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# A LexA-related protein regulates redox-sensitive expression of the cyanobacterial RNA helicase, *crhR*

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# ABSTRACT

Expression of the cyanobacterial DEAD-box RNA helicase, crhR, is regulated in response to conditions, which elicit reduction of the photosynthetic electron transport chain. A combination of electrophoretic mobility shift assay (EMSA), DNA affinity chromatography and mass spectrometry identified that a LexA-related protein binds specifically to the crhR gene. Transcript analysis indicates that lexA and crhR are divergently expressed, with lexA and crhR transcripts accumulating differentially under conditions, which respectively oxidize and reduce the electron transport chain. In addition, expression of the Synechocystis lexA gene is not DNA damage inducible and its amino acid sequence lacks two of three residues required for activity of prototypical LexA proteins, which repress expression of DNA repair genes in a range of prokaryotes. A direct effect of recombinant LexA protein on crhR expression was confirmed from the observation that LexA reduces crhR expression in a linear manner in an in vitro transcription/translation assay. The results indicate that the Synechocystis LexA-related protein functions as a regulator of redox-responsive crhR gene expression, and not DNA damage repair genes.

# INTRODUCTION

The ability to adapt to a dynamic light environment is crucial for the survival of photosynthetic organisms and includes both short- and long-term responses. Light sensing occurs either via direct mechanisms involving photoreceptor proteins, or indirectly through light-driven changes in the redox status of the electron transport chain between  $Q_A$  in photosystem II and  $Q_O$  in cytochrome  $b_6 f$  (1–4). Electron carriers in this region of the inter-photosystem electron transport chain perform essential roles in redox-sensing in higher plant chloroplasts, regulating expression of nuclearand chloroplast-encoded genes involved in photosynthesis (5–9). For example, a direct link between the redox poise of plastoquinone and chloroplast gene expression has been shown for the *psbA* and *psaAB* genes, allowing rapid cellular response to the light environment via sensing of the redox status of the electron transport chain (7). In contrast, the factors responsible for transduction of the electron transport redox poise to transcription regulation remain poorly characterized. Possible transducers identified in spinach chloroplasts include an unidentified 31 kDa dimeric protein shown to bind the *psaAB* promoter (10) and the TSP9 thylakoid-associated protein. Redox-mediated phosphorylation of TSP9 on three threonine residues releases the protein from the thylakoid membrane potentially allowing it to play a role as a signaling factor responsible for transducing plastoquinone redox poise to gene expression (11).

In prokaryotic cyanobacteria, the redox status of the electron transport chain carriers also regulates expression of a limited number of photosynthetic and non-photosynthetic genes. Expression of the RNA helicase, crhR (1), glutamine synthetase, *glnA* (12), PII protein, *glnB* (13),  $\alpha$  and  $\beta$  subunits of phycocyanin, cpcBA (14), photosystem proteins (14-16) and a transcriptional regulator (17) has been attributed to the redox poise of plastoquinone and/or cytochrome  $b_6 f$ . Members of the signal transduction pathway(s) associated with sensing and transducing changes in redox status to the transcriptional machinery also remain to be identified in cyanobacteria. Proposed mechanisms in cyanobacteria include a redox-responsive two-component signal transduction pathway (16) and the redox-sensitive transcriptional regulators, NtcA and NblS (17,18). A potential sensor is the membraneassociated sensor histidine kinase NbIS, which responds and controls photosynthesis-related gene expression in response to high light and nutrient stress (18). NblS contains a redox-sensing PAS domain potentially involved in the sensing and transduction of high light/nutrient stress induced changes in photosynthetic or cellular redox poise to as yet uncharacterized transcriptional regulators (18). While a number of redox-sensitive transcriptional regulators have been described in prokaryotic systems, the physiological electron donors have not been identified (19-23). NtcA is one transcriptional regulator whose activity has been shown to involve a complex interaction between cellular nitrogen levels, thiol group redox poise and photosynthetic electron flow although not directly correlated with the redox state of the plastoquinone pool (17). NtcA controls expression of

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This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/ by-nc/2.0/uk/) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. genes involved in nitrogen acquisition, repressing *gifA* and *gifB* and activating expression of *glnA*, *glnN* and *glnB* in response to both nitrogen levels and electron transport (24).

