Identification of stress-responsive transcription factors with protein-bound Escherichia coli genomic DNA libraries

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Abstract
Bacteria promoters along with operators are crucial elements in the control of gene expression in microbes in response to environmental stress changes. A genome-wide promoter DNA regulatory library is in demand to be developed for a microbe reporter method to monitor the existence of any given environmental stress substance. In this study, we utilized Escherichia coli (E. coli) as a model system for the preparation of both cell lysates and genomic DNA fragments. Through enriching protein-bound DNA fragments to construct luciferase reporter libraries, we found that, of 280 clones collected and sequenced, 131 clones contained either the promoter-35 and -10 conservative sequences and/or an operator transcription factor binding sites (TFBS) region. To demonstrate the functionality of the identified clones, five of 131 clones containing LexA binding sequence have been demonstrated to be induced in response to mitomycin C treatment. To evaluate our libraries as a functional screening library, 80 randomly picked up clones were cultured and treated with and without MMC, where two clones were shown to have greater than twofold induction. In addition, two arsenite-responsive clones were identified from 90 clones, one having the well-known ArsR and another having the osmotically inducible lipoprotein (OsmE1). The newly discovered osmE1 has been quantitatively validated to be induced by arsenite treatment with real-time PCR in a dose response and time course manner. This enriching protein-bound DNA luciferase reporter libraries and functional screening facilitate the identification of stress-responsive transcriptional factors in microbes. We developed functional libraries containing E. coli genomic-wide protein-bound DNA as enhancers/operators to regulate downstream luciferase in response to stress.

Keywords: Genomic DNA libraries, Luciferase assay, Transcription factors, ArsR, LexA

Introduction
Microbes are highly adaptable to environmental toxic stress such as heavy metals, pesticides, and polychlorinated biphenyls (PCBs) (Chowdhury et al. 2018; Caine 2012). The adaptation to changes in their environment is controlled by the induction or repression of gene expression (Balleza et al. 2009; Cases et al. 2003). Association or dissociation of a transcription factor (TF) to its DNA binding site is a critical step in the initiation of the transcription of its target gene (Fernandez-Lopez et al. 2015; Rogers et al. 2015). It is vital to identify and characterize genes involved in the response to an environmental stress from the entire genome. This facilitates both the understanding of the mechanisms of gene regulation as well as the identification of the key regulatory elements during environmental adaptation in the host.

Environmental genomic toxic stresses such as certain types of chemical reagents and UV irradiation can cause changes in gene expression and cellular metabolism of microbe (Foster 2007). The distinguishing feature of these genes is the presence within the promoter region of a binding sequence for transcriptional repressors, such as LexA (Butala et al. 2009) and ArsR (Chen et al.
LexA repressor normally is bound to its binding sites, repressing transcriptional expression. In response to any DNA damage, the LexA repressor undergoes dissociation from its binding sequences and activate DNA repair genes (Butala et al. 2009). ArsR is a regulatory protein that controls the expression of the genes involved in arsenical resistance via interaction with the arsenic-responsive operon (Wu and Rosen 1993). Upon arsenic binding, the protein dissociates from the promoter, subsequently activating relevant gene expression (Shi et al. 1994). Nevertheless, many toxic substances and their corresponding genes are not well characterized due to lack of simpler and more efficient methods.

Traditionally, transcription factor binding sites (TFBSs), are identified through approaches such as DNase I footprinting (Brenowitz et al. 1989) and electromobility shift assays (Hellman and Fried 2007), which are limited to the interactions between TFs and single targets. Recently, multiple TFs have been experimentally investigated using the systematic evolution of ligands by exponential enrichment (SELEX) (Ishihama et al. 2016) and chromatin immunoprecipitation with microarray (ChIP-chip) or by sequencing (ChIP-seq) (Galagan et al. 2013). Both ChIP-seq and genomic SELEX require the knowledge of stress-corresponding TFs prior to analysis, with time-consuming and tedious procedures. Recently many microbial genomes have been completely sequenced due to advances in the high-throughput genome sequencing, leading to computational methods to identify transcription factor binding sites (TFBSs) in these microbial genomes. However, computational method cannot identify the location and function of promoter region of a transcription factor (Inukai et al. 2017).

Identification of a specific target’s responsive TFBS is very helpful for the development of bacteria biosensors in detecting a chemical substance and its toxicity. However, most of the current bacteria biosensors utilize the existing substrate-induced promoter and operator regions, such as arsenite detection biosensor with GFP (Zaslawer et al. 2006) and luciferase (Chen et al. 2019) as reporters. For a new and potential toxin without knowing its associated TFs, no global reporter method has been developed yet to identify and determine the associated TFs or TFBSs that are required in the regulation of gene expression.

In this study, we present an innovative high-throughput approach to screen and discover TFBSs in response to a stress substance directly without any prior genome information. Functional libraries have been constructed with enriched protein-bound genomic DNA fragments as enhancer and operators extracted from E. coli DH5α, along with downstream luciferase reporter to facilitate functional screening. 74% of the sequenced clones were predicted to contain regulatory TFBS with BPROM program from Softberry (Solovyev and Salamov 2011). From 80 randomly screened clones upon mitomycin C (MMC) treatment, two clones were found to be induced and confirmed to contain LexA binding sites. Furthermore, when screening another 90 clones with arsenite treatment, two clones were shown to be induced and have ArsR binding site, corresponding to arsR and osmE1. In the paper we newly discovered osmE1 gene, containing an arsR binding motif. The gene expression of osmE1 was further validated by real-time RT-PCR in a dose–response and time course of arsenite-mediated induction.

