Transcription Factor SrsR (YgfI) Is a Novel Regulator for the Stress-Response Genes in Stationary Phase in Escherichia coli K-12

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Abstract: Understanding the functional information of all genes and the biological mechanism based on the comprehensive genome regulation mechanism is an important task in life science. YgfI is an uncharacterized LysR family transcription factor in Escherichia coli. To identify the function of YgfI, the genomic SELEX (gSELEX) screening was performed for YgfI regulation targets on the E. coli genome. In addition, regulatory and phenotypic analyses were performed. A total of 10 loci on the E. coli genome were identified as the regulatory targets of YgfI with the YgfI binding activity. These predicted YgfI target genes were involved in biofilm formation, hydrogen peroxide resistance, and antibiotic resistance, many of which were expressed in the stationary phase. The TCAGATTTTG C sequence was identified as a YgfI box in in vitro gel shift assay and DNase-I footprinting assays. RT-qPCR analysis in vivo revealed that the expression of YgfI increased in the stationary phase. Physiological analyses suggested the participation of YgfI in biofilm formation and an increase in the tolerability against hydrogen peroxide. In summary, we propose to rename ygfI as srsR (a stress-response regulator in stationary phase).

Keywords: transcription factor; stationary phase; biofilm formation; hydrogen peroxide sensitivity; genomic SELEX (gSELEX); Escherichia coli

1. Introduction

The growth of bacteria in a natural environment is often limited; so, within these environments, they subsequently enter a stationary phase. The long-term survival of bacteria under stressful conditions depends on the establishment of various stress defense systems. For example, stationary phase adaptation of bacteria is accompanied by marked changes in their gene expression patterns [1–3]. Escherichia coli, as an example, contains a total of approximately 4500 protein-coding sequences on its genome [4,5]. The genes required for the response to various stresses and for survival in the stationary phase have been identified by comprehensive analyses using gene deletion or overexpression strains [6,7]. At present, however, even within the model bacterium E. coli, the function of one-fourth of the species is still unknown because most stress-response genes are not expressed within laboratory conditions. Accordingly, the full set of regulators involved in the expression of stress conditions during the stationary phase has not yet been identified. In the absence of knowledge of transcription factors (TFs) and conditions affecting the expression of regulatory functions, the ordinary in vivo approach is not useful in identifying regulatory targets of hitherto uncharacterized TFs, as TFs are not always expressed [8]. At present, the function of one-fifth of E. coli TFs remains unidentified [9].
To rapidly identify the regulation targets by uncharacterized TFs, we developed an improved system of genomic systematic evolution of ligands using exponential enrichment (SELEX) [10,11], and successfully identified regulation targets for several TFs [9,12]. This gSELEX screening system using the reaction in vitro between purified test TF proteins and genome DNA fragments is especially useful in identifying regulatory targets of hitherto uncharacterized TFs, of which their expression conditions are unknown. Using the gSELEX system, we identified the regulation targets and regulatory roles for some unknown Y-TFs, including YagI (renamed as XynR), YbaO (renamed as DecR), YbiH (renamed as CecR), YbjK (renamed as RcdA), YcdC (renamed as RutR), YcjZ (renamed as PgrR), YdeO (renamed as PhhR), YdcN (renamed as SutR), YdfH (renamed as RspR), YdhM (renamed as NemR), YeaM (renamed as NimR), YedW (renamed as HprR), YugP (renamed as Dan), YiaJ (renamed as PiaR), YiaU (renamed as CsuR), and YihW (renamed as CsqR) [9,13–17]. The gSELEX analysis can also lessen the difficulty of in vivo analysis because a set of regulatory proteins involved in the regulation of a single promoter competes to bind to the overlapping DNA sites [18]. Although difficulties exist, bacterial DNA-binding TFs bind to DNA and regulate nearby genes; thus, predicting regulation target promoters, genes, and operons is possible based on the location of recognition sequences by test TFs.

As part of our research strategy to understand the function of the comprehensive set of E. coli TFs, we sought to characterize YgfI in this study, which has been recognized as one of the uncharacterized TFs in E. coli and belongs to the LysR family [9]. A systematic search was performed to identify target genes controlled by YgfI using the gSELEX-chip system [11]. As a result, a total of 10 strong YgfI-binding sites were identified throughout the E. coli genome. The predicted YgfI target genes are involved in stress responses such as biofilm formation, antibiotic resistance, and hydrogen peroxide resistance. Genes involved in these stress responses also facilitate adaptations to the stationary phase. The transcriptional regulation and physiological roles of YgfI were experimentally examined. As a result, YgfI activated a set of genes in the stationary phase. We therefore propose renaming YgfI as SrsR (a stress-response regulator in the stationary phase).

