



ORIGINAL RESEARCH COMMUNICATION

Cysteine Mutational Studies Provide Insight into a Thiol-Based Redox Switch Mechanism of Metal and DNA Binding in FurA from *Anabaena* sp. PCC 7120

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Abstract

Aims: The ferric uptake regulator (Fur) is the main transcriptional regulator of genes involved in iron homeostasis in most prokaryotes. FurA from *Anabaena* sp. PCC 7120 contains five cysteine residues, four of them arranged in two redox-active CXXC motifs. The protein needs not only metal but also reducing conditions to remain fully active *in vitro*. Through a mutational study of the cysteine residues present in FurA, we have investigated their involvement in metal and DNA binding. **Results:** Residue C¹⁰¹ that belongs to a conserved CXXC motif plays an essential role in both metal and DNA binding activities *in vitro*. Substitution of C¹⁰¹ by serine impairs DNA and metal binding abilities of FurA. Isothermal titration calorimetry measurements show that the redox state of C¹⁰¹ is responsible for the protein ability to coordinate the metal corepressor. Moreover, the redox state of C¹⁰¹ varies with the presence or absence of C¹⁰⁴ or C¹³³, suggesting that the environments of these cysteines are mutually interdependent. **Innovation:** We propose that C¹⁰¹ is part of a thiol/disulfide redox switch that determines FurA ability to bind the metal corepressor. **Conclusion:** This mechanism supports a novel feature of a Fur protein that emerges as a regulator, which connects the response to changes in the intracellular redox state and iron management in cyanobacteria. *Antioxid. Redox Signal.* 24, 173–185.

Introduction

ACQUIRING AND MAINTAINING ADEQUATE intracellular iron levels are major challenges that face most organisms, since insufficient iron impairs the correct function of essential iron proteins, whereas excess free iron can be deleterious to biomolecules due to its high reactivity, producing reactive-free radicals. The control of intracellular iron concentration in many bacteria is mainly operated by the transcriptional regulator Fur protein (11). The current model of action for this protein establishes that, under iron rich conditions, a dimer of Fur binds to specific DNA sequences (“iron boxes”) located in the promoters of iron responsive genes using Fe²⁺ as corepressor and prevents their transcription (3). However, this model does not take into account

Innovation

Ferric uptake regulator (Fur) proteins control intracellular iron concentration in most bacteria. The results of this study show for the first time for a Fur homologue that thiol/disulfide interconversion controls the activity of cyanobacterial FurA. The reduced FurA can bind its metal corepressor and hence DNA conversely. When FurA is oxidized it loses the metal and dissociates from DNA. The FurA thiol/disulfide exchange responds to alteration in the cellular redox potential. Thus, the binding of FurA to the metal corepressor mediated by its redox state may provide the basis of a concerted mechanism of iron homeostasis in response to the redox state of the cytosol in *Anabaena*.

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certain aspects of Fur-mediated regulation revealed by recent studies. In this sense, Fur participates in gene activation and iron-independent regulation of a variety of genes involved in distinct functions (6, 36).

The crystal structures of diverse Fur family members confirm the basic folding firstly described for the *Pseudomonas aeruginosa* Fur homologue, even though each Fur subclass responds to different signals (42). It consists of two well-defined domains: an N-terminal DNA binding domain and a C-terminal dimerization region. The amino acid sequences of many Fur homologues include two potential metal binding motifs, a conserved HHXHXXCXXC signature and another less conserved C-terminal CXXC motif. In fact, two or three metal atoms whose coordination involves amino acids of these motifs are observed in the crystal structures of several Fur homologues (1, 6, 9, 34, 39, 47, 48). Usually, one of the metal binding sites accommodates the regulatory metal atom, while another site has a structural character. Regarding the third site, it seems to play a role in stabilizing the dimeric form of the regulator (9).

The regulatory metal binding site appears to be conserved in all Fur and Fur-like proteins, although it shows some variability in the coordination depending on the Fur homologue and the metal. The structural metal binding site, in some Fur homologues, involves regular tetrahedral coordination of a zinc atom by four cysteine residues belonging to the two CXXC sequences present in the previously mentioned metal binding motifs (9, 28, 34). In fact, zinc coordination by the four cysteines of both CXXC motifs is suggested to be critical to stabilize the dimeric structure in *Bacillus subtilis* PerR (48). However, the existence of these CXXC sequences does not ensure the binding of structural metal. In this sense, the crystal structure of Nur, obtained under reducing conditions, indicates that both cysteines of the two CXXC motifs do not coordinate zinc, although this regulator maintains its DNA binding activity (1). In addition, *Vibrio cholerae* Fur structure does not reveal any metal binding site involving equivalent cysteines present in its primary sequence (47).

In the filamentous nitrogen-fixing cyanobacterium *Anabaena* sp. PCC 7120, three different Fur proteins have been identified: FurA (*all1691*), FurB (*all2473*), and FurC (*alr0957*) (21). FurA is a multifaceted protein that controls iron homeostasis and modulates an extensive regulon involved in a variety of cell functions (13, 14, 15). It also fulfills the features of a heme sensor protein whose interaction with this cofactor negatively affects its DNA binding ability *in vitro* (23, 41).

The FurA primary sequence contains five cysteines located at the C-terminal domain (C¹⁰¹, C¹⁰⁴, C¹³³, C¹⁴¹, and C¹⁴⁴), four of them arranged in two CXXC motifs (C¹⁰¹VKC¹⁰⁴ and C¹⁴¹PKC¹⁴⁴) (4). However, these CXXC motifs do not appear to coordinate a structural metal atom in FurA since metal analysis and electrospray ionization mass spectrometry revealed that neither zinc nor other metals are present in the purified protein (20). Interestingly, recent data have demonstrated that FurA exhibits disulfide reductase activity *in vitro* based on these CXXC motifs (5).

Previous results from our laboratory indicated that recombinant FurA exists in solution in several discrete disulfide-linked oligomeric species, being monomers and dimers the major species (20), and that four out of five cys-

teines are not reactive. In fact, determination of free thiols indicated that only one cysteine was accessible to solvent (20). Different data suggest the relevance of cysteines and their redox status in recombinant *Anabaena* FurA function. On one hand, thiol reducing agents enhance the affinity of FurA for its target DNA sequences *in vitro* (22). On the other hand, FurA reductase activity occurs depending on the redox state of cysteines present in its CXXC motifs (5). Finally, the FurA-heme interaction that involves residue C¹⁴¹ from one of the CXXC motifs undergoes redox-mediated ligand switching (41). Therefore, considering the apparent dependence of FurA activity on its oxidation state and the lack of a structural metal involving its two CXXC motifs, it can be envisaged that at least one redox-active cysteine would be required for the function of this protein. We therefore sought to identify the redox-active cysteine(s) in FurA to deepen its/their involvement in the operation mechanism of this regulator. For this purpose, mutations in the cysteines present in FurA have been performed to analyze their implication in metal and DNA interactions.

Our results evidence that a single cysteine belonging to one of the CXXC motifs of FurA is essential for both metal and DNA binding *in vitro*. This reactive cysteine, identified as C¹⁰¹, is responsible for the requirement of reducing agents by recombinant FurA to maintain optimal DNA binding activity. In addition, we have demonstrated that the redox state (thiol or disulfide) of C¹⁰¹ is modulated by C¹⁰⁴, its partner in the CXXC motif, compromising the ability of FurA to bind the metal corepressor necessary for DNA binding.