**Materials and methods**

**Preparation of cell lysate proteins**

One mL of E. coli DH5α culture was centrifuged at 10,000 g for 1 min and the pellet was resuspended in 300 μL of lysis buffer (10 mM Tris–HCl, pH 8.0, 0.1 M NaCl, 1 mM ethylenediamine tetraacetic acid (EDTA), and 0.1% (w/v) polyethylene glycol octylphenyl ether (Triton X-100)). 7.5 μL of a freshly prepared lysozyme solution (10 mg/mL in 10 mM Tris–HCl, pH 8.0, final concentration=0.25 mg/mL) was added and mixed by tapping the tube gently, and the lysis mixture was incubated for 10–20 min at room temperature. After centrifugation, the supernatant was used for filter-binding selection.

**Preparation of Genomic DNA fragments**

DH5α cells were collected through centrifugation, resuspended in 200 μL lysis buffer (10 mM Tris–HCl, pH 8.0, 1 mM EDTA, 0.5% SDS) and treated with 20 μg/mL proteinase K for 2 h at 55 °C. Genomic DNA was extracted with phenol and chloroform. The genomic DNA was digested with MnlI, 5′...CCTC(N)7...3′, which recognizes four base pairs and generates one nucleotide protruding end at the 3′ terminus, for 1 h at 37 °C. The genomic DNA fragments were subsequently purified with MinElute Reaction Cleanup Kit (QIAGEN, Hilden, Germany).

**Filter-binding selection of protein-bound DNA fragments**

Five μL cell lysate (2–10 μg) was mixed with 15 μL 2X Binding buffer (40 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES), pH 7.6, 20 mM ammonium sulfate, 2 mM dithiothreitol (DTT), 20 mM KCl, and 0.4% Tween-20). MnlI-digested 5 μL genomic DNA and 5 μL ddH2O in a PCR tube. After incubation at room temperature for 30 min, we loaded 30 μL binding mixture onto a prewashed filter assay column and incubated on ice for 20 min. The column is a nitrocellulose-based filter system, which can bind proteins and protein-DNA complex. After four times washing with Filter washing buffer to remove free DNA oligos, the bound DNA fragments...
were eluted with elution buffer (0.5% SDS). The eluted DNA fragments were subsequently used for generating libraries.

**Construction of genomic libraries**

The eluted protein-bound DNA fragments were ligated with adaptors. *MnlI* digested fragments may have multiple nucleotide possibilities at the 3′-terminus. Two basic sequences for making adaptors were selected to avoid cross hybridization with *E. coli* genome, 5′-ATG GAT AGG TCG GTGA3′ or 5′-GAC GCA CCT TGA GGC3′. The double strand adaptors were designed and synthesized to match all possible fragments generated by *MnlI*-digestion (Fig. 1) and two DNA oligos were annealed to form the double strand adaptors with different protruding ends respectively. The oligos were designed and synthesized: (F1T 5′-TCA CCG ACC TAT CCAT-T3′, F2T 5′-GCC TCA AGG TGC GTC-T3′, F1A 5′-TCA CCG ACC TAT CCAT-A3′, F2A 5′-GCC TCA AGG TGC GTC-A3′, F1C 5′-TCACCGACCTATCCAT-C3′, F2C 5′-GCCTCAAGG TGCGT-C3′, F1G 5′-TCACCGACCTATCCAT-G3′, and F2G 5′-GCCTCAAGG TGCGT-G3′). F1 and F2 were annealed with R1S: 5′-ATG GAT AGG TCG GTGA3′ or R2S 5′-GAC GCA CCT TGA GGC3′ accordingly to form eight adaptors: 5AA, 5AG, 5AC, 5AT, 3AA, 3AG, 3AC, and 3AT (Table 1). After ligation of adaptors with DNA fragments, 16 combinations were amplified by 10 PCR cycles with a forward primer introduced with XbaI sequence and a reverse primer with *HindIII* sequence. The amplified products were digested with XbaI and *HindIII* and cloned into pACYC-Luc vector, which was modified in our previous publication (Chen et al. 2019), originally derived from pACYC184 (New England Biolabs, Ipswich, MA, USA) to generate 16 libraries (AA, AT, AC, AG; TA, TT, TC, TG; CA, CT, CC, CG; GA, GT, GC, GG) listed.

![Diagram](image-url)
were treated with or without MMC, or sodium arsenite at 37 °C until the optical density (O.D.) reached 0.5. Cells were cultured for an additional 4 h with vigorous shaking. The overnight culture was diluted 1:50 in a new 96 well deep plate in the corresponding wells of 96 well deep plate with 25 μg/mL chloramphenicol, and incubated for 12–16 h at 37 °C in the corresponding wells of 96 well deep plate with MMC treatment in a time course results.