2. Results

2.1. Regulation Targets of SrsR per gSELEX-Chip Screening

To identify the set of binding sequences for SrsR, we performed gSELEX screening using an improved method [10,11]. In brief, purified His-tagged SrsR was mixed with a collection of E. coli genome fragments of 200–300 bp in length, and SrsR-bound DNA fragments were affinity-isolated. The original substrate mixture of original genomic DNA fragments formed smear bands on PAGE, but after six cycles of gSELEX, DNA fragments with a high affinity for SrsR were enriched, forming sharper bands on PAGE gels.

To identify the comprehensive set of targets directly controlled by SrsR, gSELEX fragments were labeled with Cy3, whereas the original genomic DNA library was labeled with Cy5. A mixture of fluorescently labeled samples was hybridized to an E. coli DNA tiling microarray [19,20]. The ratio of the fluorescence intensity bound to each probe between the SrsR sample and the original library DNA was measured and plotted against the corresponding position along the E. coli K-12 genome. The extent of SrsR binding correlated with its affinity for the SrsR protein. After setting the cutoff level at 200, a total of seven peaks were identified in the gSELEX-chip pattern, and three additional peaks were identified by decreasing the cutoff level to 150 (Figure 1). The highest peak was identified in the spacer region of the yfdO and yfdP genes, which are located in the CPS-53 prophage. Based on the criteria that prokaryotic TF-binding sites are located upstream of the regulatory target genes [12,21], a total of 10 genes (yfdO, yfdP, fucA, fucP, dinJ, yafL, bluR, paaZ, paaA, and cdgI) were predicted to be potential regulatory targets of SrsR (Table 1). Because some of these genes form operons, the total number of regulatory target genes was estimated to be 30 (Table 1).
Table 1. SrsR-binding sites on the *E.coli* genome.

| Peak Position (bp) | Intensity | Function | Operon | Gene | D SrsR Site | D Gene | Operon | Function | SrsR Box |
|-------------------|-----------|----------|--------|------|-------------|--------|--------|----------|----------|
| 1 246,530         | 288       | Antitoxin/DNA-binding transcriptional repressor | dinJ-yafQ | dinJ | < | > | yafL | yafL | predicted lipoprotein and C40 family peptidase | aaAGgcTTGc | 6/11 |
| 2 583,766         | 200       |                      | appY |       | > | < | ompT |       | TCcaTTTTGc | 8/11 |
| 3 1,092,430       | 278       |                      | pgaA |       | < | < | ycfB | ycfB | predicted lipoprotein and C40 family peptidase | cAGgcgTTCa | 6/11 |
| 4 1,213,450       | 207       | Transcriptional repressor | bluR | bluR | < | < | ycfF | ycfF | predicted lipoprotein and C40 family peptidase | gCAGgTTTTGc | 9/11 |
| 5 1,368,672       | 158       |                      | pglE |       | > | > | ycfN | ycfN | predicted lipoprotein and C40 family peptidase | TCacTTTTGc | 9/11 |
| 6 1,451,756       | 170       | Oxein-CoA hydrolase/3-oxo-5,6-dehydrosuberyl-CoA semialdehyde dehydrogenase | paaZ | paaZ | < | > | paaA | paaA | predicted ring 1,2-phenylacetyl-CoA epoxidase subunit | aCgGATTTc | 8/11 |
| 7 1,868,366       | 158       |                      | ycfB |       | > | > | cdgI | cdgI | c-di-GMP binding protein | aaAgTTcTGC | 7/11 |
| 8 2,471,464       | 1206      | CPS-53 (KpLE1) prophage; predicted defective phage replication protein O | yfdONMLK | yfdO | < | > | yfdP | yfdP | CPS-53 (KpLE1) prophage; predicted protein | TCaAAATTTGc, TCAGATTcTcT, TCAGATTcTcT, TCAGATTcTcT | 10/11, 10/11, 10/11, 10/11 |
| 9 2,931,960       | 270       | L-fuculose-phosphate aldolase | fucAO | fucA | < | > | fucP | fucP | FucP fucose MFS transporter | gCGgTTTTGc | 8/11 |
| 10 4,280,360      | 235       |                      | ycfE |       | > | > | actP | actP | predicted lipoprotein and C40 family peptidase | TcaAAATTGc | 9/11 |