Results

Substitution of C¹⁰¹ by serine impairs the ability of Anabaena FurA to bind DNA

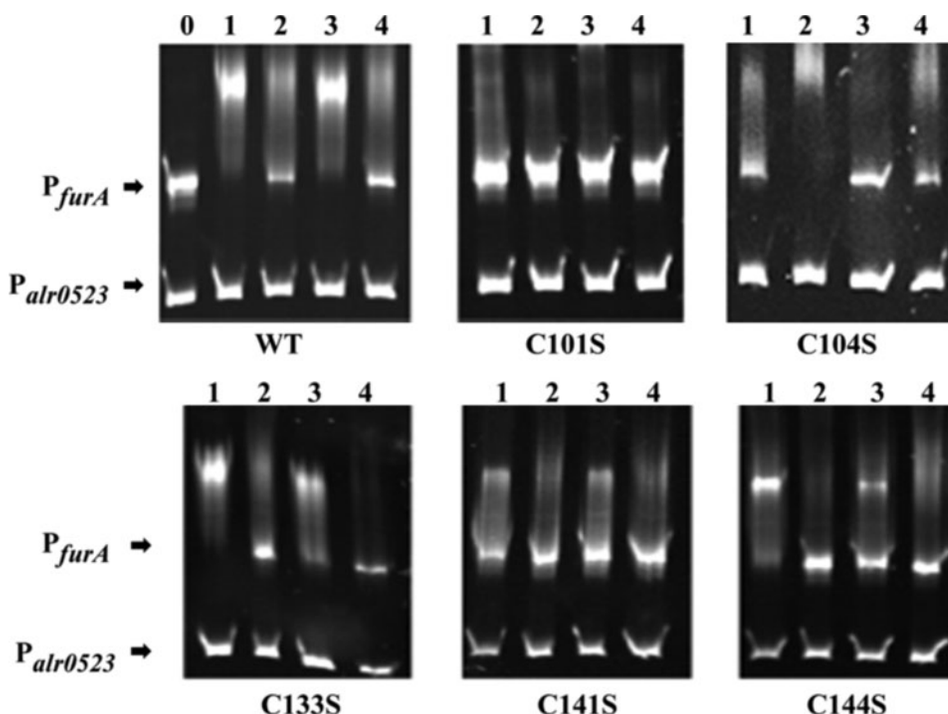
To determine whether single or multiple cysteines are important for the activity of *Anabaena* FurA *in vitro*, each of the five cysteines (C¹⁰¹, C¹⁰⁴, C¹³³, C¹⁴¹, and C¹⁴⁴) was mutated to serine. The mutations were confirmed by DNA sequencing, and after expression in *Escherichia coli*, FurA variants were purified with a yield comparable to that obtained for the wild-type protein (40).

Mutant proteins were assayed for DNA binding activity in a standard gel mobility shift assay (EMSA). Experiments were performed in the presence and absence of dithiothreitol (DTT) and/or metal, using recombinant wild-type FurA as a control. Mn²⁺ was used as a surrogate for Fe²⁺ for the *in vitro* analyses of FurA because ferrous iron readily oxidizes to Fe³⁺ in air.

Analysis of single cysteine mutants showed that C133S exhibited lower activity than wild-type FurA in the presence of Mn²⁺ or DTT (Fig. 1). Besides, C133S-DNA binding activity was negligible in the absence of both effectors (Fig. 1). Activity of C141S was also lower compared with FurA in the four conditions tested, although its dependence on metal and reducing conditions resembled to that of wild-type FurA (Fig. 1). Moreover, the highest activity of C144S was observed in the presence of DTT.

However, the most outstanding differences with respect to wild-type FurA were found in EMSA assays with mutants C101S and C104S. As shown in Figure 1, the absence of the C¹⁰⁴ residue produced a protein that only needed metal to specifically bind DNA. However, as observed for the C101S

FIG. 1. DNA binding activity of FurA (ferric uptake regulator) and its single cysteine mutants. Electrophoretic mobility shift assays show the ability of FurA wild-type or cysteine mutant proteins to bind its own promoter (P_{furA}) in the presence of a nonspecific DNA control ($P_{alr0523}$) *in vitro*. One hundred nanograms of DNA fragments free (lane 0) or mixed with 300 nM FurA proteins were separated on 6% native PAGE. Lane 1: binding assay in the presence of 1 mM DTT and 100 μ M Mn^{2+} ; lane 2: only in the presence of 100 μ M Mn^{2+} ; lane 3: only in the presence of 1 mM DTT; lane 4: in the absence of both DTT and Mn^{2+} . DTT, dithiothreitol.



mutant, the absence of the C¹⁰¹ residue prevented the binding of the protein to the consensus sequence in any assayed conditions (Fig. 1). According to these results, C¹⁰¹ seems to be essential for the sequence-specific FurA-DNA binding *in vitro*.

The loss of DNA binding ability of the C101S mutant is not due to a structural change

To know whether the loss of DNA binding activity observed for the C101S variant was associated with an important change in secondary or tertiary structures of FurA after mutation, circular dichroism (CD), Fourier transform infrared spectroscopy (FT-IR), and nuclear magnetic resonance spectroscopy (¹H 1D-NMR) experiments of wild-type FurA and C101S variant were performed.

Based on CD analysis, the substitution of C¹⁰¹ by serine caused small changes in the shape of the spectra at 222 nm, which could be due to the absorbance of cysteine residues, occurring in this region (31, 50) (Supplementary Fig. S1; Supplementary Data are available online at www.liebertpub.com/ars). The results obtained by FT-IR indicated that the content in α -helix and β -sheet was similar in the wild-type and C101S proteins (Supplementary Table 1). Finally, the ¹H 1D-NMR spectra of both species were identical (Supplementary Fig. S2), indicating the absence of major changes in the secondary and tertiary structures of the wild-type protein and its mutants. All these results strongly suggest that the inability of C101S to bind DNA is not due to a change in the protein secondary or tertiary structures after mutation of C¹⁰¹.

The reduced state of the C¹⁰¹ is essential for Mn²⁺ coordination by wild-type FurA

Structural and site-directed mutagenesis data reported in the literature indicated that cysteines are involved in metal binding sites in different Fur and Fur-like homologues (1, 6,

9, 30, 34, 39). After ruling out a structural change as the cause of the loss of activity of the C101S mutant, the ability of FurA cysteine mutants to bind the metal corepressor was assessed by isothermal titration calorimetry (ITC). Taking into account the importance of redox conditions and the presence of metal corepressor to allow purified FurA-DNA interaction (22), the binding of Mn²⁺ to wild-type FurA was studied by ITC, either in the presence or absence of DTT.

At acidic pH (pH 4), FurA did not interact with Mn²⁺, even in the presence of 1 mM DTT (not shown). At basic pH, FurA is an aggregation-prone species and this tendency rises with increasing protein concentrations (20). Therefore, the measurements were performed at basic pH in the presence of 50 mM arginine, whose use has been described as an assistant for correct folding of previously denatured proteins (37, 46) and it seems to minimize intermolecular noncovalent interactions (2).

Figure 2 shows the results of titrations of reduced and oxidized wild-type FurA with MnCl₂ in the presence of arginine. In the curve for oxidized wild-type FurA, the interaction is characterized by a small enthalpic contribution (Fig. 2A). Fitting the data to a model with a single binding site led to a dissociation constant (K_d) value of 0.28 μ M for the interaction of Mn²⁺ and wild-type FurA. However, addition of DTT increased about 20 times the heat effect (Fig. 2B) and the analysis required fitting the data to a model with two independent binding sites, providing K_d values of 50 nM and 0.84 μ M, respectively. These data indicated that the higher affinity metal binding site disappears when cysteines are in the disulfide state, suggesting that cysteines are involved in metal binding in FurA.