Real-time RT-PCR
A single DH5α colony with the OsmE1 promoter containing clone was cultured overnight and diluted at 1:50 with LB before with MMC treatment in a time course and dose response manner (the detail see in result). Total RNA was prepared with Monarch Total RNA Miniprep

Table 1 Sixteen genome libraries generated from combination of eight adaptors sequences randomly digested by MnlI restriction enzyme

| No | Genome library names | Adaptor combination | 5′ Adaptor name | 5′ Adaptor sequence | 3′ Adaptor name | 3′ Adaptor sequence |
|----|----------------------|---------------------|-----------------|---------------------|-----------------|---------------------|
| 1  | AA                   | SAA-3AA             | SAA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ A-TACCTATCCAGCCACT 5′ (F1A) | 3AA               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 2  | AT                   | SAA-3AT             | STA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ T-TACCTATCCAGCCACT 5′ (F1T) | 3AT               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 3  | AG                   | SAA-3AG             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ G-TACCTATCCAGCCACT 5′ (F1G) | 3AG               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 4  | AC                   | SAA-3AC             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F2C) | 3AC               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 5  | TA                   | STA-3AA             | STA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F1A) | 3AA               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 6  | TT                   | STA-3AT             | STA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ A-TACCTATCCAGCCACT 5′ (F1T) | 3AT               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 7  | TG                   | STA-3AG             | STA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ G-TACCTATCCAGCCACT 5′ (F1G) | 3AG               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 8  | TC                   | STA-3AC             | STA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F2C) | 3AC               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 9  | GA                   | SGA-3AA             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ G-TACCTATCCAGCCACT 5′ (F1G) | 3AA               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 10 | GT                   | SGA-3AT             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F2C) | 3AT               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 11 | GG                   | SGA-3AG             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ G-TACCTATCCAGCCACT 5′ (F2C) | 3AG               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 12 | GC                   | SGA-3AC             | SGA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F2C) | 3AC               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 13 | CA                   | SCA-3AA             | SCA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F1A) | 3AA               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 14 | CT                   | SCA-3AT             | SCA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ A-TACCTATCCAGCCACT 5′ (F1T) | 3AT               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 15 | CG                   | SCA-3AG             | SCA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ G-TACCTATCCAGCCACT 5′ (F2C) | 3AG               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |
| 16 | CC                   | SCA-3AC             | SCA             | 5′ ATGGATAGGGTGCGTGGA 3′ (R1S) | 3′ C-TACCTATCCAGCCACT 5′ (F2C) | 3AC               | 5′ GAGGCACCTTGAGGC 3′ (R2S) |

* Each adaptor was abbreviated with 5A (5′ adaptor A), 5T (5′ adaptor T), 5G (3′ adaptor G), etc. 5′ adaptors are annealed with shared sequence R1S while 3′ adaptors were annealed with shared sequence R2S. Each 5′ adaptor was combined with each 3′ adaptor to create a total of 16 genome libraries.

in Table 1. After transformation, the clones (colonies) were selected on ampicillin plates, and plasmid DNAs from 280 clones were subsequently either prepared and sequenced, or directly conducted induction luciferase screening assay.

Luciferase assay
For the screening assay, 80–90 individual colonies were picked and inoculated in 600 μL LB media supplied with 25 μg/mL chloramphenicol, and incubated for 12–16 h at 37 °C in the corresponding wells of 96 well deep plate with vigorous shaking. The overnight culture was diluted 1:50 in a new 96 well deep plate with pre-warm and fresh-prepared 600 μL LB media supplied with chloramphenicol. The diluted cells were cultured for an additional 4 h at 37 °C until the optical density (O.D.) reached 0.5. Cells were treated with or without MMC, or sodium arsenite (AsIII) at 37 °C. 20 μL of induced culture was mixed with 50 μL luciferase substrate, and the luciferase activity was measured with Veritas Microplate Luminometer (Tuner Biosystems, Sunnyvale, CA, USA). For an individual clone assay, a plasmid was transformed into DH5α, and a single colony was inoculated in 2 mL LB media 25 μg/mL chloramphenicol for 12–16 h at 37 °C in an individual tube, with the rest of steps being the same as the screening assay and treatment following the description in the results.
Kit (New England Biolabs, Ipswich, MA, USA) with DNAase treatment to remove residual DNA. Integrity of RNA was assessed by electrophoresis. RNA concentration was determined with Qubit™ RNA BR Assay Kit (Thermo Fisher Scientific, Waltham, MA, USA) in Qubit 2.0 Fluorometer according to manufacturer’s instructions. Extracted RNA (400 ng) was reverse transcribed to cDNA with AMV Reverse Transcriptase (Life Science Advanced Technology, St Petersburg, FL, USA). The primers for the target gene OsmE1 and three reference genes were designed with vector NTI (Thermo Fisher Scientific, Waltham, MA, USA) and using Primer-BLAST (NCBI, USA) and synthesized at IDT (Integrated DNA Technologies, Coralville, IA, USA). The primer specificity was confirmed by 2% agarose gel electrophoresis.

SYBR green-based real-time PCR was performed with ABI PRISM 7000 sequence detection system. 20 μl of PCR reaction was prepared based on Q5 DNA polymerase system (New England Biolabs, Ipswich, MA, USA) with 1X SYBR Green, 1X ROX dye (Roche, Basel, Switzerland), 1 μM forward and reverse primer. The amount of cDNA used in each qPCR reaction was: 1 μL for target gene osme1, 1 μl for reference genes, gryA and mGOD, and 0.6 μL of 1:100 diluted cDNA for 16S rRNA. These were pre-determined by testing serial dilutions of cDNA samples to achieve the threshold cycle (Ct) values of the three reference genes similar to that of the target gene. We ran the PCR reaction at 50 °C for 2 min and 98 °C for 5 min, followed by 40 cycles at 98 °C for 15 s, 55 °C for 30 s, and 72 °C for 30 s. A dissociation stage was then performed as follows: 95 °C for 15 s, 60 °C for 20 s, and 95 °C for 15 s. All samples were run in duplicate, and the mean Ct values for each trial were calculated. ΔCt was then calculated as the difference between target gene and the geometric mean of three reference genes. ΔΔCt was obtained by normalizing the ΔCt values of the treatments to the ΔCt value of the control without treatment. Finally, relative target gene expression values were calculated with $2^{-ΔΔCt}$ (Livak and Schmittgen 2001).