Expression of the Synechocystis DEAD-box RNA helicase, crhR (Cyanobacterial RNA Helicase-Redox), is also regulated by the redox poise of the electron transport chain (1). crhR transcripts accumulate when the electron transport chain is reduced, either from light-driven electron flow or respiratory electron flow generated by the metabolism of exogenously supplied glucose. In contrast, a reduction in electron flow, leading to oxidation of the electron transport chain, decreases crhR transcript accumulation. These results are corroborated by results obtained using electron transport inhibitors or alteration of light quality which alter *crhR* induction, confirming redox-regulated expression and identifying the redox poise of the electron transport chain between Q<sub>A</sub> in photosystem II and  $Q_0$  in the cytochrome  $b_6 f$  complex, as the potential sensor for redox-dependent regulation (1). Biochemically, CrhR exhibits enzymatic activities characteristic of RNA helicases, including RNA-dependent ATPase activity and ATP-stimulated RNA unwinding (25). In addition, CrhR also possesses ATP-dependent RNA annealing activity (26). Thus, CrhR has been proposed to regulate gene expression at the translational level through its ability to rearrange RNA secondary structures of RNA substrates, potentially of other redox-regulated gene transcripts (1,26).

As an initial step to elucidate upstream factors involved in the redox-regulated expression of the crhR gene, we have identified a LexA-related protein that controls *crhR* transcript accumulation. Treatments known to enhance *crhR* transcript accumulation reduce lexA levels and vice versa. A direct effect of the recombinant His-tagged LexA protein (rLexA) on crhR expression was confirmed from the observation that rLexA reduces crhR expression in an in vitro transcription/translation assay. LexA thus appears to function as a repressor of *crhR* transcription when *crhR* is not required i.e. under conditions which oxidize the electron transport chain. In concurrence with this conclusion, expression of the Synechocystis lexA gene is not DNA damage inducible and its amino acid sequence lacks two of three residues required for the self-cleavage activity of prototypical LexA proteins (27). We discuss the potential significance of a LexA-related repressor in the regulation of redox-responsive gene expression and, consequently, the implications of this novel role performed by LexA in Synechocystis.

#### MATERIALS AND METHODS

#### Bacterial strains and growth conditions

Synechocystis sp. strain PCC 6803 was maintained at 30°C on BG-11 agar (28) solidified with 1% (w/v) Bacto-Agar (Difco Laboratories, Detroit, MI) and grown photoautotrophically at 30°C under continuous illumination at a constant intensity of 150  $\mu$ mol photons m<sup>-2</sup> s<sup>-1</sup>. Liquid BG-11 cultures were aerated by shaking at 150 r.p.m. and continuous bubbling with humidified air. Dark conditions were created by wrapping the flasks in aluminum foil. Glucose (5 mM) was added where indicated. To induce DNA damage, cells were exposed to short-wave ultraviolet light (UV-C; 254 nm) at a dose rate of 150, 300 or 600 J/m<sup>2</sup> using a

XL-1000 UV crosslinker (Spectronics Corporation) and subsequently incubated in the dark for 1 h prior to harvesting.

*Escherichia coli* strains DH5 $\alpha$  and JM109 were used for propagation and protein expression of plasmid constructs, respectively. Cultures were grown in Luria–Bertani (LB) medium at 37°C and aerated by shaking at 200 r.p.m. Ampicillin (100 µg/ml) was added where appropriate.

#### **Plasmid constructs**

A deletion series within the *crhR* promoter/open reading frame (ORF) was created from a 3 kb EcoRI fragment encompassing the *crhR* promoter, 5'-untranslated region (5'-UTR) and the *crhR* ORF (1). Plasmid DNA was digested with NotI and SacI, and a deletion series produced using the Erase-A-Base Kit (Promega) according to the manufacturer's instructions. Two additional deletion constructs were created by restriction enzyme digestion. SpeI removed a 328 bp fragment to construct KC+125. The KC+219 construct was created by EcoRI / XmnI digestion to liberate a 2.6 kb fragment containing the *crhR* ORF downstream of +219 but lacking the promoter region. This fragment was blunt end ligated into EcoRV digested pBluescript KS+ (Stratagene).

The lexA gene was amplified by PCR to generate an in frame translational fusion in the pRSETB plasmid vector (Invitrogen). The lexA insert DNA was generated using primers LPF-27 (5'-ACTGGTGGATCCGAACCTCTCACCC-GAGCC-3') and LPF-2 (5'-GAAACAAAAAGCTTAGGA-CG-3') and Synechocystis chromosomal DNA as template. PCR were performed in a volume of 50 µl, containing 300 nM primer and 0.8 U of Expand High Fidelity enzyme mix (Roche) according to the manufacturer's instructions. The PCR program consists of 30 cycles of 1 min denaturation at 95°C, 30 s annealing at 55°C and 1 min extension at 68°C; and terminated with 4 min extension at 68°C. Purified DNA was digested with BamHI and HindIII (restriction sites underlined) and cloned into BgIII and HindIII digested plasmid DNA. The resulting plasmid pLexA expresses a recombinant LexA polypeptide (rLexA) containing a 37 amino acid residue N-terminal HIS tag. pLexA was sequenced to confirm successful in frame insertion of the lexA gene.