gSELEX was used to search for the binding sites in SrsR. Ten binding sites were identified by setting the cutoff level to 150. Column ‘SrsR site’ shown in green represent the SrsR-binding sites inside spacer regions, whereas shown in orange represent the SrsR-binding sites inside ORFs. Column ‘D’ shows the direction of the transcription unit. The potential target genes or operons of SrsR were predicted based on adjacent genes and gene orientation (shown with bold words). Grey shading indicates genes that are not potential targets.
Figure 1. Identification of SrsR-binding sites on the *E. coli* K-12 genome by gSELEX-chip. The gSELEX screening of DNA-binding sequences was performed for SrsR (renamed YgfI), a yet uncharacterized LysR family TF of *E. coli*, using purified C-terminal His-tagged SrsR and a library of DNA segments from the *E. coli* K-12 W3110 genome. Following gSELEX, a collection of DNA fragments was subjected to gSELEX-chip analysis using the tiling array of the *E. coli* K-12 genome. The cutoff level of 150 is shown by a blue line, and the list of all SrsR-binding sites from setting this cutoff level is given in Table 1. Peaks shown in green represent the SrsR-binding sites inside spacer regions, whereas peaks shown in orange represent the SrsR-binding sites inside ORFs.

Genes involved in biofilm formation, such as *bluR* (regulator of the *ycgZ-ymgA-ariR-ymgC* operon, a putative two-component system connector involved in biofilm formation), *cdgI* (putative c-di-GMP binding protein), *fucAO*, and *fucK* (fucose metabolism), are included as an abundant group in the list of predicted targets of SrsR (see below for biofilm formation assay). In addition, *yafQ* (toxin of the YafQ-DinJ toxin-antitoxin system) [22] and *yfdO* (prophage gene) [7] were reportedly involved in antibiotic resistance. Notably, some of these genes are expressed during the stationary phase [23–25]. These perceptions of SrsR target genes suggest that SrsR is involved in the stress response during the stationary phase.

### 2.2. Binding In Vitro of SrsR to the Target Sequences

The binding in vitro of purified SrsR to sequences isolated by the gSELEX screening was examined by PAGE analysis to confirm the formation of SrsR–DNA complexes. Upon increasing the SrsR addition, DNA fragments from all the major peaks of the gSELEX-chip formed SrsR–DNA complexes in an SrsR concentration-dependent manner (Figure 2a–j). Several SrsR–DNA complexes were observed for the *yfdO/yfdP* spacer probe (Figure 2h), indicating that SrsR binds to several sites in the *yfdO/yfdP* intergenic region. Generally, the target promoter, which carries multiple binding sites for the test regulator, shows highly stable TF–probe complexes. Thus, it is reasonable that the *yfdO/yfdP* spacer region showed the highest binding affinity for gSELEX screening (Figure 1). In contrast, the *lacUV5* promoter region, a reference DNA added as a negative control, did not form SrsR–DNA complexes under the same condition (Figure 2k). These results indicate the specific binding of SrsR to all 10 SrsR target sequences (Table 1).
Figure 2. Gel shift assay of SrsR–DNA complex formation. (a–k) Purified SrsR was mixed with 0.1 pmol of each target DNA probe to the SrsR-binding regions shown in Figure 1. SrsR added were (in pmol): lane 1, 0; lane 2, 0.125; lane 3, 0.25; lane 4, 0.5; lane 5, 1; and lane 6, 2. Filled triangles indicate the SrsR–DNA probe complex whereas open triangles indicate free probes.

To identify the binding sequence of SrsR, we performed a DNase-I footprinting assay using the *yfdP/yfdO* spacer as a probe. A 66 bp segment was protected from DNase-I digestion in the presence of SrsR (Figure 3a). Within this segment, four sets of 11 bp sequence TCAGATTTTGC were identified (Figure 3a). Next, a collection of 500 bp sequences from the set of 10 SrsR targets was analyzed using the MEME program [26] to identify the SrsR box sequence. We identified the same 11 bp sequence found in the *yfdP/yfdO* spacer region by the DNase-I footprinting assay (Table 1 and Figure 3b). Multiple and highly conserved SrsR boxes were found only in the *yfdP/yfdO* spacer (Table 1), in accordance with the presence of several SrsR–probe complexes observed in the gel shift assay (Figure 2h). Thus, we concluded that this 11 bp SrsR box sequence is required to ensure tight binding of SrsR.
Figure 3. SrsR binding sequence. (a) DNase-I footprinting assay for identifying the SrsR binding site on the yfdP/yfdO intergenic region. Mixtures of FITC-labeled DNA probe (1.0 pmol) of the yfdP/yfdO spacer sequence and increasing amounts of purified SrsR (0, 10, 20, 40 µM) were subjected to the DNase-I treatment. The sequence shown includes those between the initiation codon of the yfdO gene and the initiation codon of the yfdP gene. The red bar indicates the DNase-I protected region (SrsR box) by SrsR. The DNase-I protected region by SrsR is shown in black. (b) Consensus sequence of the SrsR box. Sequences of all the probes with high-level binding of SrsR were analyzed using THE MEME Suite (Available online: https://meme-suite.org/meme/ (accessed on 1 June 2021)) (see Table 1). WEBLOGO (Available online: http://weblogo.berkeley.edu/logo.cgi (accessed on 1 June 2021)) was used to perform matrix construction.