**Results**

This screening libraries consist of enriching protein-bound genomic DNA fragments and downstream luciferase reporters. The DNA fragments were generated based on the protein/DNA complex formation and protein/DNA complex separation (Fig. 1). To construct these libraries, *E. coli* genomic DNA was digested with a restriction enzyme *MnlI* that recognizes non-palindromic nucleotide sequence 5n/DNA comple (Kriukiene et al. 2005), each fragment with one protruding nucleotide at 3cleot with four possibilities: A G, C, and T. If there were DNA fragments containing promoters or TFBS and their corresponding DNA binding proteins in DH5α lysates such as Sigma 70 or TFs, protein/DNA complexes were formed. The enriched protein-bound DNA fragments were obtained and utilized to generate 16 libraries to contain all of promoter and operator regions of genomic DNA. Additionally, these libraries also are functional libraries with luciferase reporter gene. Once the TFs bind on the regulatory DNA regions of libraries, release repressor, and initiate the transcription of luciferase gene. Through measurement of luciferase activities, the clones containing regulatory DNA in response to a treatment.

In order to evaluate libraries with useful TFBS information, approximately 560 clones were obtained from the transformation of these libraries. Of these clones, we selected 280 for sequencing and generated 178 sequences with promoter region sizes around 70–300 bp. First, we analyzed these sequences with a computational analysis of promoter regions and TFBSs. Prokaryotic transcription is performed by RNA polymerase that contains four catalytic subunits and a sigma regulatory subunit. Seven total distinct sigma factors bind a set of promoter sequences and different sigma factors binding sites. The conservative sequences can be found between -10 base pairs and -35 base pairs upstream of the transcription start site in the promoter regions and TFBSs where located upstream of the promoter region acting as an enhancer or a repressor. Using the computer program BPROM (Solovyev and Salamov 2011), we found only 54 sequences having a-10 and -35 bp sigma factor. 71 have at least one TFBS and sigma factors and 6 only contain TFBS without sigma factor. A total of 131 out of 178 clones contain either promoter sequences or TFBSs, or both (shown in Table 2). The sequencing analysis showed that some promoter sequences displayed multiple TFBSs, such as elbB containing RpoD18, LexA, GLP, ArcA, FimZ, and ArgR, while some had only one TFBS such as dtpD containing only LexA. This study revealed a total of 35 unique TFs. Each TF was predicted by BPROM based on its consensus binding element, but the binding sequence on a specific promoter region may be different, which is the reason why we obtained much more the binding sequences than the number of unique TFs (Table 3).