### Generation of promoter fragments

Promoter fragments were PCR amplified from the deletion plasmids using the M13 forward primer (5'-GTAAAAC-GACGGCCAGT-3') and GWO-45 (5'-AAGCCAATGTCG-GCCAAGAG-3') (Figure 1A). PCR were performed as described above using an annealing temperature of 45°C. BssHII / BssSI digestion was used to generate fragments corresponding to KC+125 and KC+219. DNA fragments were purified from 1× TAE agarose gels using GENECLEAN<sup>®</sup>II (BIO 101).

#### DNA affinity column purification

To purify proteins binding to the *crhR* promoter region, µMACS Streptavidin magnetic separation was performed (Miltenyi Biotec). *Synechocystis* cultures (300 ml) were harvested at mid-log phase and resuspended in 2 ml cyanobacterial protein extract buffer [20 mM Tris–HCl (pH 8), 10 mM NaCl, 1 mM EDTA (pH 8) and 5 mM DTT] containing protease inhibitors (Complete Mini Protease Inhibitor Cocktail,



Figure 1. (A) crhR nested deletion series. DNA was deleted by directional digestion from the SacI site using Exonuclease III. Deleted clones are designated by their start site relative to the transcriptional start indicated as +1. DNA fragments corresponding to each deletion were generated by PCR using the M13 forward (FP) and GWO-45 primers, expect KC+125 and KC+219 which were produced by restriction digestion (KC+125: SpeI/BssS1; KC+219: XmnI/BssS1). Plasmid and crhR insert sequences are indicated by thick and thin solid lines, respectively. Scale 50 bp = 1 cm. (**B** and **C**) EMSA identification of the protein-binding region in the crhR gene. (B) Localization of the protein-binding region. <sup>32</sup>P-end-labeled DNA targets were incubated either alone (–) or with 30  $\mu$ g *Synechocystis* soluble protein extract (+). (C) Competition assays. KC-179 <sup>32</sup>P-end-labeled target DNA, containing the entire crhR promoter, was incubated with no protein or 30 µg Synechocystis soluble protein extract. Increasing amounts (0-3.0 pmol) of either specific competitor DNA (unlabeled KC-179 fragment; upper panel) or non-specific competitor DNA (unlabeled 262 bp EcoRV / PvuII fragment of pBluescript KS+; lower panel) were included in the binding reaction to determine the specificity of the protein-DNA interaction.

Roche). Cells were lysed by eight cycles of sonication for 30 s followed by 30 s cooling in an ice-water bath. Lysed cells were clarified by centrifugation and the supernatant retained. Proteins were quantified by the Bradford assay (BioRad) using BSA as a standard. Binding reactions were performed in  $1\times$  electrophoretic mobility shift assay (EMSA) buffer [10 mM Tris (pH 7.5), 50 mM NaCl, 1 mM EDTA, 5% glycerol and 1 mM DTT] containing 50 µg poly(dI-dC), 28 µg biotinylated target DNA and 6.8 mg soluble protein extract. Biotinylated target DNA was prepared by PCR amplification using the primers KCO-4 (5'-GTAA-AACGACGGCCAAGT-3') and biotinylated KCO-5 (5'-AAG-CCAATGTCGGCCAAGAG-3') and deletion plasmid DNA

as template. PCR were performed as described above using an annealing temperature of 55°C. The binding reactions were gently shaken at 4°C for 80 min, 100 µl superparamagnetic µMACS MicroBeads conjugated to streptavidin (µMACS Streptavidin kit, Miltenyi Biotec) added and incubation continued for 15 min. The µMACS column was prepared by rinsing consecutively with protein application buffer and 1× EMSA buffer. The binding reaction was applied to the µMACS column within the magnetic field of the µMACS separator, and washed consecutively with steps of increasing salt stringency (0.1 to 1 M KCl). Eluted proteins were concentrated by TCA precipitation, separated on a 10% (w/v) SDS–polyacrylamide gel and visualized by silver staining (BioRad).