2.3. Growth-Dependent Expression Level of srsR Gene

Some SrsR targets have reportedly been induced in the stationary phase, including bluR and its targets ycgZ-ymgA-ariR-ymgC [23,27], cdgl [24], and dinJ [25]. RT-qPCR analysis was, therefore, performed to measure SrsR expression levels over time. For this purpose, total RNAs were purified over time from cultured cells grown in an LB medium, and the expression level of srsR was measured. As a result, the mRNA level of srsR increased during the transition from the exponential growth phase to the stationary phase, finally reaching approximately 25 times higher levels than those of the exponential phase (Figure 4). This result indicates that SrsR is influential during the growth transition to the stationary phase and regulates a set of target genes in the stationary phase.
Figure 4. Changing of srsR mRNA level in growth curve. Total RNA was purified from the E. coli wild-type strain at 3, 6, 9, and 24 h of cultivation, and the cell growth was measured by OD$_{600}$ nm every 1 h (closed symbol). The srsR mRNA level was compared by setting the ratio against 16S rRNA as an internal control (open symbol).

2.4. Regulatory Role of SrsR in Expression of the Target Genes In Vivo

To examine the possible influence of SrsR on the target promoters detected in vitro based on SrsR-binding sites, we performed RT-qPCR analysis to determine mRNA levels in vivo for each of the predicted SrsR target genes in the presence, absence, or overexpression of SrsR (Figure 5). Total RNA was prepared from cells of wild-type E. coli K-12, the wild-type strain harboring the SrsR overexpression vector or the control empty vector, and an srsR-deleted mutant; all were grown in an LB medium, and the mRNA levels of individual srsR target genes were measured. Total RNA was prepared at both exponential and stationary phases and subjected to RT-qPCR. Except for the yfdONMLK and yfdPQ operons, the mRNA levels of targets could not be detected in all tested strains from the exponential phase cells (data not shown). In contrast, RNA of all the tested genes were detected in stationary phase cells. However, RNA decreased in the srsR mutant strain compared with the wild-type strain (Figure 5, black bar), whereas RNA from all the tested genes was increased through the SrsR overexpression (Figure 5, white bar). These results indicate that the tested genes influenced the stress response; thus, the members of the SrsR regulon participated in the stress response in the stationary phase. In summary, we propose that SrsR activates a set of genes in the stationary phase.

2.5. Regulatory Role of SrsR in Biofilm Formation

The qRT-PCR analysis suggested that SrsR influences the activation of a set of genes involved in biofilm formation, including bluR and its targets ymgA-ariR-ymgC [23,27,28], cdgl [29], fucAO [30], and fucK [31]. The involvement of SrsR as a key regulator of biofilm formation was then confirmed using the crystal violet staining assay [32]. Under the test conditions, the level of biofilm formation increased in the mutant JW5476 lacking srsR (Figure 6). In contrast, the wild-type strain harboring the SrsR overexpression vector showed markedly decreased biofilm formation, whereas the wild-type strain harboring the empty vector had no effect on biofilm formation (Figure 6).
**Figure 5.** In vivo influence of *srsR* deletion or SrsR overexpression on transcription level of the SrsR targets by RT-qPCR. *E. coli* wild-type BW25113, its *srsR*-deleted mutant JW5476, the wild-type strain harboring the SrsR overexpression vector, and its empty vector were grown in the LB medium at 37 °C. Total RNA was prepared from each strain at 24 h cultured cells and subjected to RT-qPCR analysis. RT-qPCR was repeated three times, and the average values are shown. The y-axis represents the relative level of mRNA of each SrsR target gene between the wild-type and the *srsR* mutant (black bar) as well the relative level between the wild-type harboring SrsR overexpression vector and empty vector (white bar); the ratio of 16S rRNA was set as an internal control between the compared strains. Each experiment was repeated at least three times, and the average means are shown.