In order to conduct functionality of the predicted TFBS in reporter vector, we first chose lexA as our testing target since the LexA DNA binding site was recognized to appear more frequently than others, and was predicted to be located on several gene promoter sequences, including kdo, fimZ, dtpD, and ElbB. Furthermore, LexA is widely studied and is well known to be induced by environmental stress (Maslowska et al. 2019; Kreuzer 2013). Five clones containing LexA binding sites were selected for functional tests of MMC-mediated activation of LexA: clone 137 dtpD, clone 138 elbB, clone 152...
| Gene names | Clone ID | Library | Encoded function | Insert length (bp) | TFBS |
|-----------|----------|---------|------------------|-------------------|------|
| aor       | 141      | TC      | Aldehyde ferredoxin-ductase | 279               | Sigma70 |
| araH      | 28       | GA      | L-arabinose ABC transporter permease | 167              | Sigma70, DnaA |
| arcB      | 47       | AC      | Sensor histidine kinase | 258              | Sigma 70, Ihf |
| argB      | 80       | AC      | Acetylglutamate kinase | 225              | Sigma70, ArgR2, Ihf |
| ansB      | 65, 85, 108, 114 | AG, CA CT, TT | Arsenite/antimonite:H(+) antiporter | 320 | Sigma70 |
| bgIH      | 72       | AT      | Carbohydrate-specific outer membrane porin | 244 | Sigma70, PurR |
| bsl78     | 107      | CT      | DNA (cytosine-S-)-methyltransferase | 218 | Sigma 70 |
| cas1e     | 18       | GG      | Type I-E CRISPR-associated protein | 186 | Sigma70, RpoS17 |
| casA      | 37       | GG      | CRISPR system Cascade subunit | 158 | Sigma70, RpoS17 |
| chbC      | 73       | AT      | N,N-diacylchitobiose-specific PTS enzyme IIC | 158 | Fur, RpoD17 |
| chbg      | 70, 91   | AT      | Chitin disaccharide deacetylase | 157 | Sigma70 |
| clk_1237  | 177      | TA      | ADP-ribosylglycohydrolase family protein | 124 | Sigma70 |
| ctk       | 93       | CA      | DUF4297 domain-containing protein | 135 | Sigma70, RpoD15, metJ |
| ctpA3     | 144      | TC      | Carboxyl-terminal-processing peptidase 3, chloroplastic | 158 | Sigma70 |
| cydD      | 145      | TC      | Cysteine/glutathione ABC transporter permease/ATP-binding protein | 167 | Sigma70 |
| dgc       | 122      | TT      | Diguanylate cyclases | 105 | Sigma70, RpoD17 |
| dtpD      | 137      | TC      | Dipeptide permease | 213 | Sigma70, LexA |
| eamA      | 74       | AT      | Cysteine/O-acetylserine exporter EamA | 184 | Sigma70 |
| ecm18     | 150      | TG      | Class I SAM-dependent methyltransferase | 270 | Sigma70, Crp, SoxS |
| e22563    | 35       | GG      | Selenium-dependent molybdenum hydroxylase system protein | 133 | Sigma70 |
| egc82     | 33       | GG      | D-hexose-6-phosphate mutarotase | 139 | Sigma70 |
| elbB      | 138      | TC      | Isoprenoid biosynthesis gly-oxalase | 328 | Sigma70, Irg, RpoH2, LexA, ArcA, argR, GlpR |
| f0fl      | 171      | TA      | ATP synthase subunit delta | 134 | Sigma70 |
| fmZ       | 165      | TA      | Fimbriae biosynthesis transcriptional regulator | 96 | Sigma70, PurR, LexA1, LexA2, PurR |
| frdA      | 117      | TT      | Fumarate reductase flavoprotein subunit | 357 | Sigma70 |
| frsA      | 1, 81    | GA, AC  | Esterase | 193 | Sigma70 |
| ftr1      | 3, 4, 11, 12, 13, 14, 15, 21, 27, 57, 62, 95, 97, 110, 111, 112, 126, 154 | GA, GC, GT, AC, AG, CG, CT, TC, TG | Iron permease | 145 | Sigma70, RpoD16, MetJ |
| gaf       | 6, 24    | GA, TC  | GAF domain-containing protein | 262 | Ihf |
| ghtT      | 86       | CA      | Cationic dicarboxylase symporter family transporter | 263 | Sigma70, RpoD17, OmpR |
| gshB      | 59       | AG      | Glutathione synthase | 208 | Sigma70, Fis |
| Gene names  | Clone ID | Library | Encoded function                                                                 | Insert length (bp) | TFBS |
|------------|----------|---------|-----------------------------------------------------------------------------------|-------------------|------|
| hcaE       | 30       | GA      | 3-Phenylpropionate/cinnamic acid dioxygenase subunit alpha                          | 140               | Sigma70, DnaA |
| hemK       | 147      | TC      | Peptide chain release factor N(S)-glutamate                                        | 366               | Sigma70 |
| hflC       | 63       | AG      | Protease modulator                                                                 | 320               | Sigma 70 |
| hipA       | 42       | AA      | Type II toxin-antitoxin system serine/threonine protein kinase toxin                | 230               | Sigma70, RpoD17 |
| hjr        | 22       | GG      | Holliday junction resolvase                                                        | 432               | RpoD17 |
| hsdR       | 157      | TG      | Type I restriction-modification system endonuclease                                 | 278               | Sigma70 |
| lcsA       | 153      | TG      | Outer membrane protein lcsA autotransporter precursor                              | 196               | Sigma70, RpoD17, SoxS |
| lIIA       | 55       | AC      | PTS mannitol transporter subunit                                                    | 154               | Sigma70, lhf, ArgR2, RpoD17 |
| kch        | 53       | AC      | Voltage-gated potassium channel                                                   | 160               | Sigma70, MetJ, RpoH2 |
| kdo        | 170      | TA      | 3-Deoxy-manno-octulosonate cytidylyltransferase                                     | 119               | Sigma70, LexA, RpoD18, PurR |
| kup        | 155      | TG      | Low affinity potassium transporter                                                 | 154               | Sigma70 |
| lpfC       | 136, 178 | TC, TA  | Fimbrial biogenesis outer membrane usher protein                                   | 155               | Sigma70 |
| maaFP003_1916 | 31 | GA      | Si-specific NAD(P) (+) transhydrogenase                                            | 381               | Sigma70 |
| maeB       | 94, 116  | CA, TT  | NADP-dependent oxaloacetate-decarboxylating malate dehydrogenase                   | 121               | Sigma, RpoD17, ArgR |
| mcrB       | 39       | GG      | 5-Methylcytosine-specific restriction enzyme B                                      | 128               | Sigma70, RpoD17, Irp, Fnr, NagC |
| mhpR       | 127      | TC      | DNA-binding transcriptional activator                                              | 284               | Sigma70, RpoD17, ArgR, ArcA |
| mmmC       | 69       | AT      | 5-Methylaminomethyl-2-thiouridine biosynthesis bifunctional protein                | 182               | Sigma70 |