Polypeptides of interest were identified by in-gel tryptic digestion and LC/MS/MS of the resulting peptides at the Institute for Biomolecular Design (University of Alberta). Generated LC/MS/MS data were used as queries for Mascot Daemon (Matrix Science, UK) searches of the National Center for Biotechnology Information (NCBI) non-redundant databases. A protein score greater than 73 following Mascot searches was considered significant.

## EMSA

EMSA were performed using Synechocystis soluble protein extract, E.coli soluble protein extract or recombinant LexA (rLexA) protein, and the indicated PCR-generated promoter fragments end-labeled with  $[\gamma^{-32P}]$ ATP and T4 polynucleotide kinase (New England Biolabs). Binding reactions were performed for 20 min at 37°C in 1× EMSA buffer, 1 µg poly(dI-dC), 2000 c.p.m. end-labeled DNA (~0.006 pmol) and the indicated protein concentration in a final volume of 20 µl. Reaction products were separated on a 5% TBE nondenaturing polyacrylamide gel and subjected to autoradiography. Two non-specific competitor DNAs were prepared to control for non-specific protein-binding. A vector control target was a 262 bp EcoRV / PvuII fragment containing the pBluescript KS+ multiple cloning site and a 321 bp internal fragment of the Synechocystis lexA gene, PCR amplified using primers LPF-4 (5'-ATTTGCGTTCTCCGGCC-3') and LPF-5 (5'-CTTCGATTTCCTCTTCTC-3') using an annealing temperature of 45°C as described above.

#### **Recombinant LexA expression and purification**

E.coli JM109:pLexA cultures were grown at 37°C to  $OD_{600} = 0.6$ , and LexA expression induced by addition of isopropyl-B-D-thiogalactopyranoside (IPTG) (1.0 mM) and phage (M13/T7 DE3, 5 p.f.u./cell, Invitrogen). After induction for 3 h at 37°C, harvested cells were resuspended in 1/10 volume lysis buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub>, 300 mM NaCl and 10 mM imidazole), lysed by sonication  $(6 \times 30 \text{ s inter-}$ vals), and clarified by centrifugation. The supernatant was loaded onto a Ni-NTA column (Qiagen) and incubated with gentle shaking for 60 min at 4°C. The column was washed consecutively with wash buffer (50 mM NaH<sub>2</sub>PO<sub>4</sub> and 300 mM NaCl) containing increasing amounts of imidazole (20 to 50 mM), with bound rLexA eluting in buffer containing 250 mM imidazole. Imidazole was removed from the eluted rLexA buffer by dialysis against lysis buffer lacking imidazole.

#### Northern analysis

Total RNA was isolated from *Synechocystis* by mechanical lysis, separated on a 1.2% formaldehyde gel, and transferred to a nylon membrane (Hybond N<sup>+</sup>) as described previously (29). Blots were hybridized overnight at 65°C with the appropriate probe in aqueous buffer (5× SSPE, 5× Denhardts, 0.5% SDS) and washed for 10 min at 65°C once in 1× SSPE, 0.1% SDS and once in 0.1× SSPE, 0.1% SDS. *lexA* and *crhR* DNA fragments were randomly labeled with [ $\alpha$ -<sup>32</sup>P]dCTP using random hexanucleotide primers (Roche). The probes correspond to: *lexA*, a 750 bp BgIII / HindIII fragment encompassing the entire ORF; *crhR*, a 784 bp internal BstEII fragment; and *recA*, a 1091 bp riboprobe encompassing the ORF labeled with [ $\alpha$ -<sup>32</sup>P]UTP (Promega). Membranes were stripped by incubation in boiling 0.1% SDS and probed with the *Synechocystis rnaseP* gene as a control for RNA loading (30).

#### In vitro transcription/translation assays

In vitro transcription and translation assays were performed using the Promega E.coli S30 extract system for circular DNA in a final reaction volume of 25 µl. The plasmids pCrhR (IV) and pWM3-2 (29) were used for in vitro expression of the crhR and crhC genes, respectively. pCrhR (IV) was prepared by ligating a 2.2 kb BamHI / EcoRI fragment of CS0096-9 (1) into pBluescript KS+ to remove downstream sequences encoding the argC gene. The pCrhR (IV) and pWM3-2 plasmids contain 2.2 and 2.4 kb inserts, respectively encoding the promoters, ORFs, and 5'- and 3'-UTRs of crhR and *crhC*, respectively. Reactions were performed according to manufacturer's instructions using 1 µg plasmid DNA, corresponding to 0.29 and 0.28 pmol DNA for pCrhR (IV) and pWM3-2, respectively. Reaction products were separated on a 10% (w/v) polyacrylamide gel and subjected to autoradiography. Binding reactions containing rLexA were performed according to manufacturer's instructions with an initial 5 min incubation to allow protein-binding to the crhR gene. Control reactions were performed to determine the effect of rLexA on *crhR* expression in the presence of BSA and expression of an unrelated RNA helicase, crhC (30), from its own promoter.

