Colorimetric output can be semiquantitatively obtained by observing the color reaction. We replaced gfp with lacZ as the reporter gene (its product $\beta$-galactosidase can cleave the colorless substrate X-gal to blue) (Figure 4A). We first assessed the performance of different promoters at various concentrations of arsenic using 200 μg/mL X-gal as the final concentration of the substrate (Figure 4B). $P_{\text{arsOB4}}$ showed a clear blue color among all promoters tested even in the absence of arsenic, which could easily cause false-positive results in the actual test. $P_{\text{arsOB4}}$ had an ultralow background with no arsenic, but the blue response was not as pronounced as that of $P_{\text{arsOC2}}$ when the arsenic concentration increases. We observed that the $P_{\text{lacV}}-P_{\text{arsOC2}}$-lacZ sensor showed a more pronounced blue color at 1.17 ppb than the control; however, when the arsenic content exceeded 4.69 ppb, the blue color intensity would gradually saturate and become indistinguishable. The results of OD$_{650}$ also indicated that the design of a tightly regulated promoter improved the performance of the colorimetric sensor (Figure S7A).

To analyze the effect of X-gal concentration on sensor performance, we tested the $P_{\text{lacV}}-P_{\text{arsOB4}}$-lacZ response to arsenic in different X-gal concentrations. As expected, the blue response of the sensor became more pronounced as the X-gal concentration increased while maintaining a low background (Figure 4C). Thus, the X-gal dosage can effectively adjust the sensor sensitivity to meet different detection requirements. In particular, $P_{\text{lacV}}-P_{\text{arsOB4}}$-lacZ showed a stable OD$_{650}$ for arsenic above 4.69 ppb at 300 and 400 μg/mL X-gal (Figure S7B), which is consistent with the results of the colorimetric reaction. Considering the effect of bacterial growth density on OD$_{650}$, the selection of a suitable color measurement method is important for the quantification of arsenic. For the subsequent experiments, we selected appropriate parameters that allowed the detection of arsenic down to 0–5 ppb while maintaining distinct blue differences. This will allow for a quick assessment of whether arsenic levels exceed the safety level for drinking water defined by the World Health Organization.

**Easy-to-Interpret Colorimetric Array for Arsenic Monitoring.** To enable smartphone portable analysis, we sought to design a colorimetric array based on $P_{\text{lacV}}-P_{\text{arsOC2}}$-lacZ sensors to analyze the actual samples. As shown in Figure 5A, after induction of the sensor by different samples, its color changes were recorded by a smartphone camera, and the different RGB values were read out directly with the help of a smartphone application (Color Recognizer). A functional relationship between the arsenic concentration and color

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**Figure 4.** Characterization of various arsenic-responsive promoters within a colorimetric biosensor. (A) Schematic showing the arsenic-responsive sensor module ($P_{\text{lacV}}-P_{\text{arsZ}}$-lacZ) coupled to a lacZ reporter. (B) Images of the colorimetric tests of the three different promoters under various induction levels of arsenic. (C) Images showing the dose responses of the $P_{\text{lacV}}-P_{\text{arsB4}}$-lacZ sensor to various induction levels of arsenic at different X-gal substrate concentrations.
intensity was subsequently established. In principle, the color intensity of the 24-well plates was directly proportional to the concentration of arsenic. However, when the arsenic content exceeded 5 ppb, the blue color gradually saturated and became indistinguishable, which was not conducive to the accurate assessment of arsenic. Therefore, the actual samples needed to be diluted with different dilution factors (2, 4, 6, 8, 10, and 20) first (Figure 5B). Linear relationships between the intensity of blue (255 - R) and the arsenic concentration of reference samples (0 to 5 ppb) were established by the PlacV-ParsOC2-lacZ sensor (Figure 5C). This sensor showed good linearity over the concentration range of 0 to 5 ppb (R² = 0.9928), while the LOD was as low as 0.39 ppb, which is 25.64-fold lower than the safety level of 10 ppb for drinking water. For groundwater samples, their color intensity would be within the linear range of the reference samples at the appropriate dilution factor (Figure 5D). The final concentrations of groundwater samples were calculated under different dilution factors based on the equation 255 - R = 13.65(As)/(dilution factor) + 85.6, and the maximum concentration was used as the final result. The heat map shows the mean of three biological replicates.

