INTRODUCTION

Certain metals, for example, silver (Ag), mercury (Hg), and tellurium (Te) are highly toxic to the living organisms and have microcidal property even at low concentration. Prokaryotes, fungi, and higher plants have evolved with molecular mechanisms to tolerate such harmful metal ions (Oves, Saghir Khan, Huda Qari, Nadeen Felemban, & Almeelbi, 2016). Investigations have demonstrated that mercury is...
readily accumulated by higher plants and causes severe damage to the cellular metabolism and physiology of the plants (Cargnelutti et al., 2006). Mercury hinders the photosynthetic electron transport and is exceedingly harmful as a result of the capacity of Hg\(^{2+}\) to bind with proteins (Rai, Agrawal, & Agrawal, 2016). Once, in the cell, Hg\(^{2+}\) forms covalent bonds with cysteine residues of proteins, it drains antioxidants. These mercuric ions in the aquatic environment persist for very long periods in the sediments (Randall & Chattopadhyay, 2013). However, the microorganisms evolved with a molecular mechanism to resist the inorganic and organic forms of mercurial compounds involving mer-operon.

Mercury-tolerant bacterial strains reported till date contain mer-operon in their genomes, which code for proteins involved in mercury detoxification. The main open reading frames, merR, merA, and merP, code for a transcriptional regulator, mercuric reductase, and a periplasmic mercury binding protein, respectively. In some bacterial species, additional genes such as merB code for organomercurial lyase, merD an additional transcriptional regulator, merE and merF, auxiliary transporters are present. The enzymatic reduction of the mercuric ions to elemental mercury, catalyzed by mercury reductase, is the main detoxification mechanism in bacteria (Barkay, Miller, & Summers, 2003). This key detoxification protein reduces the toxic Hg (II) to Hg (0). As a result of its low dissolvability in water and moderately high-vapour weight, the elemental mercury is discharged from the cell (Barkay et al., 2003; Silver, 1996). The biochemical basis of protection from inorganic-mercury compounds, for example, HgCl\(_2\) seems to be similar in few bacterial species. Studies have shown that a wide range of microorganisms such as Shigella flexneri, Pseudomonas aeruginosa, Serratia marcescens, Xanthomonas sp., and Staphylococcus aureus possess mer-operon. However, when mer genes were compared, the number and their operonic organization differ among various bacterial genera (Barkay et al., 2003; Hobman, Wilson, & Brown, 2000).

Cyanobacteria, which occur in almost all environmental niches contain genes that code for a putative MerR-like transcriptional regulator and MerA, mercury reductase. In Synechocystis sp. PCC6803 (hereafter referred as Synechocystis), two putative MerR family transcription factors are predicted. One of them, Slr0794 has been reported to be involved in the regulation of genes related to Ni\(^{2+}\) toxicity (García-Domínguez, Lopez-Maury, Florencio, & Reyes, 2000). Several MerR transcription factors, though they are called as mercury-resistant regulators and classified under MerR family, are also known to be involved in resistance to toxic metals other than mercury. For example, ZntR (Brocklehurst et al., 1999) in Escherichia coli and PbrR inRalstonia metallidurans CH34 (Harley & Reynolds, 1987) are MerR family members involved zinc and lead resistance, respectively. In Synechocystis, Slr1849, codes for MerA and biochemically it was demonstrated to be a mercury reductase (Marteyn et al., 2013). However, the mer-operon organization and regulation in cyanobacteria are poorly understood. Being photosynthetic bacteria, progenitors of higher plant chloroplasts and occurring in almost all habitats on the earth, it is important to unravel the regulatory mechanism of mercury tolerance in cyanobacteria. In this study, we show that the mer genes of cyanobacteria are phylogenetically distinct from other bacterial phyla. We also demonstrate a mercury-induced regulation of merA gene by a putative MerR family protein in Synechocystis.
inactivate crhR gene (Prakash et al., 2010). The Dral site in the Sp′ specific primers was underlined. The Dral-digested Sp′ cassette was cloned at the Hpal-digested pTSlr0701 DNA construct by blunt end ligation. The final plasmid, pTSlr0701:sp′ in which the slr0701 ORF was disrupted by Sp′ cassette was used to transform wild-type Synechocystis cells. The site of insertion of the Sp′ cassette was confirmed by sequencing the pTSlr0701:sp′ DNA construct using the slr0701-F primer. The segregation analysis was performed by PCR amplification to check the extent of replacement of wild-type copies of slr0701 with the pslr0701:sp′ in slr0701—mutant strain was verified by PCR amplification using genomic DNA as a template.

The slr0701′ complement strain was generated to check whether the phenotype change observed in slr0701 mutant was due to inactivation of slr0701 gene. The PCR-amplified slr0701 ORF with flanking regions as mentioned above was used for making complement. The resulting 1,340 bp DNA fragment was cloned by blunt end ligation at Smal site located within the kanamycin (kan′) cassette into a cyanobacterial replicative vector pVZ321vector (Zinchenko, Piven, Melnik, & Shestakov, 1999). The recombinant vector, pVZ-slrl0701, was introduced into slr0701-mutant by triparental mating as described in Zinchenko et al., 1999. We transferred the recombinant replicative pVZ-slrl0701 into the recipient slr0701-mutant strain by conjugal mating with donor and helper E. coli strains. E. coli-DH5α carrying pVZ-slrl0701 was served as a donor strain. The recipient, donor, and helper strains were mixed in a ratio of 10:1:1, on a membrane filter (Millipore, Catalog No: GSWP 04700), and incubated overnight on BG-11 solid agar supplemented with 5% LB under dim light for conjugal transfer. The colonies thus developed were selected with antibiotic selection pressure of 35 µg/ml of chloramphenicol (Cm r)

2.5 | Measurement of photosynthetic electron transport rate

Photosynthetic oxygen evolution was estimated in 1 ml of cell suspension (OD730 = ~1, around 5 µg/mL chlorophyll), utilizing an oxygen electrode (Oxygraph plus, Hansatech Instruments Ltd., Norfolk, England). The cells were grown photoautotrophically under continuous air bubbling by illuminating continuously with 70 µE m−2 s−1 light. PSII activity was measured in presence of 1.0 mM p-benzoquinone (PBQ) as described earlier (Sireesha et al., 2012). Photonsynthetic oxygen evolution was recorded at 1.000 µE m−2 s−1 light. Three independent cultures of wild-type and slr0701-mutant cells were estimated with or without HgCl2. Cultures were grown till 0.6 OD at 730 nm and measured the oxygen evolution and the data was used as a reference. Then 500 nM of HgCl2 was added to a final concentration of the cultures and measured oxygen evolution at 12 hr and 24 hr.