#### RESULTS

# Promoter deletion series delineates the protein-binding site within the *crhR* gene

EMSA assays were performed using nine plasmid constructs containing deletions of the *crhR* promoter (Figure 1A) to delineate the protein-binding region. Intact *crhR* promoter (KC-179) and deletions up to position +77 of the *crhR* transcript (KC+77) exhibited decreased mobility on a native PAGE gel upon incubation with *Synechocystis* protein extract (Figure 1B, lanes 1–14). The KC+125 DNA target, deleted to +125 of the transcript, exhibited a reduced amount of shift (Figure 1B, lanes 15 and 16), while deletion to +219 completely abolished the mobility shift (Figure 1B, lanes 17 and 18). Together, these results indicate that the protein-binding site is located downstream of the translational start codon (+110) in the region of DNA surrounding an SpeI site (+125). Sequence specificity of binding was shown by competition assays in the presence of increasing amounts of either specific or non-specific competitor DNA (Figure 1C). Addition of specific competitor DNA (KC-179) progressively challenged formation of the shifted complex (Figure 1C, Specific competitor). Conversely, inclusion of a similar sized fragment containing the pBluescript  $KS^+$  multiple cloning site had no effect on the mobility shift (Figure 1C, non-specific competitor). Taken together, these results indicate that at least one soluble *Synechocystis* protein interacts with the *crhR* gene in a sequence-specific manner.

# Synechocystis LexA-related protein binds within the crhR ORF

To identify the protein responsible for altered mobility of the crhR gene, DNA affinity column chromatography was performed using light-grown Synechocystis soluble protein extracts and biotinylated KC+5 DNA (239 bp). A single polypeptide with an apparent molecular weight of 28 kDa was recovered in the high stringency 1 M KCl elution (Figure 2A). The single significant hit (score 92) identified by in-gel tryptic digestion and LC/MS/MS corresponded to the Synechocystis gene sll1626, which has been annotated as encoding the transcriptional repressor LexA (http:// www.kazusa.or.jp/cyanobase/). Analysis of the deduced Synechocystis LexA amino acid sequence revealed that the sequence lacks the Ala-Gly self-cleavage site and the serine of the Ser-Lys dyad active site present in E.coli LexA, both of which are required for LexA self-cleavage [Figure 2B; (31)]. Furthermore, an SOS-like box, similar to those identified as LexA binding sites in E.coli (32,33), Bacillus subtilis (34) and Mycobacterium tuberculosis (35) could not be identified within the upstream sequence of either lexA or crhR (data not shown). However, a sequence related to the putative cyanobacterial SOS box (36), matching 7 of 9 essential residues with required spacing between essential residues, was identified within the protein-binding domain in crhR (Figure 2C). This sequence includes the SpeI site, possibly explaining the reduced shift observed with the SpeI generated KC+125 fragment (Figure 1B).

# *Synechocystis lexA*, *crhR* and *recA* transcript accumulation

crhR transcript accumulation is regulated by the redox poise of the plastoquinone pool with treatments leading to reduction of plastoquinone correlating with an increase in *crhR* transcript accumulation, whereas conditions that lead to the oxidation of the plastoquinone pool result in decreased crhR accumulation (1). Northern analysis was therefore performed to determine the relationship between lexA and crhR transcript accumulation under varying redox conditions (Figure 3A). Growth in the light (Figure 3A, lane 1), conditions favoring crhR transcript accumulation, correlate with reduced levels of lexA transcript. Conversely, growth in the dark (Figure 3A, lane 2) reduces crhR while enhancing lexA transcript accumulation. The addition of glucose (5 mM) to light-grown cells enhanced crhR and lexA transcript accumulation (Figure 3A, lane 3). crhR expression was significantly induced in response to cold stress (20°C; Figure 3A, lane 4), concomitant with the complete repression of lexA transcript accumulation. The data indicates