| motA       | 9        | GC      | Flagellar motor stator protein 98                                                 | 98                | FliA, MotAB, CheAW, CpxR |
| mshB       | 120      | TT      | Acidic protein                                                                     | 308               | Sigma70, SoxS |
| nmkF       | 20       | GG      | Chromosome partition protein                                                       | 123               | Sigma70, RpoD17, Irp, RpoH2, Fnr, NagC |
| narG       | 159      | TG      | Nitrate reductase subunit alpha                                                    | 154               | Sigma70, ArgR |
| narI       | 10       | GC      | Respiratory nitrate reductase subunit gamma 195                                    | 195               | Sigma70, Crp |
| nikC       | 45       | AA      | Nickel ABC transporter permease subunit                                            | 164               | Sigma70, RpoD16 |
| nrdD       | 175      | TA      | Anaerobic ribonucleoside-triphosphate reductase                                   | 350               | Sigma70 |
| nuoE       | 158      | TG      | NADH-quinone oxidoreductase subunit NuoE                                           | 222               | Sigma70, Crp |
| pntA       | 8, 50    | GA      | Si-specific NAD(P) transhydrogenase                                                | 415               | Sigma70 |
| rase       | 56       | AC      | 4-Hydroxybenzoate octaprenyltransfer                                              | 178               | Sigma, PurR |
| rayT       | 88       | CA      | REP-associated tyrosine transposase                                                | 248               | Sigma70 |
| Gene names | Clone ID | Library | Encoded function | Insert length (bp) | TFBS                  |
|-----------|----------|---------|------------------|--------------------|----------------------|
| rep       | 139      | TC      | ATP-dependent DNA helicase Rep | 185                | Sigma70              |
| rpoS      | 156      | TG      | RNA polymerase sigma factor  | 315                | Sigma70, Crp         |
| rppH      | 169      | TA      | RNA pyrophosphohydrolase     | 396                | Sigma70, NarP        |
| rrl       | 61       | AG      | 23S ribosomal RNA           | 153                | Crp, RpoD15          |
| sanA      | 67       | AT      | Outer membrane permeability protein | 193                | Sigma 70             |
| sdr       | 149      | TG      | Short-chain dehydrogenase   | 99                 | Sigma70              |
| secA      | 167      | TA      | Preprotein translocase subunit SecA | 161                | Sigma70, RpoD17, Ihf |
| sgr       | 132      | TC      | Helix-turn-helix domain-containing protein | 199                | Sigma70, ArgR2       |
| specF     | 90       | CA      | Ornithine decarboxylase SpeF | 128                | Sigma70              |
| spy       | 105      | CT      | ATP-independent periplasmic protein-refolding chaperone | 233                | Ihf, Fis, lrp,       |
| tesB      | 173      | TA      | Acyl-CoA thioesterase II    | 133                | Sigma70, RpoD16      |
| thiP      | 99       | CT      | Thiamine/thiamine pyrophosphate ABC transporter permease | 210                | Sigma70, OmpR        |
| tolC      | 115      | TT      | Outer membrane channel protein | 128                | ArgR                 |
| trpS/pgp  | 123      | TT      | Tryptophan-tRNA ligase Phosphoglycolate phosphatase | 115                | Sigma70              |
| tyrR      | 134      | TC      | Transcriptional regulator   | 147                | RpoD19, RpoD17, Crp, OmpR, MetR |
| ucpA      | 100      | CT      | SDR family oxidoreductase   | 133                | Sigma70, Fnr         |
| ugdH      | 121      | TT      | UDP-glucose 6-dehydrogenase | 294                | Sigma70, FlhCD, RpoH2, RpoD17, Fnr, lrp |
| uhpcC     | 16       | GG      | MFS transporter family glucose-6-phosphate receptor | 174                | Sigma70, ArgR        |
| wbbL      | 5, 29    | GA      | Beta-1,6-galactofuranosyltransferase | 200                | Sigma70              |
| wcAL      | 129      | TC      | Colanic acid biosynthesis glycosyltransferase | 290                | Sigma70              |
| weel      | 24, 75   | GT      | Beta-1,6-galactofuranosyltransferase | 217                | Sigma70, RpoD17, RpoD16,RpoD17,RpoD17 |
| wzc       | 109      | CT      | Tyrosine-protein kinase     | 179                | Sigma70              |
| ybaT      | 89       | CA      | Amino acid permease         | 171                | ArgR2, Ihf, ArcA     |
| ybjX      | 49       | AC      | DUF535 domain-containing protein YbjX | 145                | Sigma70              |
| ycbV      | 148      | TG      | Putative fimbrial-like adhesin protein | 110                | Sigma70, Crp, RpoD15, PhoB, RpoD17, lrp |
| ychE      | 131      | TC      | NAAT family transporter    | 258                | Sigma70, RpoD16, lrp, Sox5, TyrR, GipR, RpoD18 |
| ydhW      | 23, 25, 40 | GT      | Oxidoreductase             | 183                | Sigma70              |
| ydiV      | 52       | AC      | EAL domain-containing protein bacteria | 241                | Sigma70, Crp         |
| yeaW      | 101      | CT      | Carnitine monoxygenase subunit | 179                | Sigma 70             |
| yedE/fdhT | 48       | AC      | Selenium metabolism membrane protein | 110                | Sigma70              |
| yeeJ      | 119      | TT      | Inverse autotransporter adhesin | 182                | Sigma70              |
| yehH      | 135      | TC      | YehH family putative sulfite export transporter | 114                | Sigma70, OxyR, arcA, Fnr, RpoD18, TyrR, Fnr, DeoR, Ihf, ArgR2 |
(non-coding Pseudo gene), clone 170 *kdo* and clone 165 *fimZ*, which was previously reported to be regulated by LexA (Saini et al. 2009). These clone plasmids were transformed into DH5α, and inoculated and treated with 0, 0.2 and 0.5 μM MMC for 2, 4 and 16 h respectively (Fig. 2), and cell lysates were prepared for luciferase analysis. 2 h treatment did not show significant induction rate (Fig. 2a). Even though all of these clones showed the clear induction at 0.5 μg/mL MMC for 4 h treatment (Fig. 2b), the induction patterns showed a slight difference in terms of the condition for highest induction: clone 137 and clone 165 showed highest induction at 0.2 μg/mL for a 16 h treatment (Fig. 2c), while clone 138, clone 152 and clone 170 showed highest induction at 0.5 μg/mL for 4 h treatment (Fig. 2b). The results of the clones containing LexA binding sequences were confirmed to be induced by MMC with luciferase assays, since the LexA binding sequences in these clones come from different gene promoter regions, which may affect the responding pattern of MMC treatment.