2.6 | Measurements of chlorophyll a fluorescence

We recorded chlorophyll a fluorescence of Synechocystis cell suspensions with a continuous excitation PEA fluorometer (PEA, Hansatech, King’s Lynn, Norfolk, UK). The PEA fluorometer provides continuous excitation at 650 nm (3,000 µE m−2 s−1; ∆λ = 22 nm). It perceives fluorescence at wavelengths above 700 nm (50% transmission at 720 nm) and records it continuously from 10 µs to 300 µs. The fluorescence curves were recorded using wild-type and slr0701-mutant cells as described (Sireesha et al., 2012). Cultures were grown till OD at 730 nm reached to 0.6, and then measured the chlorophyll a fluorescence. These data were used as a reference. Then 500 nM of HgCl2 was added to a final concentration of the cultures and measured fluorescence at 12 hr and 24 hr.

2.7 | Transcript analysis of mercury responsive genes

Wild-type and slr0701-mutant cells were cultivated at 70 µE m−2 s−1 of light. 50 ml of the cells were killed immediately by the addition of 50 ml equal volume of cold 5% w/v, phenol in ethanol, and then total RNA was isolated as described earlier (Srikumar et al., 2017). The RNA was treated with DNase I (Cat. No. 89836, Thermo Fischer Scientific) to remove the DNA contaminants. One-microgram of RNA from total RNA was converted to cDNA using Takara Kit (Cat. no. 6110A). qRT-PCR was done using the SYBR® Premix Ex Taq™ II (TliRNase H Plus; feline. no. RR820A). Every reaction was completed in a 25 µl volume containing 12.5 µl of Power SYBR Green Master Mix, 0.2 µM of specific primers.

(Table 1) and 5 µl of dilute cDNA was kept in triplicates for running qRT-PCR (Mx3005P, Agilent Technologies). The instrument was conditioned for 95°C for 10 min, and then 40 cycles of 30 s at 95°C, 60 s at 60°C, and 60 s at 72°C. For every reaction, the melting
curves were investigated and the PCR product was analyzed on an agarose gel with the end goal to affirm the specificity of the RT-PCR. Expression levels were normalized using the gap1 gene as an internal reference.

### 2.8 Overexpression and purification of Slr0701

PCR amplification of slr0701 ORF was performed, with the specific primers slr0701-ExF (5′-tgacGG TAG CGT GAG CAT TAT GTT GGC/ TTT CTCGAG CAA CGT CTC ATT TAA CCA AC-3′) and slr0701-ExR (5′-gacGTA GAG GGC ACA AGT CAA CTG) to generate a DNA fragment of 500 bp. The PCR product was purified and then cloned into pET-28a (+) at the Nhel and EcoRI restriction cloning sites. The C-terminally 6x-His-labeled Slr0701 protein was overexpressed in Rosetta™ (DE3), which had been transformed with pET-slr0701 and was induced with 0.4 mM final concentration of isopropyl β-D-1-thiogalactopyranoside (IPTG). The E. coli cells were pelleted with centrifugation at 4,000 rpm for 5 min at 4°C. The cell-pellet was resuspended in the lysis solution containing 6 M urea, 50 mM Tris, 300 mM NaCl, 20 mM Imidazole/pH 8 to washout impurities, and undesirable protein. 6x-His–Slr0701 protein was eluted in an elution solution containing 6 M urea, 50 mM Tris, 300 mM NaCl, and 200 mM Imidazole/pH 8. These elutes were analyzed on 12% SDS-PAGE. The Slr0701 protein was pooled in elution solution of pH 8 and serially dialysed at 4°C for seven times using the initial solution containing 50 mM Tris, 300 mM NaCl, and 100 mM Imidazole. At each round of buffer replacement, urea and imidazole concentrations were gradually decreased to attain the final buffer concentration of 50 mM Tris and 300 mM NaCl.

### 2.9 Western blotting analysis

Western blotting experiment was performed as presented earlier with some modifications (Prakash et al., 2010). Polyclonal antibodies induced in rabbit against 6x-His–Slr0701 protein was utilized as primary antibody, and an HRP-linked antibody induced in goat against rabbit IgG was utilized as the secondary antibody. Wild-type cells were grown with and without 500 nM HgCl₂ and the cells were harvested for protein extraction at different time intervals. Protein samples were separated using SDS-PAGE and were blotted onto PVDF membrane (Cat. No. IPVH00010, Immobilon-P; Merck Millipore) with a semi-dry transfer apparatus (TE77-PWR semi-dry transfer unit, GE Healthcare). The levels of Slr0701 were detected immunologically with the ECL Plus, immunoblotting system (Cat. No. 1705060-61, Bio-Rad). We used anti-rabbit secondary antibodies conjugated to horseradish peroxidase (1:5,000) for detection. Blot was scanned using Bio-Rad chemidoc (XRS®) and analyzed with Image Lab software (Bio-Rad).

### 2.10 Prediction of cis-regulatory elements using MEME suite

Intergenic DNA regions of cyanobacterial orthologs obtained from blastP were taken and submitted to MEME (Motif Extraction by Multiple Expectation Maximizations) version 4.3.0. MEME was run using the default parameters to find maximum of three motifs per sequence with the distribution of zero or one occurrence per sequence (Bailey & Elkan, 1994).