CONCLUSIONS

In summary, to address the issues of high background and low induction fold of arsenic whole-cell biosensors, we proposed a de novo promoter design approach for transcription-factor-regulated promoters with long footprint sites. The design approach is time-saving compared to error-prone PCR and high-throughput screening methods. We mined potential promoters within the ABS and developed improved arsenic whole-cell biosensors with low leakage and high signal output. The fold change of the optimized sensor was increased from the initial 25.27 × to 187.52 × in the presence of 300 ppb arsenic. Compared to previous studies, we constructed PlacV-ParsOC2 and PlacV-ParsOC2-lacZ sensors that showed a superior detection limit and linear range (Table S7). Excess ArsR may be detrimental to the detection limit and sensitivity of the sensor, while our refactored promoter greatly improves the repression efficiency of ArsR to Pars and avoids the use of excess ArsR to control the basal expression. Hence, this approach enables the repression of background noise and increase of fold change upon signal emission without impairing.
the detection limit and sensitivity. In addition, excess ArsR increases the nonspecificity of the sensor. Therefore, the de novo designed promoter provides a new alternative approach for the development of arsenic sensors with a low detection limit, high signal-to-noise ratio, and high specificity. The use of different reporter genes to construct the sensor showed different characteristics, with the gfp reporter requiring a higher expression to obtain a visible output signal though having a wider linear range, while the lacZ reporter facilitates the development of a device-free detection platform due to the chromogenic response though with a narrower linear range and requiring a lower promoter leakage. Low leakage and high signal output promoters allow for a more flexible selection of reporter genes and are the basis for obtaining highly sensitive sensors. Finally, a highly sensitive colorimetric sensor array was built to accurately assess arsenic contaminant levels from 0 to 100 ppb and had a low detection limit of 0.39 ppb. Although the current sensor requires 5 h of incubation to generate a sufficiently strong colorimetric output for visualization, this is mainly associated with the restricted diffusion and transport of substrates across the cell membrane. This can be optimized to generate a much faster response by cell-free systems or lysing cells. \(^{34,35}\) The color difference arising from the colorimetric array assay can be quickly captured by a smartphone without sophisticated equipment, thus facilitating its potential use as an easy-to-integrate and low-cost environmental monitoring tool in the field. This study provides a new approach for designing tightly regulated promoters with an aim for developing highly sensitive whole-cell biosensors. It will facilitate the on-site application of whole-cell biosensors for precise arsenic detection.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.analchem.2c00055.

Schematic diagram of the p-TT-P\(^{-}r30\)-arsR plasmid construct (Figure S1); schematic diagram of the p-TT-P\(^{-}r30\)-arsR-ParsXX (named P\(^{-}J109\)-ParsXX) plasmid construct (Figure S2); validation of the effective length of the ArsR binding site (Figure S3); results showing the ABS sequence alignment with the RNA polymerase binding site (Figure S4); different fluorescent sensors showing a linear response to arsenic (Figure S5); sensor specificity and temperature effect on the sensor performance (Figure S6); absorbance of diverse colorimetric sensors at 650 nm monitored across various arsenic concentrations (Figure S7); primers used in this study (Table S1); sequences of the genetic constructs used in this study (Table S2); plasmid constructs used in this study (Table S3); RNAP binding site library containing eighteen −35 sites and thirty-six −10 sites (Table S4); best fits for the characterized responses of the arsenic sensors with diverse promoters (Table S5); accuracy and reliability of the constructed biosensors for real samples (Table S6); and comparison of previously reported arsenic sensors with this study (Table S7) (PDF).

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Author Contributions

Sheng-Yan Chen, Bang-CE Ye, and Baojun Wang conceived the project. Sheng-Yan Chen performed all research with the help of Renjie Li. Yan Zhang performed the characterization experiments of the fluorescent biosensor. Sheng-Yan Chen, Bang-CE Ye, and Baojun Wang analyzed data and wrote the manuscript with comments from all co-authors.

**Notes**

The authors declare no competing financial interest.

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