Figure 2. Isolation and characterization of a crhR regulatory protein by affinity chromatography and LC/MS/MS. (A) A 28 kDa polypeptide interacts with the crhR ORF. A single polypeptide was isolated by DNA affinity chromatography using KC+5 as the target. Non-specifically bound proteins were removed by increasing KCl washes. Silver staining of eluted proteins separated by a 10% SDS-PAGE reveals a single polypeptide in the 1 M KCl elution. LC/MS/MS identified this polypeptide as the Synechocystis gene annotated as encoding LexA. Lane 1, 250 mM KCl wash; lane 2, low molecular weight standards (BioRad); lane 3, 1 M KCl wash. (B) Amino acid sequence analysis of the Synechocystis (gray) and E.coli (black) LexA proteins. Residues essential for E.coli LexA self-cleavage in response to DNA damage are indicated as follows: \* = Ala-Gly self-cleavage site; + = Ser and # = Lys indicate the Ser-Lys dyad active site. (C) Alignment of the putative LexA binding region of the crhR gene (SpeI site is underlined) with the consensus cyanobacterial LexA binding sequence (32). Conserved residues are bolded. The LexA binding sequence within the crhR matches at 7 of 9 conserved residues with appropriate spacing.

differential regulation of *crhR* and *lexA* expression in response to alterations in the redox status of the electron transport chain, implying that LexA functions as a regulator of *crhR* expression.

LexA association with the *E.coli* SOS response and repression of DNA damage repair gene expression warranted comparative analysis of *Synechocystis lexA*, *recA* and *crhR* expression in response to DNA damage. DNA damage was induced by UV-irradiation, and the resulting expression patterns were examined by northern analysis (Figure 3B). In contrast to *E.coli*, following UV-irradiation neither *Synechocystis lexA* nor *recA* expression (Figure 3B, lanes 2–4) was induced above basal levels detected in dark grown cells (Figure 3B, lane 1). In fact, in contrast to other prokaryotic systems, *recA* was expressed at very low levels



**Figure 3.** Transcript analysis. Total RNA (30 µg) was isolated from *Synechocystis* cells grown as indicated. RNA was separated on a 1.2% formaldehyde agarose gel, transferred to Hybond N<sup>+</sup> and hybridized with the indicated <sup>32</sup>P-labeled probe. (A) *lexA*, *crhR* and *rnaseP* transcript accumulation following incubation in the light. Lane 1, 3 h light; lane 2, 3 h dark; lane 3, light plus 5 mM glucose; lane 4, cold stress for 3 h (20°C). (B) *recA*, *lexA* and *rnaseP* transcript accumulation in response to increasing levels of UV-irradiation. Lane 1, 1 h dark; lane 2, UV irradiated with 150; lane 3, 300; lane 4, 600 J/m<sup>2</sup> followed by a 1 h incubation in the dark.

under all conditions tested, requiring riboprobe detection and extended exposure times. *lexA* transcript accumulation was also not altered by DNA damage-induced by mitomycin C (data not shown). Similarly, expression of *crhR* was not UV-inducible (data not shown); rather, it followed the expected decrease in transcription that occurs in wild-type cells in the dark (1). The lack of induction of the *Synechocystis recA*, *lexA* and *crhR* genes following DNA damage suggests these gene products are not required during the cellular response to DNA damage.

#### Synechocystis LexA interacts with the crhR gene

Recombinant His-tagged LexA (rLexA) was purified to near homogeneity and used to test interaction with the KC+5 crhR promoter fragment (Figure 4). KC+5 crhR promoter DNA mobility was reduced by incubation with total Synechocystis protein extracts (Figure 4A, lane 8 versus lane 1). Mobility of the KC+5 DNA target was also altered by incubation with purified rLexA, with alteration of target DNA mobility exhibiting rLexA concentration dependence (Figure 4A, lanes 2-7). E.coli soluble protein extracts did not alter crhR DNA target mobility, indicating that E.coli proteins do not bind the crhR gene (Figure 4A, lane 9). Sequence-specific binding was demonstrated by competition assays in the presence of increasing concentrations of either specific or nonspecific competitor DNA. Addition of unlabeled specific competitor (KC+5) challenged formation of the shifted complex at all concentrations tested (Figure 4B, lanes 2-5), with addition of ≥50-fold excess of unlabeled target abolishing shift of the DNA target. In contrast, incubation with nonspecific competitor DNA, an internal lexA fragment similar in size to the specific competitor, did not significantly alter mobility shift at comparable concentrations (Figure 4B,





Figure 4. LexA-related protein-binding analysis. EMSA using recombinant LexA (rLexA) were performed to confirm interactions between LexA and the *crhR* gene. (A) rLexA concentration curve. Increasing concentrations of rLexA were incubated with <sup>32</sup>P-labeled KC+5. As controls, rLexA was also incubated with *Synechocystis* (lane 8) and *E.coli* (lane 9) soluble protein <sup>83</sup>P-labeled KC+5 and the indicated fold excess of either specific competitor DNA (unlabeled KC+5; lanes 1–5) or non-specific competitor DNA (internal *lexA* fragment; lanes 6–10).

lanes 7–10). Together, the results indicate that recombinant LexA interacts with the crhR gene in a sequence-specific manner.