### 2.11 Gel retardation assays

The slr0701 and slr1849 upstream DNA regions were PCR amplified using oligonucleotide primers pslr0701-FP-B: 5’CCC TCC CAT TTC TTC TCT GTT CAC ACT C3′ and pslr0701-RP-B: 5’GGG GTT AGG TAG GGC ACC AAG TTA C3′, and pslr1849-FP-B: 5’CGT GAA TGT AAA TGA AAA AAC AAG CG3′ and pslr1849-RP-B: 5’ACT TAG AAT TGC TGA TTG CCG TAT TAC 3′. The sizes of amplified PCR products corresponding to slr0701 and slr1849 upstreams were 231 bp and 230 bp, respectively. The binding reactions were performed in the 20 µl reaction volume containing 80 ng of upstream DNA and 4 µl of the binding buffer with different concentration of 6x-His-Slr0701. The mixtures were incubated for 30 min with and without 50 µM HgCl₂ at 25°C. The reaction mixtures were loaded on a non-denaturing 6% polyacrylamide gel. Electrophoresis was carried out at 4°C and 200 V in 0.5X Tris-borate/EDTA. Gels were washed and incubated in a clean plastic container with sufficient 1X SYBR® Green EMSA staining solution about 50 ml with continuous, gentle agitation at 50 rpm for ~20 min, protected from light. For visualization, gel was washed twice with 150 ml of ddH₂O for ~10 s to remove excess stain. The gel was scanned using Bio-Rad chemidoc (XRS®) and analyzed with Image Lab software (Bio-Rad).

### 2.12 Biomolecular interaction analysis by SPR

All the experiments in surface plasmon resonance (SPR) analysis were performed at 25°C using Biacore T200 instrument (GE Healthcare Life Sciences). To study the interactions under real-time conditions between 6x-His-Slr0701 protein and P<sub>slr0701</sub> a DNA fragment was immobilized on the CM5 chip (GPI immobilization kit, GE Healthcare). The SPR experiment was carried out at 25°C using a flow rate of 30 µl/min.

### Table 1: Primers used for qRT-PCR

| Gene     | Forward primer (5′-3′)     | Reverse primer (5′-3′)  |
|----------|---------------------------|-------------------------|
| slr0701  | GTC TATTCCGTCTGCAAAGG     | CTTTGATTCACCTTTCCAGCTG |
| slr1849  | TGATTTCGCCGCTTGGATGG     | TCCCGAAATCTTTGAGCAGAG   |
| slr0884 (gap1) | TCGCCTACCGGTGGAGCC | GGGCACAATGGCCTCAACAA |

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covering the predicted conserved DNA binding motif by MEME, was synthesized (Integrated Device Technology, Inc.). Biotinylated oligonucleotide containing a sequence 5’BiolsG/TAC TAC AAT ATC CCT ATA CTT AGG TAT AAG GTT GTA GGT TGA TAT AG3’ and the complementary sequence 5’CTA TAT CAA CCT ACA ACC TTA TAC CTA AGT ATA GGG AAT ATA TGG TAG TA3’ were hybridized to obtain double-stranded DNA duplex, P_{slr0701} SA sensor chip (GE Healthcare Bio-Sciences AB, Uppsala, Sweden) was kept at room temperature for at least 30 min before use. The instrument was first primed three times with HBS_EP+ running buffer. The flow cell 3 (FC 3) was used as the reference cell (without biotinylated ligand) and flow cell 4 (FC 4) was used for the immobilization. According to the manufacturer’s instructions before immobilization the SA chip was conditioned for three consecutive injections of 1 minute duration each with 1 M NaCl in 50 mM NaOH (activation buffer), followed by immobilization of the DNA. Reference cells were washed with 50% isopropanol in 1 M NaCl and 50 mM NaOH. After surface activation of the sensor chip, immobilization of 500 nM of biotinylated P_{slr0701} dissolved in a final volume of 128 µl HBS_EP+ buffer was performed.

Slr0701 protein was chosen as an analyte to study the protein–DNA interactions in HBS_EP+ running buffer. The method was standardized with varying concentration of Slr0701 from 2 to 32 µM with and without 62 µM of mercury chloride. The experiment was performed in triplicates. At the end of each dissociation period, the sensor chip was regenerated using 10 mM Glycine-HCl pH 2.0 for a contact time of 30 s, with a flow rate of 30 µl/min followed by a subsequent wash with running buffer for each cycle, passing through FC3 and FC4 in series, recording the reference subtracted signal. Biacore T200 evaluation software version 2.0 was used for data analysis, 1:1 Langmuir model fit was used for determining the binding kinetics.