To further demonstrate the feasibility of direct library screening without prior information, 80 clones were randomly selected from libraries. We chose treatment conditions of 0.5 μg/mL MMC for 4 h treatment (Fig. 2b), the induction patterns showed a slight difference in terms of the condition for highest induction: clone 137 and clone 165 showed highest induction at 0.2 μg/mL for a 16 h treatment (Fig. 2c), while clone 138, clone 152 and clone 170 showed highest induction at 0.5 μg/mL for 4 h treatment (Fig. 2b). The results of the clones containing LexA binding sequences were confirmed to be induced by MMC with luciferase assays, since the LexA binding sequences in these clones come from different gene promoter regions, which may affect the responding pattern of MMC treatment.

To validate our direct screening function of libraries, we utilized *arsR* as another screening target, which we have widely studied in our recent publication (Chen et al. 2019). Another 90 clones from libraries were cultured and treated with 5 μM arsenite for 2 h based our previous optimal conditions. Nine clones showing high luciferase activities (> 600 RLU) (Fig. 4a) were then selected and analyzed thoroughly with individual arsenite induction assay. Two clones, clone 12 and 68, were confirmed to have greater than twofolds induction. The plasmids were prepared from clone 12 and 68 and then subjected to sequencing. Through NCBI BLAST search, clone 12 revealed *osmE1* and clone 68 revealed *arsR*. Both clones were unable to be analyzed with the promoter prediction program BPROM as the program does not contain ArsR binding sequences, although Arsenite-mediated induction of ArsR is well-documented (Chen et al. 2017, 2019; Bose et al. 2006; Kostal et al. 2004). The *arsR* binding site on ArsR found in this study TTA AAT CAT ATG CGT TTT TGGTT was the identical to the published one (Xu et al. 1996). The potential ArsR binding site on *osmE1* was predicted to be GCTTGAAAAAGCGCCCAATG based on reported consensus sequence, tTGxxxx xx xxxxCAa (Busenlehner et al. 2003) shown in Fig. 5.

Since *osmE1* is not well studied and is newly discovered in our study, this gene induction by arsenite treatment needs to be further investigated. To analyze arsenite-mediated induction of *osmE1* gene expression,

| Gene names   | Clone ID | Library | Encoded function                          | Insert length (bp) | TFBS       |
|--------------|----------|---------|-------------------------------------------|--------------------|------------|
| *yfeX*       | 54       | AC      | Porphyrinogen peroxidase                   | 167                | Sigma70    |
| *yhdP*       | 46       | AA      | AsmA2 domain-containing protein            | 200                | Sigma70    |
| *yidR*       | 125      | TC      | DUF3748 domain-containing galacturonate catabolism protein | 174                | Sigma70    |
| *yihG*       | 151      | TG      | Putative acyltransferase                   | 186                | Sigma70    |
| *ymeE*       | 98       | CG      | Bestrophin family inner membrane protein   | 120                | Sigma70    |
| *ypfG*       | 64       | AG      | DUF1176 domain-containing protein          | 328                | Sigma70, RpoH3 |
| 26           | GT       | Hypothetical protein                       | 211                | Sigma70, ArgR2, Crp |
| 152          | TG       | Non-coding, Pseudo genes                   | 186                | Sigma70, Fis, Fnr, LexA |
| 161          | TG       | Unknown                                       | 626                | Sigma70    |

Table 2 (continued)
we utilized real time RT-PCR quantitative measurement in time and dose course. For dose response assays, DH5α cells were treated with 0, 0.04, 0.08, 0.16, 0.31, 0.63, 1.25, 2.5, 5, and 10 μM arsenite for 2 h. Total RNAs were prepared, and reverse transcribed to cDNA. SYBR Green PCR reactions were performed in duplicate, and the mean Ct values for each trial were calculated. As shown in Fig. 6a, the treatment with 2.5 μM of arsenite yielded the highest induction of osmE1 gene expression. Next, we examined the time-course response of OsmE1 gene expression to 2.5 μM arsenite for periods of 0, 15, 30, 60, and 120 min. The samples were collected at the indicated time points and quantification of osmE1 gene expression normalized using the references. The results revealed that the 120 min treatment yielded the highest induction, ninefolds, of osmE1 gene expression (Fig. 6b).

**Discussion**

Bacteria biosensors act as a new class of detectors to produce a detectable signal upon activation of a promoter reporter gene induced by specific stimuli, which have been used for monitoring environmental pollutants such as heavy metals or pesticides (Gutiérrez et al. 2015). The key component of whole-cell biosensors is
Fig. 3 Directly functional screening of mitomycin C-responsive clones. Eighty clones randomly selected from generated library, cultured them with 0.5 μM mitomycin C treatment for 4 h, and subject to luciferase analysis (a). Six clones with luciferase activities > 550 RLU were selected for induction assay with and without mitomycin C treatment (b).