#### LexA represses crhR gene expression in vitro

An *in vitro* transcription and translation system was used to confirm LexA regulation of crhR gene expression from its native promoter. As shown in Figure 5A, CrhR protein accumulation decreased in response to increasing rLexA concentration. Quantification of these results indicated that the rLexA inhibition of CrhR expression was linear with respect to rLexA concentration (Figure 5B). The specificity of repression was demonstrated by the lack of change in the levels of the plasmid-encoded  $\beta$ -lactamase protein, a non-LexA regulated protein. Similarly, crhR expression was unaffected by incubation in the presence of 1 pmol BSA (Figure 5C), a protein concentration at which rLexA significantly altered crhR expression. Furthermore, in vitro transcription and translation of a second cyanobacterial RNA helicase, crhC (29,30), was also unaffected by 1 pmol rLexA (Figure 5C). Together, these results indicate that LexA specifically regulates *crhR* transcription in a negative fashion.

#### DISCUSSION

We have begun investigation of the signal transduction pathway transducing the redox poise of the electron transport system to the transcription apparatus in cyanobacteria. In this



Figure 5. In vitro transcription/translation. In vitro reactions were used to investigate the nature of the LexA regulatory relationship with the crhR gene. (A) CrhR protein (55 kDa) accumulation in the presence of increasing rLexA concentration. A plasmid (pCrhR IV) containing the complete crhR gene, including 289 bp upstream of the translation start, was incubated in a transcription/translation mixture in the presence of increasing rLexA (0-40 nM). (B) Quantification of rLexA effect on CrhR expression. CrhR expression in the presence of increasing rLexA concentration was quantified from triplicate, independent replicates similar to the data shown in (A) using ImageQuant (Molecular Dynamics). Standard deviations from the means are shown. (C) Specificity of rLexA regulation. As control reactions, the transcription/translation efficiency of CrhR was evaluated in the presence of a non-specific protein, BSA (1 pmol). In addition, accumulation of the temperature-regulated cyanobacterial RNA helicase, CrhC (24,25) was determined in the presence and absence of rLexA (1 pmol). The relative level of CrhC expression is shown below the figure, as determined using ImageQuant.

paper we show that a LexA-related protein regulates expression of the redox-responsive RNA helicase, *crhR*. This identification implies a novel function for LexA in *Synechocystis*, a conclusion consistent with previous studies suggesting that LexA may regulate expression of carbon metabolism and bidirectional hydrogenase genes in *Synechocystis* (37–39).

The LexA binding site is located downstream of the crhR transcription start site, requiring sequences surrounding +125 of the crhR transcript. This localization suggests a regulatory mechanism for the Synechocystis LexA-related protein that differs from LexA regulation of DNA damage inducible genes in E.coli and other bacteria, where the LexA binding site (SOS box) surrounds the transcriptional start (32,33). Similar regulatory element arrangements, where transcription factors bind downstream of the transcription start site, have been observed in the cyanobacterium Synechococcus sp. strain PCC 7942 (40). In fact, the light responsive transcription of the psbA and psbD gene families in Synechococcus requires enhancer elements located downstream of the transcription start (40-42). The LexA DNA binding site within the crhR ORF is therefore consistent with regulatory proteinbinding sites localized in other genes whose expression is known to be regulated by either light or redox signals. Unfortunately, the DNA binding proteins interacting with these other sites remain to be identified.