To investigate the location of the two Mn²⁺ sites in FurA, metal binding affinities were determined for all single cysteine FurA mutants under reducing conditions (Supplementary Fig. S3). Four out of five mutants (C104S, C133S,

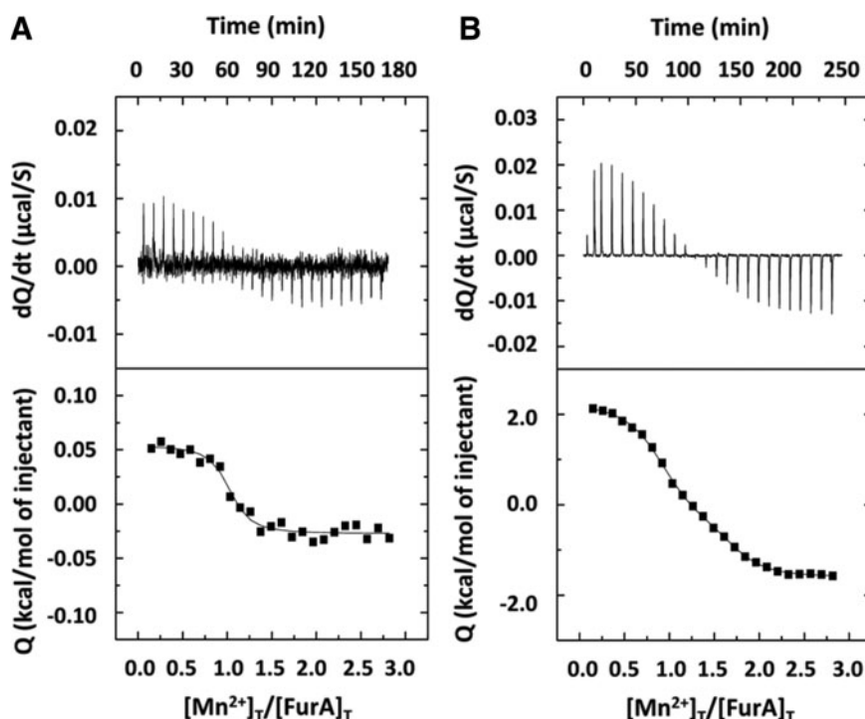


FIG. 2. ITC experiment following Mn^{2+} binding to oxidized or reduced FurA. Top sections show raw data obtained on a MicroCal VP-ITC calorimeter at 25°C. Aliquots of 300 μM MnCl_2 were added over 20 μM FurA in 50 mM Tris/HCl, 50 mM arginine, and pH 9 in the absence (A) or presence (B) of 1 mM DTT. Lower sections show the ITC isotherms of the data integrated and fitted to a single-site model (A) or two independent-site models (B). ITC, isothermal titration calorimetry.

C141S, and C144S) showed Mn^{2+} binding affinities in the same order of magnitude as that of the wild-type protein (Table 1), in contrast to C101S variant where the mutation of C¹⁰¹ led to lose the Mn^{2+} higher affinity site. The lack of the higher affinity site was also observed in a double mutant C101/133S (not shown). However, major changes were not observed regarding the second Mn^{2+} binding site. Thus, C¹⁰¹ residue is clearly important for FurA to coordinate Mn^{2+} *in vitro*. Moreover, since there are no significant changes in the secondary or tertiary structures of C101S, we suggest that the loss of the higher affinity metal binding site is largely responsible for the inability of C101S to bind DNA.

According to the results shown so far, we can conclude that the higher affinity metal binding site is lost in oxidized FurA and the C¹⁰¹ residue is essential for the binding of the metal atom. Therefore, it appears that, in purified FurA, C¹⁰¹ is part of a disulfide bridge and its redox state is responsible for the protein to coordinate the metal corepressor, since C101S

mutant cannot bind DNA in the presence of metal under reducing conditions (Fig. 1).

Recombinant FurA contains two intramolecular disulfide bridges and one free cysteine

Since sulfenic acids can serve as intermediates between thiols and disulfides, we tested the possibility that FurA cysteines could be oxidized to sulfenic acids using 7-chloro-4-nitrobenz-2-oxa-1,3-diazole (NBD-Cl). This electrophilic compound reacts with thiols and sulfenic acids, but the conjugation products have different absorption wavelengths. Upon reaction of NBD-Cl with thiol groups, an absorption peak at about 420 nm can be registered, whereas the peak shifts to 347 nm when the compound reacts with sulfenic acids (38). This approach showed that no sulfonylation occurs in prerduced FurA untreated or treated with H_2O_2 upon exposure to NBD-Cl (Supplementary Fig. S4).

Trapping experiments of free thiol groups were performed to determine the presence of disulfide bonds in the recombinant wild-type FurA. The thiol modifying agent 4-acetamido-4'-maleimidylstilbene-2,2'-disulfonic acid (AMS) reacts with free cysteines in the protein increasing its molecular weight in 540 Da per AMS molecule added (8).

As observed in Figure 3, reduced wild-type FurA displayed five free cysteines with a size shift of 2700 Da. Two bands were observed in the oxidized state, one corresponding to FurA dimers with an intermolecular disulfide bridge (no AMS molecules) and another band corresponding to monomers showing a weight consistent with the uptake of one AMS molecule. A control of recombinant FurA with no DTT subjected to AMS alkylation showed an identical band pattern to that of oxidized FurA (not shown), confirming that the purified protein is obtained in the oxidized state. According to these results, recombinant wild-type FurA monomers contain two intramolecular disulfide bridges and a single free

TABLE 1. DISSOCIATION CONSTANTS (K_D) ESTIMATED BY ITC ANALYSES FOR THE INTERACTION OF FURA PROTEINS WITH Mn^{2+} , FITTING THE EXPERIMENTAL DATA TO THE THEORETICAL EQUATION BASED ON ONE (K_{D1}) OR TWO (K_{D2}) BINDING SITE MODELS

Protein	Site 1 (nM)	Site 2 (μM)
FurA	$K_{d1} = 50$	$K_{d2} = 0.8$
C101S	—	$K_{d2} = 0.5$
C104S	$K_{d1} = 14$	$K_{d2} = 3.4$
C133S	$K_{d1} = 47$	$K_{d2} = 3.0$
C141S	$K_{d1} = 32$	$K_{d2} = 10.2$
C144S	$K_{d1} = 71$	$K_{d2} = 1.8$

Relative errors: 10–15% for K_D , 5% for ΔH and $-\Delta S$, and 2% for ΔG .

Fur, ferric uptake regulator; ITC, isothermal titration calorimetry.

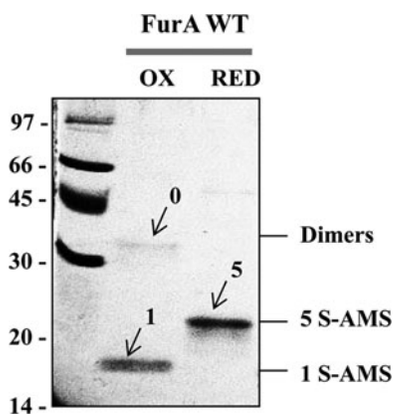


FIG. 3. *In vitro* redox state of FurA. DTT treatment, free thiol alkylation with AMS, and electrophoretic analysis of FurA were performed as indicated in the Materials and Methods section. The resulting proteins resolved on 12.5% nonreducing SDS-PAGE gels were stained with Coomassie blue. The presence of dimers and monomers with 1 (1 S-AMS) or 5 (5 S-AMS) free thiols alkylated with AMS is indicated. OX, oxidized protein; RED, reduced protein; SDS-PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis.

cysteine that is prone to forming an intermolecular disulfide bond with the free cysteine of another monomer. This is in good agreement with previous data obtained by analyzing the free thiol content in FurA with the Ellman's reagent (20).

C¹⁰¹ seems to be responsible for dimerization of recombinant wild-type FurA

Sensitivity of recombinant FurA and its cysteine mutants to reducing conditions were characterized using nonreducing sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE). FurA mutant proteins preincubated with 10 mM DTT were predominantly found as monomers, although a weak band appeared at the position expected for dimers, likely resulting of hydrophobic interactions (Fig. 4). Thus, reduced proteins contained a very little amount of intermolecularly cross-linked dimeric species, confirming that dimerization of wild-type FurA and cysteine mutant species was mainly due to intermolecular disulfide bridges. The

differences in the hydrophobicity of the resulting proteins after mutation of cysteines, which are supposed to be in the surface of the protein, could account for the trimer bands observed after reduction with DTT (33).