2.13 Analysis of P_{slr0701} and P_{slr1849} promoter activity

To monitor the activity of promoters, we fused LuxAB, reporter gene coding for luciferase to the promoter of slr0701 and slr1849. The modified pSyn2030-2031P_{psr} vector was used to construct the desired promoter clones within slr2030-slr2031 neutral site. The vector was linearized using inverse PCR technique with specific primers FP: 5’ AGA AGG AGC TGC AGA TCT CAT ATG C3’ and RP: 5’ GAG AGC TTG ATC GGC AGC TAA G3’ and the upstream DNA fragments were PCR amplified with the infusion primers specific to pslr071FP: 5’ GAT CAA CGT CTC ATT ACA GTC GAG AAC TAA GAC AA3’, pslr071RP: 5’ GAT CAA CGT CTC ATT ACA GTC GAG AAC TAA GAC AA3’ and pslr1849 FP: 5’ GAT CAA CGT CTC ATT ACT TAG AAT TGC TGA TTT CC3’, pslr1849RP: 5’ ATT TCC AAA CTT CAT TTA AGG ATT AAT TGT TTA ATG3’ 100 µl wild-type and slr0701-mutant Synechocystis cells were transformed with 1 µg of plasmid DNA containing chloramphenicol (Cm) gradient plate. After 10 days, the colonies from the gradient plate were picked and plated onto BG-11 agar plate containing chloramphenicol at a concentration of 10 µg/ml. For complete segregation, Cm-resistant colonies were grown at increasing Cm concentrations (up to 50 µg/mL) and finally transferred into the liquid BG-11 medium. In case of pslr0701-mutant cells, screening was performed in the presence of spectinomycin. Luminescence of the luciferase reactions was induced by the addition of 500 nM HgCl₂ to the cyanobacterial cell suspension. Light emission was monitored in a photomultiplier based luminometer (BioOrbit, Labsystems) following the addition of N-decanal (50 mM decanal in methanol/water (50%, v/v), the final concentration in the suspension was 1 mM). We used the LuxAB plasmid without P_{slr0701} and P_{slr1849} in the wild-type and slr0701-mutant as a control to evaluate the metal-induced regulation of slr0701 and slr1849 promoters.

2.14 CD spectroscopy

The upstream of slr0701 was amplified by PCR using specific primers FP 5’CCC TCC CAT TTC TCT GTC TCT ACT AC3’ and RP 5’GGG GTT AGG TAG GAG ATC GCC AAA TTA C3’ from the genomic DNA of Synechocystis. A 231 bp PCR amplified product was purified and used for CD spectral analysis. CD spectra were recorded with a Jasco J-1500 spectropolarimeter. Far UV-CD spectra of 20 µM of Slr0701 were recorded in the presence of varying concentrations of HgCl₂ ranging from 20–200 µM. The ellipticity value at 222 nm was plotted against the concentration of HgCl₂. Further, the near UV-CD spectra of Slr0701 protein was analyzed in the presence of DNA, without and with 25 µM of mercuric chloride. The protein and DNA concentrations used were 5 µM each. The spectra were collected with a scan speed of 20 nm/min and an average of three scans was calculated.

3 RESULTS

3.1 Diversity in the genetic organization of mer genes in cyanobacteria

To identify the mercury responsive genes in Synechocystis, sequences of well-characterized mer proteins from certain gram-positive and gram-negative bacteria were used as queries and blastP search was performed against Synechocystis genome database as described in the experimental methods. The best hits of blastP search using well-characterized MerR and MerA were a putative merR transcriptional regulator and a mercury reductase encoded by ORF numbers slr0701 and slr1849, respectively. The evolutionary relationship of Slr0701 and Slr1849 with their respective orthologs is shown in Appendix information Figure A1. The phylogenetic trees demonstrate their presence across different bacterial and cyanobacterial genera. The homologs for Slr0701 and Slr1849 proteins were identified mainly in freshwater cyanobacterial species and some other non-photosynthetic bacteria isolated from soil as well. Though several of these cyanobacterial species were originally isolated from freshwater ponds, they were observed to exist in soil crusts also (Tirkey & Adhikary, 2005). It was also observed that both, MerR and MerA homologs
from cyanobacterial species formed a separate clade (Figure A1). Homologs for Slr0701 and Slr1849 proteins were also present in Geobacter pickeringii, a metal-reducing isolate from sedimentary kaolin deposits (Shelobolina et al., 2007), and Acidithiobacillus prosperus, isolated from geothermally heated sea sediments (Khaleque, 2017). The orthologs from G. pickeringii formed a separate clade along with the cyanobacterial mer genes. Notably, Slr0701 and Slr1849 of Synechocystis showed 53.2% and 46% sequence similarity to a well-studied B. megaterium MerR and MerA homologs, respectively (Wang et al., 1989). Interestingly, Slr1849 showed 70.2% and 43.1% sequence similarity with the MerA of metal-reducing bacterial species, G. pickeringii and A. prosperus, respectively. But, the putative regulator, Slr0701, showed only 37.3% and 44.8% similarity to MerR of these metal-reducing bacterial species. This suggests that the cyanobacterial MerA might have horizontally transferred and independently acquired from metal reducing Geobacter species.

In bacteria, proteins involved in resistance to mercury are encoded by mercury responsive transcription factor; merR, an inorganic-mercury reductase; merA, alternative transcription factor; merD, periplasmic transporters; merP, merT and merC, and organo-mercurial lyase; merB. These genes are organized as mer operon in most of the bacterial species reported to date (Barkay et al., 2003). Comparison of the organization of mer genes from a diverse range of bacterial species has revealed considerable similarity in their genetic organization (Figure A2). Almost all these operons contain a regulatory gene, merR. In gram-positive bacteria, the mer genes including merR form a single operon. Downstream to the regulator, merR other mer structural genes merA, merD, merP, and merT are located in gram-positive bacteria. In some gram-positive bacteria, an additional ORF, merC, is also part of the operon. In some gram-negative bacteria, merR is divergently located to other mer structural genes which form an operon without merR. In certain bacterial species, such as B. megaterium and Pseudomonas sp. K62, an organo-mercurial lyase, merB was also reported as a part of the mer-operon (Kiyono, Omura, Inuzuka, Fujimori, & Pan-Hou, 1997; Wang et al., 1989). In some bacterial genomes, more than one copy of the structural gene for merA, merB and merR were reported (Khaleque, 2017; Kiyono et al., 1997; Wang et al., 1989). In Synechocystis, only merA and merR genes exist and were distantly located in the genome and all other mer genes, such as merC, merD, merP, merT, merG, merE, and merB were not detected. It is important to note that not only in Synechocystis, but also in all other cyanobacterial species reported to date, merR and merA are distantly located in their genomes unlike in other bacterial species (Figure A2). However, the mechanism of gene regulation in response to mercury metal toxicity has not been reported in any cyanobacteria. Since homologs of the Slr0701 and Slr1849 appear to be well conserved among different aquatic cyanobacteria and relatively more similar to the metal-reducing bacterial species, it appears likely that the Slr0701 might be a regulatory gene in metal stress response.