**Figure 6.** Level of biofilm formation. *E. coli* K-12 wild-type, the isogenic *srsR* mutant JW5476, the wild-type strain harboring SrsR overexpression vector, and its empty vector were grown in LB medium at 37 °C for 24 h in a plastic tube. The level of biofilm formation was measured by crystal violet staining. The relative level of biofilm represents the mean ± SD of three experiments. Statistically significant differences were obtained using the Student’s t-test with multiple comparisons’ correction and are represented by line over bars with associated *p* values.
The level of biofilm formation has reportedly increased in gene deletion strains of ymgA, ariR, and ymgC, which are repressed by BluR [23,27], and also cdgI [29] grown in LB medium. Moreover, strains overexpressing YmgA or YmgB activate the RcsCDB repressor, which influences the formation of curli and fimbriae [23]. The mRNA levels of bluR and cdgI decreased 1.6 and 5.9-fold in the srsR mutant strain and increased 7.2 and 4.1-fold following the overexpression of SrsR, respectively (Figure 5). Together, these findings support the conclusion that SrsR influences the repression of biofilm formation.

2.6. Regulatory Role of the SrsR in Hydrogen Peroxide Sensitivity

The highest level of SrsR-binding activity was observed in the yfdO/yfdP intergenic region (Figure 1) within the CPS-53 prophage locus. Deletion of the prophage CPS-53 strain has reportedly increased hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) sensitivity [6]. Moreover, this report described that the deletion of yfdK, yfdO, and yfdS in the CPS-53 prophage reduced viability against oxidative stress induced by H\textsubscript{2}O\textsubscript{2}. In accordance with our findings, the mRNA level of the yfdONMLK operon in the stationary phase decreased by 3.7-fold as detected by yfdO mRNA and 15.6-fold as per yfdK mRNA in the srsR mutant strain; however, it increased by 10.7-fold as detected by yfdO mRNA and 1052-fold as per yfdK mRNA in the SrsR overexpression strain (Figure 5).

Based on these findings, we tested the role of SrsR in H\textsubscript{2}O\textsubscript{2} resistance. Both the E. coli wild-type strain and its single-gene srsR-deletion strain were grown in LB medium until the exponential phase, and then exposed to 30 mM H\textsubscript{2}O\textsubscript{2} (Figure 7a). Following a 15 min exposure to H\textsubscript{2}O\textsubscript{2}, the level of cell survival in the wild-type strain was 1.3%, whereas that of the srsR mutant strain decreased to 0.18%, thereby indicating that the presence of SrsR is needed for H\textsubscript{2}O\textsubscript{2} resistance. The effect of SrsR was confirmed by testing the SrsR overexpression vector. The wild-type strain harboring the vector led increased cell viability to 19% under the same conditions, but no effect occurred with the empty vector pCA24NAgp (Figure 7a).

To confirm the regulatory role of SrsR in H\textsubscript{2}O\textsubscript{2} resistance, mRNA levels of the wild-type strain and srsR mutant strain in the exponential phase were determined using Northern blot analysis. The yfdK probe should be included in the yfdONMLK operon transcript, which is shorter than the 2904-nucleotide 23S rRNA on the membrane (Figure 7b). The level of the yfdONMLK transcript was detected in the wild-type strain harboring the SrsR overexpression vector and was further increased by the addition of IPTG for SrsR induction. Since Northern blotting analysis was not sensitive enough to detect the transcript levels of the operon in wild-type and srsR deletion strains, next, we performed RT-qPCR analysis. RT-qPCR analysis showed the mRNA level of the yfdONMLK operon in the exponential phase decreased by 9.5-fold as detected by yfdO mRNA and 2.8-fold as per yfdK mRNA in the srsR mutant strain, whereas it increased by 12.4-fold as detected by yfdO mRNA and 2.2-fold as per yfdK mRNA in the SrsR overexpression strain (Figure 7c). These results suggest that SrsR activates the yfdONMLK operon not only during the stationary phase but also in the exponential phase. The H\textsubscript{2}O\textsubscript{2} sensitivity of the yfdK mutant decreased by approximately 103 times (Figure 7a). Similarly, the single-gene yfdO and yfdS deletion strains exhibited the same level of sensitivity (data not shown), thus confirming the influence of the yfdONMLK operon in hydrogen peroxide sensitivity. This finding is in accordance with previously published observations [6], while the srsR mutant showed a 10-fold increase in sensitivity compared with the wild-type strain (Figure 7a). Although less severe than the yfdK deletion mutant, increased sensitivity to oxidative stress was also observed in the srsR mutant, a regulator of the yfdKONMLK operon, suggesting that SrsR activates the operon and increases the tolerability against hydrogen peroxide.
Figure 7. Sensitivity test against hydrogen peroxide. (a) Survival test for the *E. coli* wild-type BW25113, its srsR-deleted mutant JW5476, the wild-type strain harboring SrsR overexpression vector, its empty vector, and yfdK mutant JW2350. Each strain was inoculated until the OD$_{600}$ reached 0.4 (exponential phase) and then subjected to 30 mM H$_2$O$_2$ for 15 min. The relative level of sensitivity represents the mean ± SD of three experiments. Statistically significant differences were obtained using the Student’s t-test with multiple comparisons’ correction and are represented by line under bars with associated $p$ values. (b) *E. coli* wild-type BW25113 and its srsR-deleted mutant JW5476, the wild-type strain harboring its empty vector, and SrsR overexpression vector with or without IPTG were grown in an LB medium at 37°C until the OD$_{600}$ reached 0.4 (exponential phase). Total RNA was prepared from both wild-type and the srsR mutant and subjected to Northern blot analysis. The mRNA of the yfdONMLK transcription unit detected with yfdK DIG-labeled hybridization probes is shown. The amounts of total RNA analyzed were examined by measuring the intensity of ribosomal RNAs. (c) The same RNAs as in panel (b) were subjected to RT-qPCR analysis. The y-axis represents the relative level of mRNA of yfdO and yfdK between the wild-type and the srsR mutant (black bar) as well the relative level between the wild-type harboring SrsR overexpression vector and empty vector (white bar); the ratio of 16S rRNA was set as an internal control between the compared strains. Each experiment was repeated at least three times, and the average means are shown.