In contrast, in the absence of DTT, the C101S variant migrates mainly as a monomeric species and no tetramer band is observed, unlike the rest of mutants (Fig. 4), suggesting that the formation of tetramers in purified wild-type FurA is dependent on the prior existence of dimers *in vitro*. From this finding, it follows that in C101S variant, intermolecular disulfide bond formation is disfavored with respect to the other cysteine mutants and it is reasonable to assume that the four cysteines in this variant (C¹⁰⁴, C¹³³, C¹⁴¹, and C¹⁴⁴) form intramolecular disulfide bridges.

Considering the oxidation state of the cysteines in FurA and the inability of the C101S mutant to bind DNA in any conditions tested (Fig. 1), we suggest that the low DNA binding activity of FurA with metal but not DTT could be attributed to its inability to bind the metal corepressor, as a consequence of the formation of a disulfide bridge between C¹⁰¹ from different monomers, preventing metal binding.

Characterization of intramolecular disulfide bridges in wild-type FurA monomers

The behavior shown by FurA and its C101S mutant led us to consider that residues C¹⁰⁴, C¹³³, C¹⁴¹, and C¹⁴⁴ in recombinant wild-type FurA are involved in intramolecular disulfide bonds.

Analyzing the position of these residues in the primary sequence of FurA contributed to shed some light on the possible match. Taking into account that the formation of a disulfide bond between the cysteines of a CXXC motif is strongly favored when this sequence is within an N-terminal helical position (26), we used the PSIPRED secondary structure prediction server to assess whether the CXXC sequences in FurA were helical (25, 29). According to this method, C¹⁴¹ and C¹⁴⁴ are part of a CXXC motif (C¹⁴¹PKC¹⁴⁴) predicted to be helical, and therefore, both residues are likely to form a disulfide bridge (Supplementary Fig. S5).

Several evidences contributed to support the existence of an intramolecular disulfide bond between C¹⁴¹ and C¹⁴⁴. First of all, electrophoretic analyses in the presence or absence of DTT were carried out to evaluate the contribution of

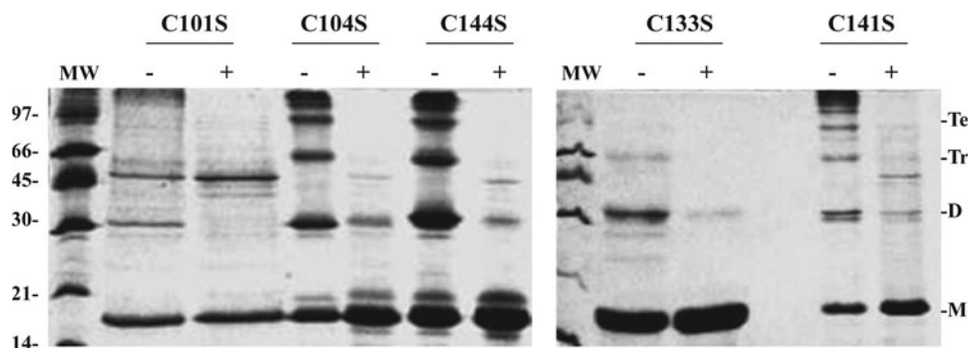


FIG. 4. Nonreducing electrophoretic analyses of the oligomeric state of FurA and its single cysteine mutants. The 17% nonreducing SDS-PAGE gels stained with Coomassie blue show the oligomerization *via* disulfide bonds of different FurA variants preincubated with 10 mM DTT as reducing agent (+) or without any treatment (-). MW, molecular weight markers (97, 66, 45, 30, 20, and 14 kDa). The presence of monomers (M), dimers (D), trimers (Tr), and tetramers (Te) is indicated.

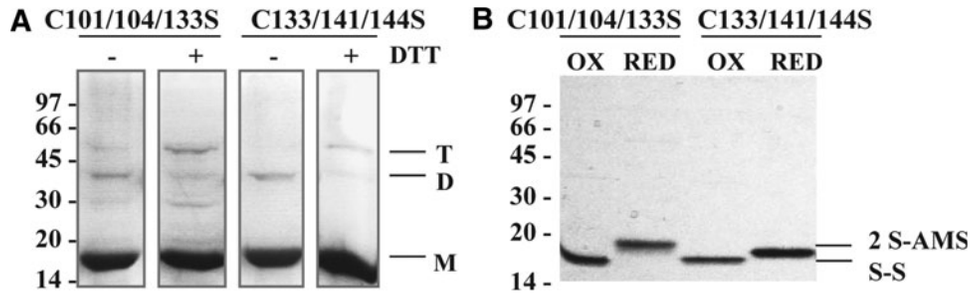


FIG. 5. Disulfide bonds in FurA triple cysteine mutants. (A) Nonreducing SDS-PAGE and Coomassie blue staining of C101/104/133S and C133/141/144S FurA variants with (+) or without (-) 10 mM DTT treatment. The presence of monomers (M), dimers (D), and trimers (T) is indicated. (B) Free thiol alkylation experiments with AMS of oxidized (OX) or reduced (RED) FurA triple cysteine mutants resolved on a nonreducing 12.5% SDS-PAGE gel and stained with Coomassie. The bands named S-S correspond to the presence of an intramolecular disulfide bond in both oxidized proteins, whereas the bands named 2 S-AMS indicate the presence of two reduced thiols alkylated with two AMS molecules (shift size 1080 Da).

disulfide bridges to the oligomerization of the triple C101/104/133S mutant. Results shown in Figure 5A indicated that, in the absence of DTT, monomers were the predominant form in this mutant. Moreover, AMS trapping experiments shown in Figure 5B indicated that the remaining cysteines (C^{141} and C^{144}) were disulfide linked when this mutant was oxidized. Furthermore, the motifs $C^{142}KEC^{145}$ in *Helicobacter pylori* Fur or $C^{105}KNC^{108}$ in the *Campylobacter jejuni* homologue, which are equivalent to *Anabaena* FurA $C^{141}PKC^{144}$ motif, are also localized in a α -helix according to the crystal structures of both regulators (6, 9). However, in *H. pylori* or *C. jejuni* Fur, these cysteines are reduced since they coordinate the structural metal. Finally, *Anabaena* FurA $C^{141}PKC^{144}$ sequence is similar to the CXXC motif of *E. coli* DsbA ($C^{30}PHC^{33}$), where these cysteines remain naturally oxidized forming a disulfide bridge (44, 52).

Accordingly, if C^{141} and C^{144} are intramolecularly disulfide bonded and C^{101} is mainly involved in the dimerization of FurA, C^{104} and C^{133} would be responsible for the second intramolecular disulfide bridge observed in the AMS trapping experiments with FurA. To gather experimental evidence on the existence of this intramolecular disulfide bond, thiol trapping experiments were performed with a triple C101/141/144S mutant containing C^{104} and C^{133} .

The thiol content in this mutant was monitored with AMS using different reduced DTT/oxidized DTT, trans-4,5-Dihydroxy-1,2-dithiane (DTT_{red}/DTT_{ox}) ratios. As observed in Figure 6, only one band was observed under the most reducing conditions (2S-AMS) that corresponded to the addition of two AMS molecules. This result indicated that the protein was completely reduced. However, several bands were observed as the DTT_{red}/DTT_{ox} ratio diminished. These bands corresponded to different species namely oxidized protein (S-S), protein with one AMS molecule (S-AMS), and different intermolecularly disulfide-bonded dimer combinations (Dimers S-S) that disappeared along the titration. This experiment clearly demonstrated that, in the triple C101/C141/C144S mutant, C^{104} and C^{133} can form an intramolecular disulfide bond *in vitro* and that intramolecularly oxidized cysteines coexist with reduced and intermolecularly oxidized cysteines.