3.2 Complete targeted inactivation of a gene, slr0701 coding for putative MerR

To elucidate the regulatory role of Slr0701, the ORF was inactivated as shown in the schematic representation (Figure 1a). The extent of replacement of wild-type copies of the slr0701 with that of disrupted copies of slr0701::sp' was confirmed by comparing the sizes of the amplified PCR products. When the genomic DNA from wild-type was used as a template for PCR amplification with specific primers (slr0701-F and slr0701-R), a PCR product of 1,340 bp representing the slr0701 ORF was amplified including the upstream and downstream regions. In contrast, when the genomic DNA isolated from slr0701-mutant were used as a template with the same set of primers, a 3,423 bp PCR product was amplified corresponding to the wild-type fragment of 1,340 bp including the inserted Ω-spectinomycin gene (sp') cassette of 2083 bp (Figure 1b).

3.3 slr0701-mutant is more sensitive to HgCl2 treatment compared to wild-type

Since, slr0701 codes for a putative MerR type transcriptional regulator, we investigated the effect of trace metals on the viability of slr0701-mutant cells in comparison with wild-type. Initially, to find the involvement of Slr0701 in metal resistance, we tested the growth of wild-type and slr0701-mutant strain in which slr0701 gene

**FIGURE 1** Strategy for disruption of the slr0701 gene in the genome of Synechocystis. (a) Schematic representation of the genotype of the slr0701 mutant. A 1,340 bp DNA fragment covering slr0701 gene and a 2083 bp DNA fragment containing Sp' gene are drawn to scale. The sp' and slr0701 ORFs are shown in the filled and open arrows, respectively. Small arrows indicate positions of PCR primers used to amplify the region and to verify gene replacement. FP: Forward primers; RP: Reverse primers. (b) PCR analysis of wild-type and slr0701-mutant cells with the primers indicated in (a). M represents 1-kb DNA ladder (Invitrogen™ Life Technologies; Cat. No. 15615-016)
was inactivated on a solid BG-11 media containing various metals as described in experimental procedures. Both wild-type and \textit{slr0701}-mutant cells revealed similar profiles of growth at different metal ions tested except the growth on the BG-11 solid plate containing HgCl\textsubscript{2} (data not shown). In order to confirm that the slow growth phenotype seen in \textit{slr0701}-mutant was due to HgCl\textsubscript{2}, we compared the growth profiles of wild-type, \textit{slr0701}-mutant and \textit{slr0701}\textsuperscript{+}-complemented cells in the absence and presence of HgCl\textsubscript{2} (Figure 2). All the three strains exhibited similar growth profiles when serially diluted cultures were spotted on a BG-11 agar plate without HgCl\textsubscript{2}. However, with increasing concentrations of HgCl\textsubscript{2} in the BG-11 solid agar plate, \textit{slr0701} mutant exhibited a slow growth phenotype compared to wild-type. At 500 nM of mercuric chloride, almost no growth of the mutant cells was observed in the spot area, at which 5-times diluted cells were deposited (Figure 2). The complementation of \textit{slr0701}-mutant cells with a functional Slr0701 expressed from pVZ321-slr0701 restored growth, and appeared similar to that of wild-type growth in the presence of mercuric chloride. We also tested the effect of HgCl\textsubscript{2} using the liquid cultures. We observed a significant difference in the growth profiles of wild-type and \textit{slr0701}-mutant cultures, when incubated with 500 nM HgCl\textsubscript{2}. However, at 750 nM concentration both the strains exhibited similar growth phenotypes (Figure A3). This indicated that mercury-induced expression of \textit{slr0701} gene seems to be crucial for \textit{Synechocystis} cells to tolerate inorganic mercury. The functional complementation of \textit{slr0701}-mutant cells by the \textit{slr0701} gene explains that the slow growth phenotype observed in the presence of mercury was due to the inactivation of \textit{slr0701}.

3.4 | Effect of HgCl\textsubscript{2} on photosynthetic activity due to mutation in \textit{slr0701}

We measured the PSII activity in the wild-type and \textit{slr0701}-mutant cells to analyze the extent of damage caused by mercury on photosynthetic performance due to inactivation of \textit{slr0701} gene. We used both oxygraph as well as relative Chl \textit{a} fluorescence kinetics to analyze PSII activity. At optimal growth conditions, no significant difference in the PSII activity was observed (from water to PBQ) between wild-type and \textit{slr0701}-mutant cells (Figure 3a). The PSII activity was decreased in both the WT and \textit{slr0701}-mutant cells when incubated with 500 nM HgCl\textsubscript{2} for 12 and 24 hr, respectively. There was about 40% loss observed in the PSII activity of wild-type cells at 24 hr of incubation with mercuric chloride, whereas the \textit{slr0701}-mutant cells showed 90% decrease in its PSII activity at the same point of incubation time (Figure 3a). Both wild-type and \textit{slr0701}-mutant cells exhibited the characteristic OJIP fluorescence transient upon illumination of dark-adapted cells. Minimal fluorescence (Fo) was observed to be almost constant with the increase in time of incubation with HgCl\textsubscript{2}, whereas variable fluorescence (Fv) and maximal fluorescence (Fm) was observed to be decreased significantly (data not shown). The Fv/Fm ratio, which reflects the activity of photosystem II was decreased in both the wild-type and \textit{slr0701}-mutant cells when incubated with 500 nM HgCl\textsubscript{2}. About 17% decrease in Fv/Fm ratio was observed in wild-type cells at 24 hr of incubation whereas the \textit{slr0701}-mutant cells showed about 48% decrease in Fv/Fm ratio at the same point of incubation time (Figure 3b). A decrease in the fluorescence yield due to mercury treatment can be attributed to inhibition of electron flow at the oxidising site of PS II (Lu & Vonshak, 2002). Such inhibition of electron flow, that is, reduced PSII activity due to incubation of cells with HgCl\textsubscript{2} was much higher in the \textit{slr0701}-mutant than wild-type cells. Data from both oxygraph and Chl \textit{a} fluorescence kinetics are consistent and a rapid decline in PSII activity under HgCl\textsubscript{2} stress in \textit{slr0701}-mutant clearly indicates severe damage in the photosynthetic machinery as compared to wild-type cells.