A combination of DNA affinity chromatography and mass spectrometry identified the protein interacting with the crhR gene as being related to LexA. Northern blot analysis showed that Synechocystis lexA transcripts accumulate when cells are grown under conditions correlating with the repression of crhR accumulation. Based on these results, it appears that LexA functions as a negative regulator of crhR expression. Negative regulation was confirmed using an in vitro transcription/translation assay, which demonstrated that LexA binding interferes with *crhR* expression possibly through interference with promoter recognition and/or transcription initiation. Based on these results, it appears that Synechocystis LexA functions as a repressor of crhR expression. LexA activity is well studied in E.coli and other prokaryotes where it regulates expression of  $\sim 20$  unlinked genes associated with DNA damage repair, the SOS regulon, which include recA and lexA (27). Derepression occurs following DNA damage, and requires RecA-stimulated LexA autocleavage and subsequent derepression of lexA, recA and other regulon members (27). Induction following DNA damage ranges between regulon members; recA and lexA are induced 10 and 2- to 5-fold, respectively (43-45). Similarly, DNA damage caused by UV-irradiation or mitomycin C treatment strongly induces recA transcript and protein accumulation in another cyanobacterium, Anabaena variabilis (46,47). Levels of both the recA transcript and its protein remain elevated until the damaging agents are removed and/or the DNA repaired, as observed for E.coli recA transcripts (27). In contrast, expression of the Synechocystis recA and lexA genes was not induced by UV-irradiation. Rather, our results show that recA and lexA levels decrease following UV-treatment, in agreement with other studies (39,48). These results indicate that the DNA damage induction of recA is variable among not only cyanobacteria but also prokaryotes in general. Our observation that lexA is not induced following UV-irradiation further suggests that Synechocystis LexA is not required for survival following DNA damage, and therefore potentially regulates expression of genes not associated with DNA repair. We therefore refer to this protein as being LexA-related.

The discrepancies in *lexA* regulation may also imply differences at the protein level, where LexA self-cleavage may not be required for derepression of gene expression. This appears to be the case, as Synechocystis LexA possesses modifications in two sites important for LexA function in *E.coli*; an altered cleavage site, and the absence of the nucleophilic serine of the Ser-Lys dyad. In E.coli, LexA self-cleavage and derepression of the SOS regulon requires a catalytic serine/lysine dyad and an Ala-Gly cleavage bond (31). In the absence of these residues, as indicated by mutational studies, LexA self-cleavage in *E.coli* is defective (31). These modifications to the Synechocystis protein have been previously noted as a potential explanation for the absence of a 'cvanobacterial' SOS box within the upstream regions of Synechocystis DNA repair genes (36), and further imply an unique cellular function for the Synechocystis LexA protein.

Novel roles for LexA have been implied in other bacteria, including Mycobacterium tuberculosis (49,50) and Deinococcus radiodurans (51), although the alternative function has not been identified. In D.radiodurans, RecA protein levels remain unchanged regardless of the lexA status (52), which is unexpected if LexA is required to regulate recA expression and is similar to the results reported here. DNA damage induction of repair genes in *M.tuberculosis* also occurs predominately via a LexA- and RecA-independent mechanism as shown by mitomycin C induction of DNA repair gene expression in recA mutants (49,50). Evidence for separation of recA expression from lexA regulation may also exist in higher plant chloroplasts, which possess a DNA damage-induced recA homologue (53), while lexA has not been reported to be encoded by plant genomes. This evidence suggests that conservation of the LexA/RecA regulation of the SOS response may be less widespread than previously anticipated, and furthermore, homologues of these proteins may fulfill different roles in their respective hosts.

crhR encodes an RNA helicase proposed to regulate RNA metabolism through its modification of RNA secondary structure (1,26). The redox-responsive regulation of crhR expression suggests that its cellular capacity to catalyze RNA secondary structure modifications is regulated by the redox status of the electron transport chain. The observed induction of crhR resulting from respiratory electron flow suggests that crhR may regulate the function of RNAs associated with photosynthesis (light harvesting and/or carbon metabolism) (1) or the cellular response to the predominating light/redox environment. The implication of *crhR* in carbon metabolism is consistent with both a proposed role for LexA in the regulation of carbon uptake and utilization genes (39) and observations demonstrating reduction in lexA transcript accumulation following a downshift in inorganic carbon availability (54). Further to our hypothesis suggesting LexA may ensure cells express the necessary gene products to respond to a dynamic light environment, one of the proposed functions for the bidirectional hydrogenase, a recently identified LexA-activated gene (37,38), is as an electron valve during photosynthesis (55). LexA regulation of crhR and the hoxEFUYH bidirectional hydrogenase may ensure continued maximal photosynthetic capabilities in response to changing cellular redox conditions.

Identification of a LexA-related protein as the regulator of crhR transcription provides unique insights into the

suggest a unique regulatory role for *Synechocystis* LexA in regulating gene expression in response to environmental cues other than DNA damage. These insights also imply the ubiquitous nature of the Lex/RecA DNA repair dogma is not conserved in *Synechocystis*, raising questions regarding the mechanisms by which DNA repair gene expression is regulated in this organism.

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