The mutational study presented here suggests that recombinant dimeric FurA contains an intermolecular disulfide bridge $C^{101}-C^{101}$ and two intramolecular bonds $C^{104}-C^{133}$ and $C^{141}-C^{144}$ *in vitro*. Interestingly, previous data obtained in our laboratory indicated the occurrence of a redox-active disulfide bond between C^{101} and C^{104} (5). Moreover, non-reducing SDS-PAGE and AMS trapping experiments using

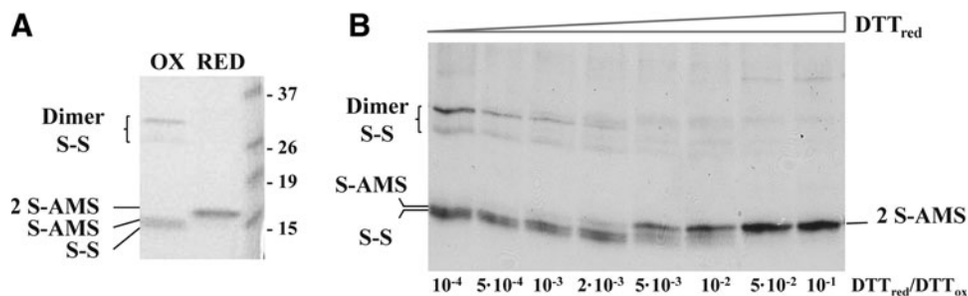


FIG. 6. C^{104} and C^{133} intramolecular disulfide bond. (A) Fully oxidized (OX) or reduced (RED) FurA triple mutant C101/141/144S was obtained as indicated in the Materials and Methods section and subjected to AMS treatment to be used as controls. (B) Oxidized and reduced fractions of the FurA triple mutant C101/141/144S were obtained by incubation with variable DTT_{red}/DTT_{ox} ratios and subjected to thiol trapping treatment with AMS. Proteins were resolved by nonreducing SDS-PAGE and Coomassie stained. The oxidized protein contained an intramolecular bond (band S-S), although some molecules exhibit a free thiol that reacted with one AMS (band S-AMS) or was intermolecularly disulfide bridged in dimers (band Dimer S-S). The reduced protein reacted with 2 AMS molecules (band 2 S-AMS).

the triple C133/141/144S mutant retaining these cysteines confirmed the presence of an intramolecular bridge C¹⁰¹-C¹⁰⁴ in the protein (Fig. 5A, B). All together, we conclude that cysteines C¹⁰¹, C¹⁰⁴, and C¹³³ of FurA can form disulfide bonds in different combinations, namely C¹⁰¹-C¹⁰⁴ or C¹⁰⁴-C¹³³.

The redox status of C¹⁰¹ is modulated by C¹⁰⁴

Mutations of any of these cysteines probably intramolecularly bridged in wild-type FurA (C¹⁰⁴-C¹³³ or C¹⁰¹-C¹⁰⁴) would release its partner to form a different disulfide bond, either intra- or intermolecularly.

This possibility prompted us to analyze the behavior of FurA and its single cysteine mutants by cross-linking experiments. Results shown in Figure 7 indicate that mutant C104S has a high tendency to dimerize, suggesting that residues C¹³³ and C¹⁰¹ are not prone to forming an intramolecular disulfide bridge. Proof of this is that the C104S variant requires only metal to reach maximal DNA binding activity (Fig. 1), and therefore, C¹⁰¹ is reduced in this mutant. These results suggest that dimers observed in the C104S mutant, unlike in wild-type FurA, would involve residues C¹³³ from two monomers and C¹⁰¹ would be free to interact with metal and hence with DNA. On the contrary, mutant C133S shows a low tendency to dimerize (Fig. 7), suggesting that C¹⁰¹ can be disulfide bridged to C¹⁰⁴ in this protein. As expected, C133S needed both metal and reducing conditions to display maximal DNA binding activity (Fig. 1), suggesting that C¹⁰¹ was actually intramolecularly disulfide bonded in this mutant and not available to directly coordinate the metal corepressor.

Previous results presented along with this article have shown that mutant C101S contained a lower amount of dimeric species than the rest of single cysteine mutants (Fig. 4 and 7), suggesting that C¹⁰⁴ and C¹³³ tend to form an intramolecular

bond *in vitro*. This mutant is unable to bind DNA, since it lacks C¹⁰¹ to coordinate the metal corepressor (Fig. 1).

In summary, the above results indicate that the redox state of C¹⁰¹ is affected by the presence or absence of C¹⁰⁴. When C¹⁰⁴ is present in a mutant lacking C¹³³ (C133S), C¹⁰¹ is oxidized, but C¹⁰¹ is reduced in a mutant lacking C¹⁰⁴ (C104S), although C¹³³ is present. Taking into account that C¹⁰⁴ can form intramolecular disulfide bonds with either C¹⁰¹ or C¹³³, a mechanism of *Anabaena* FurA activation–deactivation based on an intramolecular disulfide interchange can be envisaged. When C¹⁰¹ is bound to C¹⁰⁴, FurA is inactive because it is unable to bind the metal corepressor and hence DNA. If C¹³³ forms a disulfide bridge with C¹⁰⁴, then C¹⁰¹ is released from the C¹⁰¹-C¹⁰⁴ disulfide bond, and consequently, FurA is activated.

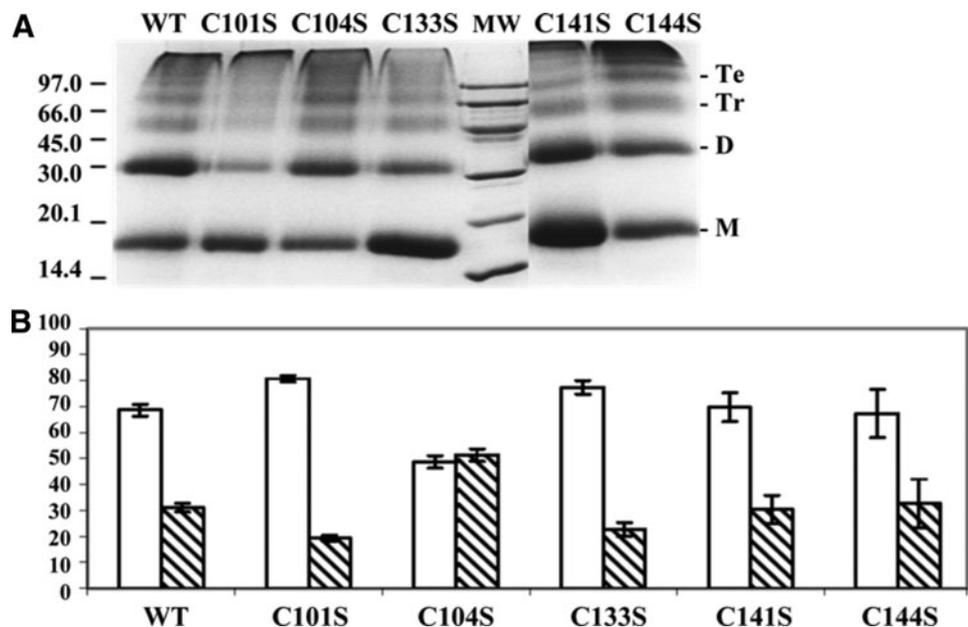
FurA thiol/disulfide redox state responds to cellular redox alterations

To determine whether thiol/disulfide couples in FurA respond to changes in intracellular redox conditions, AMS trapping experiments were used to analyze the redox states of FurA cysteines in *Anabaena* PCC 7120 cultures at different growth phases or subjected to redox stress.

Under early exponential growth conditions, FurA was mainly reduced. Stationary growth conditions, which typically involve bacterial protein oxidation as a consequence of nutritional or oxidative stresses (10), entailed the conversion of most of the cysteines to the disulfide state (Supplementary Fig. S6).