3. Discussion

A set of stress-response genes exists in bacterial genomes [2,3]. Some of these genes are expressed for survival during the stationary phase. To detail the mechanisms of stress adaptation, it is essential to clarify the regulatory roles of stress-response genes. In this study, we identified that the hitherto uncharacterized *E. coli* transcription factor SrsR (renamed from YgfI) is a local regulator of a group of stress-response genes, including biofilm formation, hydrogen peroxide sensitivity, drug resistance, and cold shock stress. The expression level of SrsR increased in the stationary phase and during activation of the set of stress-response genes; we, therefore, propose to rename *ygfI* to *srsR* (stress-response regulator in stationary phase).

Several genes have been identified in biofilm formation in *E. coli* K-12 [33–35], whereas only a limited number of regulators have been identified in the deceleration of biofilm formation. One thoroughly characterized regulator inhibiting biofilm formation influences
the repression of the csgDEFG operon, which encodes a set of genes involved in the formation of curli fimbriae. The level of biofilm formation decreases, owing to the repression of the csgD promoter, where CpxR, FliZ, H-NS, MqsA, BtsR, RcsAB, and RstA act as repressors [18,36]. Another deceleration system of biofilm formation includes BluR as a repressor of the ycgZ-ymgA-ariR-ymgC operon, which inhibits biofilm formation via the cascade for activating rcsDCB and repressing curli and fimbriae formation [23]. Similar to these transcription factors, SrsR regulates a set of genes including bluR as a new inhibitor of biofilm formation during the stationary phase (Figure 6).

While SrsR suppressed biofilm formation, SrsR increased the tolerability against hydrogen peroxide (Figure 7). E. coli carries several regulators for activating a set of genes for oxidative stress, including OxyR, SoxR, and SoxS [37]. Recently, it was reported that Cys25 of OxyR, which is S-nitrosylation in planktonic cells, plays an important role in suppressing biofilm formation [38]. Such transcriptional regulation by SrsR and OxyR suggests the importance of switching between the biofilm and planktonic lifestyles as well as conferring oxidative stress tolerance to planktonic cells in bacterial adaptation to the stressful environment.

For Enterobacteriaceae, including E. coli, survival and infection in the host gut is important to their survival strategy. In contrast, the host produces reactive oxygen species such as hydrogen peroxide to eliminate pathogens [39,40]. This study showed that not only the yfdONMLK operon, which is significant for oxidative stress tolerance, but also the fucAO operon and fucPIKUR operon, which are required for survival in the host gut [30,31], were activated by SrsR (Figure 5). These findings suggest that SrsR may be a crucial transcription factor for survival strategies of E. coli in the host gut.

The SrsR regulon may also be involved in adaptation to nitrogen starvation conditions, since it had been reported that srsR is repressed by Nac (nitrogen assimilation control), a transcription factor that is expressed under nitrogen starvation conditions [41]. SrsR belongs to the LysR family, which includes the maximum number (47 members) of TFs in E. coli [9]. The TF of the LysR family contains a DNA-binding domain at the C-terminus and a co-inducer-binding domain at the N-terminus. Further studies are needed to identify a co-inducer to elucidate further physiological functions of SrsR.