3.5 | Inactivation of \textit{slr0701} changes the mercury-induced expression \textit{mer} genes

The \textit{slr0701} and \textit{slr1849} mRNA levels were analyzed by real-time PCR in the wild-type and \textit{slr0701}-mutant cells. The expression of both \textit{slr0701} and \textit{slr1849} genes was upregulated during incubation of the cells with HgCl\textsubscript{2}. Incubation of wild-type cells with 500 nM

\textbf{FIGURE 2} \textit{slr0701} is involved in mercury resistance. Tolerance of wild-type, \textit{slr0701}-mutant and \textit{slr0701}\textsuperscript{+} strains to mercury was examined. As indicated serially diluted cultures were spotted onto solid BG-11 agar plate and also on HgCl\textsubscript{2} supplemented BG-11 agar plates with the indicated concentrations. Plates were photographed after 5 days of incubation under continuous illumination of 70 \textmu{E} m\textsuperscript{-2} s\textsuperscript{-1} of photons.
HgCl₂ resulted in upregulation of mRNA levels to 12.2 ± 0.27 of slr0701 and 1.5 ± 0.13 of slr1849 within 10 min. While the slr0701 mRNA levels were maintained to be high throughout the period of incubation, slr1849 mRNA levels were gradually increased to 3.81 ± 0.27 fold by 120 min (Figure 4a). In contrast, induced expression of neither slr0701 nor slr1849 was observed during the incubation of slr0701-mutant cells with HgCl₂. In fact, there was a little, but the significant downregulation of the expression of these genes was noted in the slr0701-mutant (Figure 4b). The results indicate that upon inactivation of slr0701 the expressions of its own gene and slr1849 gene were altered.

### 3.6 Slr0701 protein levels were enhanced due to the presence of HgCl₂

Slr0701 was induced and purified as a 6x-His–Slr0701 protein from *E. coli* Rosetta™ (DE3). The 6x-His–Slr0701 protein was resolved as a single protein band at ~17.3 kDa on SDS-PAGE. Under semi-denaturing condition (without 2-mercaptoethanol), Slr0701 protein appeared near 30 kDa, indicating its probable existence as a dimer (Figure 5a). The Slr0701 protein was detected in soluble fraction, but not in the insoluble fraction of the lysate (Data not shown). The anti-Slr0701 antibody recognized Slr0701 at a molecular mass nearly at 16 kDa in the wild-type cells. Slr0701 protein was not recognized in slr0701-mutant as anticipated. Figure 5b shows the immunodetection of Slr0701 in the wild-type. There was a gradual increase in the Slr0701 protein levels during the course of incubation with HgCl₂, which is in consistent with the upregulated gene expression of slr0701 (increase in slr0701 mRNA levels) as was analyzed by qRT-PCR.

### 3.7 Slr0701 binds to upstream of slr0701 and slr1849

As the expression of distantly located slr0701 and slr1849 genes were affected by the slr0701 inactivation, a common cis-regulatory binding element for Slr0701 protein is expected in the upstream of these genes. Using MEME motif discovery tool, we identified a common conserved cis-regulatory element not only in the upstream of slr1849 and slr0701 genes, but also in the upstream of their homologs from other cyanobacterial species (Figure 6a).
The binding of Slr0701 to the upstream of slr0701 and slr1849 with Slr0701 protein was studied using EMSA. Purified 6x-His–Slr0701 protein retarded the electrophoretic mobility of the DNA fragments having P<sub>slr0701</sub> and P<sub>slr1849</sub> cis-acting elements, and in-tensities of retarded DNA bands were increased with the amount of protein used for incubation (Figure 6b,c). Importantly, in the presence of 50 µM of HgCl<sub>2</sub>, the intensity of retarded DNA-protein complex was more compared to the intensity of the same in the absence of HgCl<sub>2</sub>. This result indicated that Hg<sup>2+</sup> enhances the binding of Slr0701 to its target DNA binding site. In order to confirm the specificity of binding to slr1849 and slr0701, we used an unrelated DNA upstream (sll1920) in the presence and in the absence of HgCl<sub>2</sub> (Figure A4). The 6x-His-Slr0701 did not bind to the unrelated-DNA fragment confirming its specificity to the upstream of slr1849 and slr0701.

### 3.8 Binding kinetics of Slr0701 in the upstream of slr0701

We further analyzed the interaction between the upstream of slr0701 and Slr0701 protein using SPR, as there was a significant increase in Slr0701 binding to P<sub>slr0701</sub> in the presence of HgCl<sub>2</sub>. We performed a binding analysis with varying concentrations of Slr0701 protein against the immobilized P<sub>slr0701</sub> DNA (Figure 7a). When the titration was repeated in the presence of 62 µM mercuric chloride, there was an increase in the response (RU). It is clear that association rate (k<sub>a</sub>) of the Slr0701 with immobilized DNA fragment was faster and the dissociation rate (k<sub>d</sub>) was observed to be slower in the presence of HgCl<sub>2</sub> (Figure 7b). The equilibrium dissociation constant (K<sub>d</sub>) evaluated from the kinetic traces showed that Slr0701 has nearly 1,000 times greater affinity to the promoter in the presence of HgCl<sub>2</sub> (Figure 7c). This complements with the EMSA results where mercury enhanced the binding between the Slr0701 protein and upstream of slr0701 (Figure 6b).