When cells in early exponential phase were subjected to oxidative stress generated by the addition of 0.5 mM H₂O₂, the proportion of reduced cysteines decreased. Estimates by densitometry indicated that around 60% of the fully reduced protein converted to forms bearing three or only one reduced cysteines (Supplementary Fig. S6). Thus, FurA thiol/

FIG. 7. Oligomerization of FurA and its single cysteine mutants. (A) Proteins were dark incubated with 0.5% glutaraldehyde, resolved on a 17% reducing SDS-PAGE gel, and stained with Coomassie blue. Monomers (M), dimers (D), trimers (Tr), and tetramers (Te), as well as the MW, are indicated. (B) The ratios of monomers (*white bars*) and dimers (*striped bars*) exhibited by wild-type FurA and the cysteine mutants are represented. Band intensities from the Coomassie-stained gels were quantified by measuring the density of the bands on Gel Doc 2000 (Bio-Rad) and analyzed by Multi-Analyst software (Bio-Rad). Values in bar graphs represent means ± standard deviations for three separate experiments performed.



disulfide state responds to changes in *Anabaena* sp. PCC 7120 intracellular redox conditions.

Discussion

The lack of structural Zn²⁺ in *Anabaena* FurA provides its cysteines with a great reactivity, not detected in other family members (5). Cysteines in FurA play a key role in the different functions, including the interactions with its metal corepressor and DNA. In this study, the status of FurA cysteines, as well as their contribution to the activity of this global regulator, has been investigated.

Our data suggest that four out of five cysteines in recombinant FurA are engaged in the formation of two intramolecular disulfide bridges, C¹⁰⁴-C¹³³ and C¹⁴¹-C¹⁴⁴. The remainder cysteine (C¹⁰¹) easily oxidizes in the air, leading to the formation of an intermolecular disulfide bond (C¹⁰¹-C¹⁰¹) that appears to be responsible for most dimers observed in recombinant FurA. This behavior is comparable to that

shown by *P. aeruginosa* Fur, where C⁹², the only cysteine in the protein, forms intermolecular disulfide bonds on air oxidation (32). Interestingly, this residue aligns with C¹⁰¹ in *Anabaena* FurA (Fig. 8). However, the formation of (C¹⁰¹-C¹⁰¹) dimers in *Anabaena* FurA leads to the loss of DNA binding ability *in vitro*, unlike *Pseudomonas* Fur where the presence of dimers is batch dependent and does not affect DNA binding (32).

The behavior of recombinant wild-type FurA *in vitro* is consistent with our *in vivo* results showing that this protein is mainly a monomer with a single free cysteine in the cytoplasm of *Anabaena* sp. PCC 7120 cultures at the stationary phase, suggesting that two intramolecular disulfide bridges are also formed *in vivo* (5).

The FurA C¹⁰¹ residue is highly conserved in Fur homologues and it belongs to a CXXC (C¹⁰¹VKC¹⁰⁴) motif present in a large number of Fur proteins as well (Fig. 8). However, the importance of this cysteine varies among different homologues. For instance, it is essential for the functionality of

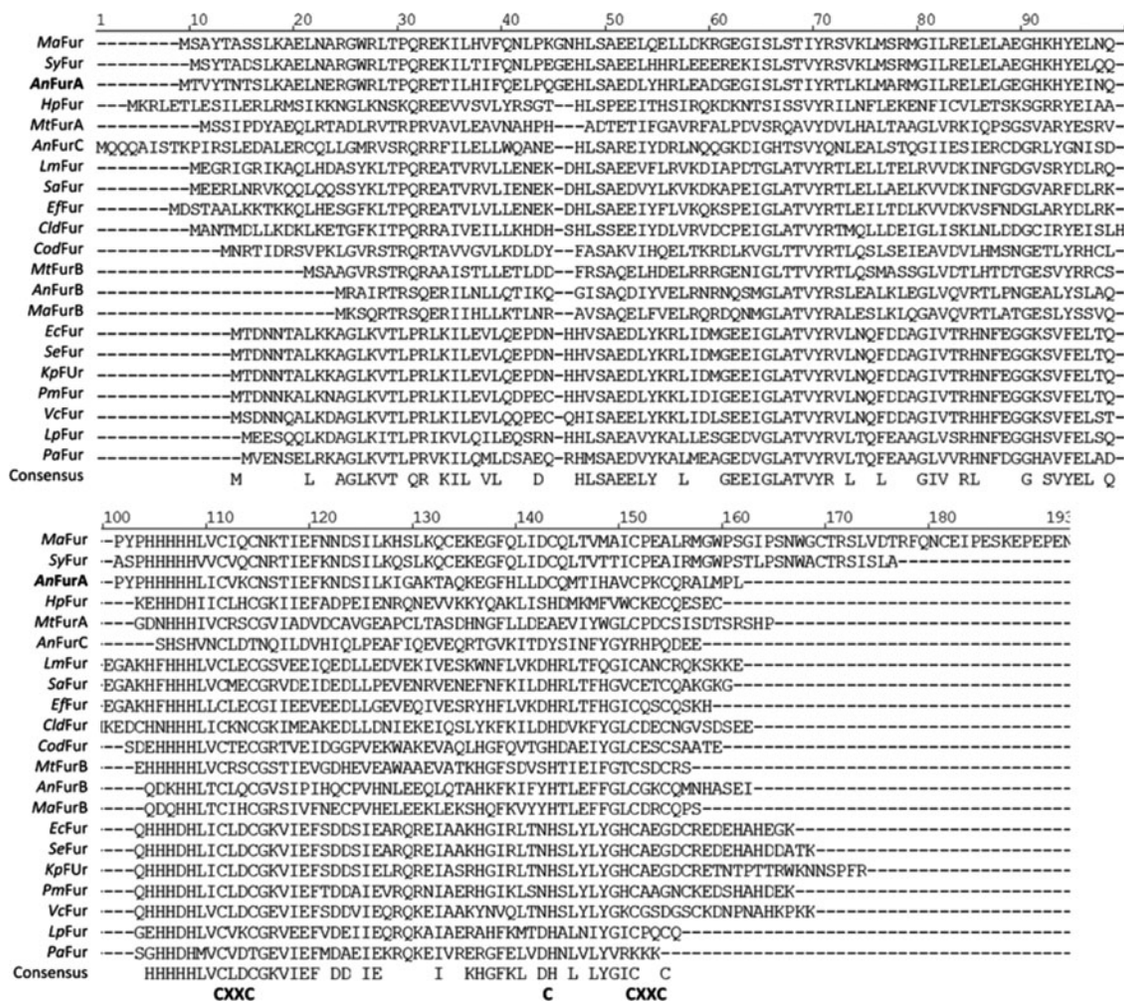


FIG. 8. Multiple sequence alignment of Fur proteins present in different bacteria. The abbreviations correspond to the following microorganisms: *Ma*: *Microcystis aeruginosa* PCC 7806, *Sy*: *Synechocystis* PCC 6803, *An*: *Anabaena* PCC 7120, *Hp*: *Helicobacter pylori* 26695, *Mt*: *Mycobacterium tuberculosis* H37Rv, *Lm*: *Listeria monocytogenes* EGDe, *Sa*: *Staphylococcus aureus* ATCC BAA-39, *Ef*: *Enterococcus faecalis* ATCC 29212, *Cld*: *Clostridium difficile* ATCC 43255, *Cod*: *Corynebacterium diphtheriae* ATCC 13812, *Ec*: *Escherichia coli* BL21, *Se*: *Salmonella enterica* Typhimurium LT2, *Kp*: *Klebsiella pneumoniae* ATCC 43816, *Pm*: *Proteus mirabilis* HI 4320, *Vc*: *Vibrio cholerae* CECT512, *Lp*: *Legionella pneumophila* Philadelphia-1, *Pa*: *Pseudomonas aeruginosa* ATCC 27853.

E. coli and *H. pylori* Fur proteins (9, 27, 39), while *P. putida* Fur lacks cysteines (32). Our results indicate that C¹⁰¹ and its particular redox state play an essential role in the coordination of the metal corepressor, which ultimately controls FurA-DNA binding activity *in vitro*. In fact, ITC experiments show that reduced recombinant FurA contains two metal binding sites of different affinities. The high affinity site (K_d 50 nM) disappears when recombinant wild-type FurA is oxidized. The C101S variant also loses the high affinity metal binding site under reducing conditions, suggesting an essential role for C¹⁰¹ in metal coordination, conditional on its redox state. Parallel EMSA experiments show that, when C¹⁰¹ is missing, FurA does not interact with DNA even in the presence of metal under reducing conditions, thus connecting FurA-DNA binding and its ability to interact with metal through C¹⁰¹.