Fig. 4 Directly functional screening of arsenite-responsive clones. Ninety clones from library were randomly selected, cultured with 5 μM arsenite treatment for 2 h, and subject to luciferase analysis (a). Nine clones with luciferase activities > 600 RLU were then selected for arsenite induction assay (b).
the reporter (Gui et al. 2017), consisting of a promoter/operator and a reporter gene. Therefore, it is crucial to find a responding promoter/operator in a high throughput method from surviving microbes in an environment containing a target pollutant. The current bacteria reporter biosensors are only for the known toxin substance-induced TFBS constructed reporter system, and cannot be used for discovering a TF and the associated TFBS for a novel substance.

This study presents a novel approach to enriching protein-bound genomic DNA fragments for the construction of luciferase libraries conducting directly functional screening to identify substance-responsive TFBS elements. This dramatically reduces time and labor in the screening of unknown TFBS elements in response to a potential toxin substance. It has been widely known that there are around 300 TFs and seven sigma factors in the E. coli genome (Pérez-Rueda and Collado-Vides 2000; Tripathi et al. 2014). Our protein bound enriched DNA libraries displayed 131 TFBS containing clones from screening 280 clones based on sequencing analysis and bacteria TFBS prediction software BPROM, and identified two well-studied ArsR (Chen et al. 2017, 2019; Bose et al. 2006; Kostal et al. 2004) and FimZ (Saini et al. 2009) among these TFs, which demonstrating our libraries are highly enriched with useful TFBS information. In addition, through luciferase assay, the same TF (such as LexA) on the promoter region with different binding sequences were shown to have various induction patterns, therefore the libraries can not only obtain a specific TF binding motif, but also provide multiple promoter associated binding sequences with different induction patterns, which may offer possibilities to develop more sensitive and selective stress substance screening system. Through direct functional screening, we were able to obtain MMC-responsive lexA clones and As-responsive arsR and osmE1 clones. These results showed that our functional libraries can be utilized to efficiently screen and discover the responsive clones under stress substance stimulation. Our library screening does not require the prior knowledge of the target microbial genome or any known transcription factor, therefore our libraries have great potential to be used for identifying a specific TF binding site of a given substance, and developing functional screening methods for unknown microbes with very limited physiological and genomic information.

Studies demonstrate that arsenite can mediate ArsR induction, which is well-documented in literature (Chen et al. 2017, 2019; Bose et al. 2006; Kostal et al. 2004). ArsR, belonging to the Smt/ArsR family, is a regulatory protein that controls the expression of the genes involved in arsenical resistance via interaction with the arsenic-responsive operon (Chen et al. 2017). Due to the abundant presence of ArsR binding sequences in microbial chromosomes, the alignment of these binding sequences via comparison and analysis leads to the identification of a binding consensus sequence (Saini et al. 2009). SmtB/ArS Binding sequences share a conserved 12-2-12 palindrome (Kostal et al. 2004). Our recent study indicated that among the inverted repeat, TC and GA are critical to ArsR binding (Chen et al. 2019). Interestingly, we found that OsmE1 is also a target capable of regulation.
by arsenite, although this has been shown in only one previous study (Patel 2005). This study reported that the identification of the arsenic binding-protein fractions with arsenic analysis revealed two low molecular weight proteins, which one of them being OsmE1. Cells under arsenate stress conditions could allow the expression of osmE1. Further studies need to determine how many genes are induced under arsenic stress, how they are regulated by arsenite, and what function they play in response to arsenic stress.

Our E. coli protein-bound DNA enriched functional library technology can easily be adapted to mammalian TFBS identification; however, mammalian transcriptional regulation is much more complicated than bacteria transcriptional regulation as there are more than 2000 TFs for mammals (Brivanlou and Darnell 2002). Luciferase-based screening may be time-consuming to assay individual clones. GFP reporter can replace luciferase reporter to construct libraries, so that the differentially expressed reporter genes can be easily identified through fluorescence-activated cell sorting (FACS) to sort the interesting population in response to a certain treatment. Our protein-bound enriched functional library technology has a wide application for TFBS identification of unknown transcriptional regulation in prokaryotic and eukaryotic system.

Acknowledgements
Not applicable.

Authors’ contributions
XL contributed to experimental design. XJ performed clone library screening. MY, YF, and YW contributed to data analysis. XL, MY, GS, and JG were involved with study design and overseeing the experiments. The manuscript was written by XL, and all authors commented on the manuscript. All authors read and approved the final manuscript.

Funding
This work was supported by the High-level Leading Talent Introduction Program of GDAS (2016GDASRC-0208) and the Science and Technology Planning Project of Guangzhou City (201707020021) to XL, National Natural Science Foundation of China (91851202) and the Science and Technology Project of Guangdong Province (2019B110205004) to MX.

Availability of data and materials
All data and materials are available.

Ethics approval and consent to participate
This article does not contain any studies with human participants and animals performed by any of the authors.

Consent for publication
All authors consent the publication.

Competing interests
The authors declare that they have no conflict of interest.

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Received: 10 October 2020 Accepted: 18 October 2020
Published online: 02 November 2020

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