### 3.9 HgCl<sub>2</sub> induces transcription of mercury responsive genes

We monitored the activities of P<sub>slr0701</sub> and P<sub>slr1849</sub> promoters in the wild-type and slr0701—mutant cells in the presence and the absence of HgCl<sub>2</sub> in order to examine the role of HgCl<sub>2</sub> and Slr0701 in transcriptional activation of merA. The P<sub>slr0701</sub> and P<sub>slr1849</sub> promoters were independently fused to a luxAB reporter gene and introduced into both Synechocystis wild-type and slr0701-mutant
These modified strains were used for monitoring activities of the promoters. A weak luminescence signal was detected in the wild-type cells for P<sub>slr0701</sub> and also for P<sub>slr1849</sub> promoters even in the absence of HgCl<sub>2</sub> indicating their constitutive expression (Figure 8a). This result is consistent with previous reports where merR promoter found to be a weak promoter (Lund & Brown, 1989). The P<sub>slr0701</sub>‐lux<sub>AB</sub> in the wild‐type and in slr0701‐mutant showed 4.3 ± 2.4 and 1.3 ± 0.1 relative luminescence units (RLU) prior to the treatment with HgCl<sub>2</sub>. The P<sub>slr1849</sub>‐lux<sub>AB</sub> showed 4 ± 1.3 and 0.6 ± 0.3 RLUs prior to HgCl<sub>2</sub> treatment indicating that the basal activity of P<sub>slr0701</sub> and P<sub>slr1849</sub> promoters even in the absence of mercury. In the wild-type cells, after the addition of HgCl<sub>2</sub> there was an increase in luminescence signal suggesting that HgCl<sub>2</sub> enhances the expression of mer genes in Synechocystis (Figure 8a). Upon addition of HgCl<sub>2</sub> to wild-type cells, within 30 min the relative luminescence units were increased by four-fold (22.2 ± 2.0) and three-fold (10.3 ± 1.3) for P<sub>slr0701</sub> and P<sub>slr1849</sub>, respectively indicating that both the promoters could get activated immediately in the presence of HgCl<sub>2</sub> (Figure 8a). However, P<sub>slr1849</sub> activity was gradually increased as compared to the P<sub>slr0701</sub>, which is consistent with the qRT‐PCR results (see Figure 4a).

In contrast, the slr0701‐mutant cells harbouring plasmid DNA constructs having either P<sub>slr0701</sub> or P<sub>slr1849</sub> promoter fused to lux<sub>AB</sub> genes did not show any luminescence signal either in the absence or in the presence of HgCl<sub>2</sub> suggests that Slr0701 protein is necessary for mercury‐induced activation of its own gene expression as well as slr1849 transcription (Figure 8b).

### Circular dichroism analysis of structural changes in Slr0701

The structural changes in Slr0701 upon interaction with HgCl<sub>2</sub> were analyzed using CD. In the far‐UV region (200–250 nm), a gradual decrease in negative ellipticity was observed, when the protein Slr0701 was titrated with increasing concentration of HgCl<sub>2</sub>. This suggests the relaxation in the secondary structure of the protein upon binding to HgCl<sub>2</sub> (Figure 9a). The ellipticity changes followed at 222 nm upon addition of HgCl<sub>2</sub> in Figure 9b. The initial changes saturating at 100 µM of HgCl<sub>2</sub> might be attributed to the binding of HgCl<sub>2</sub> whereas further sharp decline in the negative ellipticity could arise from denaturation of the protein.

In addition, the tertiary structural change in Slr0701 protein was analyzed in the near‐UV region (250–300 nm) upon binding with P<sub>slr0701</sub> with and without HgCl<sub>2</sub>. As compared to DNA—Protein alone, there was a significant increase in ellipticity in the presence of HgCl<sub>2</sub> suggesting stronger binding of the protein with DNA in the presence of HgCl<sub>2</sub> (Figure 9c).
Cyanobacterial species are widespread in both aquatic and terrestrial habitat and often expose to various toxic metals. Although the genome sequence information revealed the presence of a putative MerR (a transcriptional regulator) and MerA (mercury reductase) like genes in cyanobacteria, the organization of mer genes and regulation has not been studied. Interestingly, merA and merR genes are distantly located from each other in the cyanobacterial genomes, unlike in other bacterial phyla which contain five different mer genes forming an operon (Figure A2). In Synechocystis, slr0701 and slr1849 code for a putative transcriptional regulator (MerR) and a mercury reductase (MerA), respectively. The Slr0701 and Slr1849 proteins show greater similarities to homologs from Geobacter than well-characterized homologs from Bacillus indicates that cyanobacterial mer genes might have horizontally transferred from these metal reducing species (Figure A1). Lack of several mer related genes and existence of only inorganic-mercury reductase, MerA, a putative MerR transcription factor and their unique gene organization in all cyanobacterial species prompted

**FIGURE 9** Circular dichroism analysis of Slr0701. (a) Far UV-CD spectra of Slr0701 (20 µM) measured in varying concentrations of HgCl₂. (b) The change in ellipticity of Slr0701 at 222 nm upon increasing the concentration of HgCl₂ representing secondary structural loss in the protein. (c) Near UV-CD spectra of Slr0701, P_{slr0701}, P_{slr0701}+Slr0701, and P_{slr0701}+Slr0701+HgCl₂

**FIGURE 10** Schematic representation of mercury-induced merA gene regulatory mechanism by a putative transcriptional regulator, Slr0701 in Synechocystis sp. PCC6803. (a) Slr0701 and Slr1849 genes are located far apart from each other in Synechocystis genome. Slr0701 is constitutively expressed and regulates its own gene expression. Slr0701 is transcribed from a weak promoter. (b) When cyanobacterial cells experience inorganic Hg²⁺, Slr0701-Hg²⁺ complex binds to cis-regulatory element located upstreams of its own ORF as well as slr1849 with greater affinity and leads to an induced expression. Thus Hg²⁺ bound Slr0701 activates the mercury-dependent expression of Slr1849. Slr1849 being an inorganic-mercury reductase converts Hg²⁺ to Hg⁰ and the volatile mercury is sent out of the cell