According to our results, oxidized *Anabaena* FurA has a unique Mn²⁺ binding site with a K_d value of 0.28 μ M, assumed to correspond to the low affinity metal binding site in reduced FurA (0.84 μ M). This second metal binding site could be equivalent to the third metal binding site that, aside from the regulatory and structural ones, has been detected in the crystal structures of *H. pylori* Fur or *B. subtilis* Zur, where it is proposed to stabilize protein dimers (9, 35).

Many Fur proteins also contain a second pair of cysteines close to the C-terminus of its primary sequence, in some cases as part of a second CXXC motif. The existence of two CXXC motifs in the primary sequence of Fur has been related to the existence of zinc (6, 9, 12, 28, 48). In *Anabaena* FurA, as occurs in *Streptomyces coelicolor* Nur, both CXXC motifs are present but they do not coordinate a structural metal (20). Regarding FurA, as previously mentioned, *in vivo* results show that two intramolecular disulfide bridges are formed (5). It indicates that the presence of intramolecular disulfide bridges is important for the protein activity *in vivo*.

According to our mutational study *in vitro*, different intramolecular disulfide bonds can be found in *Anabaena* FurA, involving cysteines C¹⁰¹, C¹⁰⁴, and C¹³³. On one hand, residues C¹⁰¹ and C¹⁰⁴ belonging to the C¹⁰¹VKC¹⁰⁴ motif can be disulfide bridged in a mutant lacking C¹³³ residue. On the other hand, C¹³³ seems to be disulfide bonded to C¹⁰⁴ in C101S, while C¹⁰¹ appears not to be disulfide bridged to residue C¹³³ in C104S.

Taking into account a recent definition of redox switch (45), it is not unreasonable to speculate about a mechanism of FurA functioning based on a thiol–disulfide redox switch, involving these cysteines and controlling the redox state of C¹⁰¹. In this way, the oxidation of the C¹⁰¹VKC¹⁰⁴ motif would result in the loss of FurA metal binding ability, causing a reduction of DNA binding affinity with consequent derepression of regulated genes. It does not exclude the existence of a second thiol–disulfide switch functioning either independently or in coordination with the previous one, affecting the cysteines of the second C¹⁴¹PKC¹⁴⁴ motif. These cysteines form a redox-sensitive disulfide bridge in a triple C101/104/133S mutant containing C¹⁴¹ and C¹⁴⁴ (5). C¹⁴¹ is also involved in *Anabaena* FurA–heme coordination (41).

In this regard, a recent study has described a thiol–disulfide redox switch that controls the interaction between heme and the ligand binding domain of Rev-*erbb* (18). Moreover, an NMR study of different single and double mutants of the heme oxygenase HO-2 involved in signaling and regulatory

processes demonstrates that, analogous to our observations, the redox state of a given cysteine out of the three cysteines in HO-2 is sensitive to the absence of one or both cysteines (49).

The finding that different disulfide-bonded forms of FurA exist *in vivo*, generally with two and sometimes one or even no intramolecular disulfide bridges, whose ratio depends on the redox state of the cytoplasm of the cyanobacterium (Supplementary Fig. S6) (5), supports the idea that disulfide shuffling in FurA is important to control its function. In this sense, according to our results, an early exponential phase *Anabaena* culture contains FurA mainly in the reduced state. Exposure of this culture to oxidative stress causes the oxidation of FurA cysteines *in vivo* and presumably its inactivation. This is confirmed by different studies that show that transcription of FurA-repressed target genes, namely *isiA* and the flavodoxin-encoding gene *isiB* is induced under oxidative stress conditions in different cyanobacterial strains (19, 51). In particular, Yousef *et al.* show that transcription of major iron-regulated genes, such as *isiA*, *isiB*, and *irpA* among others, is induced by oxidative stress within a few minutes by treatment of cells with hydrogen peroxide or methyl viologen (51). Thus, it appears that the cysteines of FurA form a physiological relevant redox rheostat that responds to the intracellular redox potential.

The FurA thiol-based switch mechanism that we propose is new for this kind of metalloregulators and likely specific for cyanobacterial Fur, since it relies on the presence of C¹³³, only conserved in Fur homologues from cyanobacteria (Fig. 8). C¹³³ would be responsible for maintaining C¹⁰⁴ in the oxidized state, avoiding the formation of a C¹⁰⁴–C¹⁰¹ disulfide bond that would lead to the inactivation of FurA. The reverse process would be necessary to activate FurA again (Fig. 9). The mechanism also explains the lack of structural zinc in this regulator, since C¹⁰¹ and C¹⁰⁴ are part of the redox switch and cannot be found simultaneously reduced to coordinate zinc.

This behavior described for FurA is comparable to that shown by transglutaminase 2 (TG2), a protein tightly regulated through three identified allosteric modifiers: Ca²⁺ ions, guanine nucleotides, and an allosteric disulfide bond whose thiol/disulfide exchange is proposed to control the conversion of the inactive TG2 to the active form (7).

It would be expected that a redox stimulus would account for a disulfide bond exchange in *Anabaena* FurA that would

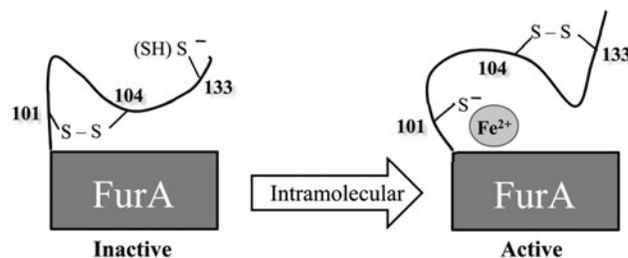


FIG. 9. Proposed mechanism of the redox-mediated FurA activity. When the protein is transcriptionally active, C¹³³ and C¹⁰⁴ form an intramolecular disulfide bond and C¹⁰¹ can coordinate the metal corepressor. Upon an intracellular redox status, a thiol–disulfide interchange would generate a new intramolecular disulfide bond between C¹⁰¹ and C¹⁰⁴. This oxidation event would simultaneously occur with metal release and FurA inactivation. The reverse process would be necessary to activate FurA again.

trigger some sort of change, allowing the protein to bind or release the metal corepressor and in turn DNA. Cleavage and formation of the corresponding disulfide bridges should be reversible so that the perturbation of the balance between reduced and disulfide-bonded forms could be the key of FurA regulation. The way by which the expected disulfide bridge between C¹⁰⁴ and C¹³³ would reform to activate FurA is an important issue. The presence of CXXC motifs in different redox states in FurA suggests that this regulator could be a redox partner of thioredoxin. However, we did not retrieve any interaction between FurA and thioredoxin or another partner entailing thiol–disulfide exchange using pull-down experiments complemented with bacterial two hybrid assays (5). Instead, we detected some interacting partners of FurA, namely the hypothetical protein All1140 that contains an Ami_3 domain related to N-acetylmuramoyl-L-alanine amidase activity, the histone-like DNA binding protein HU, and the phosphoribulokinase enzyme. Up to now, no redox partners of FurA have been identified, but work is currently underway to determine the implication of different photosynthetic electron carriers in the disulfide bond exchange in this cyanobacterial regulator.

Materials and Methods

Protein preparation

The *furA* wild-type gene (*all1691*) from *Anabaena* sp. PCC 7120 was cloned into pET28a (Novagen: Merck, Darmstadt, Germany), as previously described (4). Several pET28a-*furA* variants containing cysteine-to-serine substitutions were generated by site-directed mutagenesis or purchased from Mutagenex, Inc. (Piscataway, NJ). DNA sequences were confirmed by DNA sequencing. All the proteins were overexpressed and purified using the high-recovery one-step purification method previously described (40). Proteins were spectrophotometrically quantified using the experimental molar extinction coefficient at 276 nm 13,760 M⁻¹ cm⁻¹ (20).