4. Materials and Methods

4.1. Bacterial Strains and Plasmids

Escherichia coli K-12 W3110 type-A [42] was used as the DNA source to construct the SrsR expression plasmid. The E. coli K-12 W3110 type-A genome was also used to construct the DNA library needed for gSELEX screening. E. coli DH5α cells were used to perform plasmid amplification, and E. coli BL21 (DE3) cells were used as the SrsR expression. E. coli BW25113 [43], its single-gene knockout mutants (JW5476 for srsR and JW2350 for yfdK) [44], and the expression plasmid from the ASKA clone library were obtained from the E. coli Stock Center (National Bio-Resource Center, Saitama, Japan). Plasmid pPET21 was used to construct the SrsR expression plasmid pSrsR. Cells were grown in an LB medium at 37 °C with constant shaking at 150 rpm. When necessary, 20 µg mL⁻¹ kanamycin or 30 µg mL⁻¹ chloramphenicol was added to the medium. When observing the effect of SrsR overexpression, IPTG at a final concentration of 0.5 mM was added from the start of inoculation. Cell growth was monitored by measuring turbidity at 600 nm.

4.2. Purification of SrsR Protein

Plasmid pSrsR for expressing and purifying SrsR was constructed according to the standard procedure [10]. In brief, SrsR coding sequences were PCR-purified using the E. coli K-12 W3110 genome DNA as a template and inserted into the pET21a (+) vector (Novagen, Darmstadt, German) between NdeI and NotI sites, leading to the construction of pSrsR. The expression plasmid, pSrsR, was transformed into E. coli BL21 (DE3) cells. Transformants were grown in an LB medium, and SrsR was expressed by adding IPTG to the middle of the exponential phase. The SrsR protein was purified by affinity purification
using a Ni-nitrilotriacetic acid (NTA) agarose column. The affinity-purified SrsR protein was stored, frozen, in the storage buffer at \(-80^\circ\text{C}\) until use. Protein purity was greater than 95\%, as determined by SDS-PAGE.

4.3. Genomic SELEX (gSELEX) Screening of SrsR-Binding Sequences

The gSELEX screening was performed as previously described [10,11]. Briefly, a mixture of DNA fragments of the \(E.\ coli\) K-12 W3110 genome was prepared by sonication of purified genomic DNA and cloned into a multi-copy plasmid, pBR322. For each gSELEX screening, the DNA mixture was regenerated by PCR. For gSELEX screening, 5 pmol of the mixture of DNA fragments and 10 pmol SrsR were mixed in a binding buffer (10 mM Tris-HCl, pH 7.8 at \(4^\circ\text{C}\), 3 mM magnesium acetate, 150 mM NaCl, and 1.25 mg/mL bovine serum albumin). The SELEX cycle was repeated six times to enrich the SrsR-binding sequences. Mapping of SELEX fragments along the \(E.\ coli\) genome was also performed using a gSELEX-chip system with a 43,450-feature DNA microarray [45]. The genomic SELEX sample obtained using SrsR was labeled with Cy3, whereas the original genomic DNA library was labeled with Cy5. Following hybridization of samples to the DNA tiling array (Agilent technology, Santa Clara, CA, USA), the Cy3/Cy5 ratio was measured, and the peaks of the scanned patterns were plotted against the positions of the DNA probes along the \(E.\ coli\) K-12 genome.

4.4. Gel Shift Assay

A gel shift assay was performed according to the standard procedure [46]. Probes for the SrsR-binding target sequences were generated by PCR amplification using a pair of primers (Table S1a) and Ex Taq DNA polymerase (TaKaRa, Kyoto, Japan). A mixture of each probe and SrsR was then incubated at \(37^\circ\text{C}\) for 30 min in the gel shift buffer. After adding the DNA loading solution, the mixture was directly subjected to PAGE. DNA in the gels was stained with GelRed (Biotium, Fremont, California, USA) and detected using LuminoGraph I (Atto, Tokyo, Japan).

4.5. DNase-I Footprinting Assay

A DNase-I footprinting assay was performed under standard reaction conditions [47]. Probes for the \(yfdP/yfdO\) sequence were generated by PCR amplification using a pair of primers (Table S1b) and Ex Taq DNA polymerase (TaKaRa). For footprinting, 1.0 pmol of FITC-labeled probes was incubated at \(37^\circ\text{C}\) for 30 min with various concentrations of purified SrsR in 25 \(\mu\text{L}\) of a solution containing 10 mM Tris-HCl (pH 7.8), 150 mM NaCl, 3 mM magnesium acetate, 5 mM CaCl\(_2\), and 25 mg/mL bovine serum albumin. Following incubation, DNA digestion was initiated by adding 5 ng DNase-I (TaKaRa). Then, following digestion for 30 s at \(25^\circ\text{C}\), the reaction was terminated through an addition of 25 \(\mu\text{L}\) of phenol. DNA was precipitated from the aqueous layer by ethanol, dissolved in a formamide dye solution, and analyzed by electrophoresis on a 6% polyacrylamide gel containing 8 M urea.