**4 | DISCUSSION**

Cyanobacterial species are widespread in both aquatic and terrestrial habitat and often expose to various toxic metals. Although the genome sequence information revealed the presence of a putative MerR (a transcriptional regulator) and MerA (mercury reductase) like genes in cyanobacteria, the organization of mer genes and regulation has not been studied. Interestingly, merA and merR genes are distantly located from each other in the cyanobacterial genomes, unlike in other bacterial phyla which contain five different mer genes forming an operon (Figure A2). In Synechocystis, slr0701 and slr1849 code for a putative transcriptional regulator (MerR) and a mercury reductase (MerA), respectively. The Slr0701 and Slr1849 proteins show greater similarities to homologs from Geobacter than well-characterized homologs from Bacillus indicates that cyanobacterial mer genes might have horizontally transferred from these metal reducing species (Figure A1). Lack of several mer related genes and existence of only inorganic-mercury reductase, MerA, a putative MerR transcription factor and their unique gene organization in all cyanobacterial species prompted
We inactivated the slr0701 that code for a putative MerR transcriptional regulator (Figure 1). Mercury is known to damage photosynthesis in cyanobacteria and higher plants (Bernier, Popovic, & Carpenter, 1993; Tangahu et al., 2011). We analyzed the extent of damage caused by HgCl$_2$ on photosystem II activity. The slr0701-mutant strain showed a severe reduction in photosynthetic activity and became relatively more sensitive to HgCl$_2$ than wild-type cells emphasizing its role in Hg$^{2+}$ detoxification (Figures 2 and 3). Rapid and gradual upregulation of slr0701 and slr1849 mRNA levels were observed in the presence of HgCl$_2$ (Figure 4).

This induced gene expression was observed to be due to the binding of Slr0701 protein to the cis-regulatory element in the upstream of its own ORF and slr1849 followed by transcriptional activation (Figures 5, 6 and 10). Moreover, the cis-regulatory element is found to be well conserved among most cyanobacterial mer homologs. In the presence of HgCl$_2$, Slr0701 protein showed faster association and slower dissociation constants to the target DNA binding element (Figures 7 and 10). The slr0701 and slr1849 are constitutively expressed. Upon addition of HgCl$_2$, mercury-bound Slr0701 enhances the transcription of these genes (Figures 8 and 10). In addition HgCl$_2$ induced conformational changes in the protein and enhanced affinity to DNA leads to upregulation of gene expression (Figures 9 and 10). In the slr0701-mutant, the mercury-induced expression of mercury reductase, merA was not observed due to inactivation of slr0701 gene. Hence, the slr0701-mutant became more sensitive to Hg$^{2+}$ than wild-type. Cyanobacteria contain only a well-conserved transcriptional regulator, MerR and a mercury reductase, MerA. They are found to be distantly located in the genome which is a unique feature when compared to other bacterial phyla. However, MerR could still regulate the mercury-dependent expression of MerA. The mechanism of gene regulation involving MerR is well-conserved among all bacterial phyla.

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**CONFLICT OF INTERESTS**

None declared.

**AUTHORS CONTRIBUTION**

D.K.S performed majority of the experiments and wrote the paper. L.B generated the mutant strain. K.T.N and D.K.S performed CD spectral experiments together. N.P.P analyzed the CD and SPR results and edited the paper. J.S.S.P conceived the idea for the project and wrote the paper with D.K.S.

**ETHICS STATEMENT**

None required.

**DATA ACCESSIBILITY**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Phylogenetic relationship of Slr0701 and Slr1849 with their closely related orthologs. The complete amino acids sequences of Slr0701 and Slr1849 and their orthologs were taken, phylogenetic relationship was deduced. All investigated cyanobacterial and bacterial strains are presented in the figure. The well-studied *Bacillus megaterium*, MerR and MerA with five other proteins from different organisms more closely related to these Mer proteins were also included. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (500 replicates) is shown next to the branches. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances are given in the number of amino acid substitutions per site as indicated by the scale bar representing 0.7 for *slr0701* (a) and 0.3 for *slr1849* (b) substitutions per amino acid.

**FIGURE A1**
FIGURE A2 Comparison of mer operon organization in different bacterial and cyanobacterial species. merR (●), Transcriptional regulator; merG (●), merT (●), merP (●), merC (●), merE (●), periplasmic proteins; merA (●), mercury reductase; merB (●), organo-mercurial lyase and merD (●) alternative transcriptional regulator. The reading frames are shown as arrows to indicate ORFs. The mer genes shown below the bold dash line belong to cyanobacterial species. (//) indicates that the genes are located distantly from each other in the genome.

FIGURE A3 Effect of HgCl₂ on growth of Synechocystis wild-type and slr0701-mutant strains. For monitoring the growth in the presence of HgCl₂, cultures were initially allowed to grow till 16 hr as described in methods, and then HgCl₂ was added to a final concentration of 250 nm, 500 nm and 750 nm to the respective culture tubes. Optical density at 730 nm was measured at regular intervals. Filled symbols represent growth of wild-type Synechocystis cultures and open symbols represent slr0701-mutant cultures. The mean of three experimental values is presented with standard error.

| MerR (µg) | 0.5 | 1.0 | 0.5 | 1.0 |
|-----------|-----|-----|-----|-----|
| P_slr1920 (80 nmoles) | +   | +   | +   | +   |
| HgCl₂ (50 µM) | −   | −   | −   | +   |

FIGURE A4 Binding of Slr0701 protein to the upstream DNA fragment of slr1920 ORF. A gel mobility shift assay was performed with 0.5 µg of 6x-His-Slr0701 protein and 80 nmoles of upstream DNA fragment of slr1920 in a binding reaction. A 201 bp DNA upstream of slr1920 starting from −1 to −201 bp with respect to the translation start site was used in the binding reaction.