Spectroscopic analyses

Ultraviolet–visible (UV-vis) absorption spectra were recorded on a Cary 100 UV-Vis double beam spectrophotometer (Agilent Technologies, Santa Clara, CA).

CD spectra were collected on a Jasco J815 spectropolarimeter (Jasco International Co. LTD., Tokyo, Japan) fitted with a thermostated cell holder and interfaced with a Peltier unit. Spectra were acquired at a scan speed of 50 nm/min with a response time of 2 s and averaged over four scans at 25°C. Far-UV measurements were performed using 10 μM of FurA proteins in 10 mM sodium acetate buffer pH 4 in 0.1 cm pathlength quartz cells (Hellma, Forest Hills, NY). All spectra were corrected by subtracting the proper baseline. The molar ellipticity ($[\theta]$) was expressed in deg · cm² · dmol⁻¹.

For FT-IR studies, the proteins (250 μg) were lyophilized and dissolved in 30 μL of deuterated water. No pH corrections were applied for isotope effects. Protein samples were placed between a pair of CaF₂ windows separated by a 50-mm-thick spacer, in a Harrick demountable cell at 25°C. Spectra were acquired on a Bruker FT-IR spectrophotometer (Bruker Optics, Inc., Billerica, MA), equipped with a DTGS detector, and thermostated with a Braun water bath. The cell

container was filled with dry air. Five hundred scans per sample were taken, averaged, apodized with a Happ–Genzel function, and Fourier transformed to give a final resolution of 2 cm⁻¹. The signal-to-noise ratio of the spectra was better than 10,000:1. Baselines were subtracted, and contributions of side chains removed as described (16, 17). Only in those cases where there were no distortions in the spectra, the resulting difference spectra were used for analysis. To determine the amount of secondary structure components, the Amide I band in D₂O was decomposed into its constituents by curve fitting, using a line shape that was a combination of Gaussian and Lorentzian functions (43).

The NMR experiments were acquired on a Bruker Avance DRX-500 spectrometer. Samples of FurA proteins (0.3–1.4 mM) were prepared in a 10 mM deuterated sodium acetate buffer pH 4 in 5-mm NMR tubes at 25°C. Spectra were acquired using 16 K data points, averaged over 512 scans and with a spectral width of 7801.69 Hz (13 ppm).

Isothermal titration calorimetry

Metal binding to FurA proteins was assayed in a high-sensitivity isothermal titration VP-ITC microcalorimeter (MicroCal, Northampton, MA). Protein samples and reference solutions were properly degassed and carefully loaded into the cells to avoid bubble formation during stirring. Experiments were performed with 20 μM freshly prepared FurA protein solutions in 50 mM Tris buffer and 50 mM arginine pH 9 at 25°C. Samples were titrated with 300 μM MnCl₂ in the calorimetric cell. Control experiments were performed under the same experimental conditions. The heat due to the binding reaction was obtained as the difference between the reaction heat and the corresponding heat of dilution, the latter estimated as an adjustable parameter in the analysis. The association constant and the enthalpy change were obtained through nonlinear regression of experimental data to a model for either a single ligand binding site or two independent ligand binding sites.

Electrophoretic mobility shift assays

Gel retardation assays were performed as previously described (22) using the promoter region of *furA* (P_{*furA*}) as target DNA and the promoter of *alr0523* (P_{*alr0523*}) as non-specific DNA. Both promoters were obtained by PCR amplification using oligonucleotides 5'-CTCGCTAGCAA TTTAACAAC-3' and 5'-GCCTTGAGCGAAGTATTTGTG-3' for P_{*furA*} and 5'-GTCTGTATGGATTAACACTATC-3' and 5'-CTGCCTGTTCAACTACTTTGG-3' for P_{*alr0523*} and then purified by the GFX PCR DNA and Gel Band Purification Kit (GE Healthcare, Buckinghamshire, United Kingdom). A 300 nM protein solution was incubated with 100 ng for each target and nonspecific DNA regions in the reaction buffer containing 10 mM Bis-Tris pH 7.5, 40 mM KCl, 5% glycerol, and 0.05 mg/mL bovine serum albumin for 20 min before being loaded on native 6% PAGE gels. Depending on the tested conditions, 1 mM DTT and/or 100 μM MnCl₂ were added to the reaction mixtures.

AMS trapping of reduced thiols

Proteins (25 μM) were incubated for 2 h at 30°C in a redox buffer containing 100 mM *trans*-4,5-dihydroxy-1,2,dithane

(oxidized DTT, DTT_{ox}; Sigma-Aldrich, St. Louis, MO) and 1 μ M or 20 mM (DTT_{red}; Sigma-Aldrich) to obtain oxidized or reduced FurA proteins, respectively. Reactions were stopped by adding 5% (v/v) trichloroacetic acid (TCA; Sigma-Aldrich), and proteins were precipitated by centrifugation at 13,300 g for 10 min at 4°C. After washing with cold acetone, air-dried protein pellets were resuspended in 10 μ L of thiol modification buffer containing 50 mM Tris/HCl pH 7.5, 1% (w/v) SDS (Sigma-Aldrich), and 20 mM AMS (Molecular Probes; Life Technologies Ltd., Paisley, United Kingdom). Proteins in nonreducing Laemmli buffer were resolved on 12.5% (w/v) polyacrylamide [30:0.8 (w/w) acrylamide/bisacrylamide] SDS-PAGE gels and stained with Coomassie Blue R-250 (Sigma-Aldrich) (24).

For *in vivo* trapping experiments, 10 mL of exponentially growing *Anabaena* cultures was treated with 0.5 mM hydrogen peroxide for 30 min. A culture without treatment was used as control. Both cultures were then incubated with TCA to a final concentration of 1% (v/v) for 30 min. Proteins were precipitated by centrifugation, washed with cold acetone, and resuspended in 25 μ L of thiol modification buffer. After 1 h of treatment, samples were subjected to 12.5% nonreducing SDS-PAGE and revealed by Western blotting with rabbit polyclonal antibodies raised against FurA.

Analysis of the oligomeric state of FurA proteins in vitro

Oligomeric species were analyzed using glutaraldehyde as a chemical crosslinker. Proteins (40 μ M) were incubated in the dark with 0.5% (v/v) glutaraldehyde (Sigma-Aldrich) for 30 min, and reactions stopped by adding Laemmli buffer and boiling. Samples were finally subjected to 17% SDS-PAGE and stained with Coomassie Blue R-250. The effect of the reducing agent was assessed by incubating 40 μ M protein solutions with 10 mM DTT for 30 min. Nonreducing Laemmli buffer was added to the samples before boiling. Proteins were separated by 17% SDS-PAGE followed by Coomassie Blue R-250 staining.

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Author Disclosure Statement

No competing financial interests exist.

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Abbreviations Used

[θ] = molar ellipticity
 AMS = 4-acetamido-4'-maleimidylstilbene-2,2'-disulfonic acid
 CD = circular dichroism
 D = dimers
 DTT = dithiothreitol
 DTT_{ox} = oxidized DTT, trans-4,5-Dihydroxy-1,2-dithiane
 DTT_{red} = reduced DTT
 EMSA = electrophoretic mobility shift assay
 FT-IR = Fourier transform infrared spectroscopy
 Fur = ferric uptake regulator
 ITC = isothermal titration calorimetry
 K_d = dissociation constant
 M = monomers
 MW = molecular weight markers
 NBD-Cl = 7-chloro-4-nitrobenz-2-oxa-1,3-diazole
 NMR = nuclear magnetic resonance
 OX = oxidized protein(s)
 PAGE = polyacrylamide gel electrophoresis
 Palr0523 = promoter region of the *alr0523* gene
 P_{furA} = promoter region of the *furA* gene
 RED = reduced protein(s)
 SDS = sodium dodecyl sulfate
 TCA = trichloroacetic acid
 Te = tetramers
 TG2 = transglutaminase 2
 Tr = trimers
 UV-vis = ultraviolet-visible region