4.6. Consensus Sequence Analysis

To analyze the SrsR-binding sequence, an SrsR-binding sequence set identified by the gSELEX-chip was analyzed using the MEME program [26]. Sequences were aligned, and a consensus sequence logo was created by WEBLOGO (Available online: http://weblogo.berkeley.edu/logo.cgi (accessed on 1 June 2021)).

4.7. RT-qPCR Analysis

RT-qPCR was performed according to standard procedure [48]. \(E.\ coli\) cells were inoculated into an LB medium at \(37^\circ\text{C}\) under aeration, with constant shaking at 150 rpm. After inoculation for 24 h, the total RNA was extracted using an ISOGEN solution (Nippon Gene). Total RNAs was transcribed to cDNA with random primers using the THUNDERBIRD SYBR qPCR RT Set (TOYOBO, Osaka, Japan). Quantitative PCR (qPCR) was
conducted using the THUNDERBIRD SYBR qPCR Mix (TOYOBO) and a LightCycler 96 system (Roche, Barsel, Switzerland). The primer pairs used are described in Table S1c. The cDNA templates were serially diluted 4-fold and used in the qPCR assays. The qPCR mixtures, each containing 10 µL of THUNDERBIRD SYBR qPCR Mix (TOYOBO), 1 µL of each primer (5 µM stock), 7 µL of water, and 1 µL of cDNA, were amplified under the following thermal cycle conditions: 95 °C treatment for 2 min, at 45 cycles of 10 s at 95 °C, and 20 s at 55 °C, and then incubation for 20 s at 72 °C. The 16S rRNA expression level was used to normalize the varying levels of the test samples, and the relative expression levels were quantified using the Relative Quantification software provided by Roche. The results are presented as the average of three independent experiments.

4.8. Biofilm Assay

The crystal violet staining method was used as previously described in another study [32]. E. coli cells were grown in an LB medium at 37 °C in a 2.0 mL tube. Following incubation for 24 h, planktonic cells were discarded, the tube was washed twice with PBS (-) and subsequently stained with 0.1% crystal violet for 20 min at room temperature. After extensive washing with H2O, biofilm-bound crystal violet was extracted with 500 µL of 70% ethanol and measured at OD595.

4.9. Hydrogen Peroxide Test

E. coli cells were inoculated in an LB medium at 37 °C under aeration with constant shaking at 150 rpm until the OD600 reached 0.4, and then hydrogen peroxide was added to the culture to a final concentration of 30 mM. Following this addition, incubation continued for 15 min, and then the cells were diluted and spread on LB agar plates. Untreated hydrogen peroxide-treated cells were also plated. The plates were incubated at 37 °C overnight, and then the number of colonies on the plates was counted. The survival rate was calculated as the ratio of colony numbers between hydrogen peroxide-treated cells and untreated cells.

4.10. Northern Blot Analysis

Total RNAs were extracted from the exponential phase E. coli cells (OD600 = 0.4) by ISOGEN solution (Nippon Gene). RNA purity was checked by electrophoresis on 1.5% agarose gel with formaldehyde present, followed by staining with GelRed. Northern blot analysis was performed as described in previous research [49]. DIG-labeled probes were prepared by PCR amplification using W3110 genomic DNA (50 ng) as a template, with a pair of primers (Table S1d), DIG-11-dUTP (Roche) and dNTP, as substrates, gene-specific forward and reverse primers, and Ex Taq DNA polymerase. Total RNAs (3 µg) were incubated in a formaldehyde-MOPS (morpholinepropanesulfonic acid) gel-loading buffer for 10 min at 65 °C for denaturation, subjected to electrophoresis on formaldehyde-containing 1.5% agarose gel, and then transferred onto nylon membrane (Roche). Hybridization was performed using a DIG easy Hyb system (Roche) at 50 °C overnight with a DIG-labeled probe. To detect the DIG-labeled probe, the membrane was treated with anti-DIG-AP Fab fragments and CDP-Star (Roche), and the image was scanned using LuminoGraph I (Atto).

Supplementary Materials: The following supporting information can be downloaded at https://www.mdpi.com/article/10.3390/ijms23116055/s1.

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Abbreviations

SELEX (systematic evolution of ligands by exponential enrichment), TF (transcription factor), PAGE (polyacrylamide gel electrophoresis), DNase (deoxyribonuclease), RT-qPCR (reverse transcription-quantitative polymerase chain reaction), LB (Luria–Bertani), SrsR (stress-response regulator in stationary phase), NTA (Ni-nitrilotriacetic acid), MOPS (morpholinepropanesulfonic acid), DIG (digoxigenin).

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