Biogenesis of Iron-Sulfur Clusters in Photosystem I

HOLO-NfuA FROM THE CYANOBACTERIUM SYNECHOCOCCUS SP. PCC 7002 RAPIDLY AND EFFICIENTLY TRANSFERS [4Fe-4S] CLUSTERS TO APO-PsaC IN VITRO

Revised for publication, May 5, 2008, and in revised form, July 29, 2008. Published, JBC Papers in Press, August 11, 2008, DOI 10.1074/jbc.M803395200

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The NfuA protein has been postulated to act as a scaffolding protein in the biogenesis of photosystem (PS) I and other iron-sulfur (Fe/S) proteins in cyanobacteria and chloroplasts. To determine the properties of NfuA, recombinant NfuA from Synechococcus sp. PCC 7002 was overproduced and purified. In vitro reconstituted NfuA contained oxygen- and EDTA-labile Fe/S cluster(s), which had EPR properties consistent with [4Fe-4S] clusters. After reconstitution with $^{57}$Fe$^{2+}$, Mössbauer studies of NfuA showed a broad quadrupole doublet that confirmed the presence of [4Fe-4S]$^{3+}$ clusters. Native gel electrophoresis under anoxic conditions and chemical cross-linking showed that holo-NfuA forms dimers and tetramers harboring Fe/S cluster(s). Combined with iron and sulfide analyses, the results indicated that one [4Fe-4S] cluster was bound per NfuA dimer. Fe/S cluster transfer from holo-NfuA to apo-PsaC of PS I was studied by reconstitution of PS I complexes using P700-FX core complexes, PsaD, apo-PsaC, and holo-NfuA. Electron transfer measurements by time-resolved optical spectroscopy showed that holo-NfuA rapidly and efficiently transferred [4Fe-4S] clusters to PsaC in a reaction that required contact between the two proteins. The NfuA-reconstituted PS I complexes had typical charge recombination kinetics from $[F_{A}/F_{B}^{-}]$ to P700$^{0}$ and light-induced low-temperature EPR spectra. These results establish that cyanobacterial NfuA can act as a scaffolding protein for the insertion of [4Fe-4S] clusters into PsaC of PS I in vitro.

In the photosynthetic electron transport chains of cyanobacteria and chloroplasts of higher plants, iron-sulfur (Fe/S)$^4$ clusters not only serve as the terminal electron-accepting cofactors in photosystem (PS) I but they also act as electron transfer cofactors in the cytochrome $b_{f}$ complex and in soluble ferredoxin. PS I is a large, membrane-protein complex that catalyzes the light-induced transfer of electrons from plastocyanin or cytochrome $c_{6}$ in the thylakoid lumen to ferredoxin or flavodoxin in the cytoplasm (1, 2). As indicated by the x-ray crystal structure, the PS I reaction center of Synechococcus elongatus contains 12 protein subunits, which bind 96 chlorophyll (Chl) a molecules, 22 $\beta$-carotenes, 2 phylloquinones, 3 [4Fe-4S] clusters, and 5 lipids (3). The PS I electron transport chain consists of P700 (Chl a—Chl a$^{+}$ dimer), A$_{0}$ (Chl a monomer), A$_{1}$ (phylloquinone), and three [4Fe-4S] clusters, which are the terminal electron acceptors denoted F$_{A}$, F$_{A'}$, and F$_{B}$. PsaC and PsaB each provide two cysteine (Cys) ligands to the intrasubunit [4Fe-4S] F$_{X}$ cluster, and the F$_{A}$ and F$_{B}$ terminal electron acceptors are bound to a 9-kDa soluble protein, PsaC, which forms a ridge on the stromal surface of PS I with PsaD and PsaE (4). Our previous studies on PS I biogenesis have shown that the insertion of Fe/S clusters is a key step in its post-translational maturation (5, 6). The inability to assemble the F$_{X}$ cluster in a rubA mutant destabilizes PS I and prevents the association of PsaC, PsaD, and PsaE with the membrane-associated portion of the complex. Studies of pscA mutants with modified Cys ligands for F$_{A}$ and F$_{B}$ also point to the importance of Fe/S clusters in the biogenesis and stability of this complex (2).

The biosynthesis of Fe/S clusters is a complex and highly regulated metabolic process. Three major systems for Fe/S cluster biosynthesis have been identified in various organisms: the NIF, ISC, and SUF systems (7, 8). In cyanobacteria and in the chloroplasts of algae and higher plants, oxygen evolution accompanying water oxidation by PS II is one of the major processes associated with the light reactions. Perhaps for this reason, an oxygen-resistant system is essential for Fe/S cluster biogenesis in such organisms (see Ref. 9 for a review). The first evidence connecting the SUF system to PS I biogenesis was obtained through the characterization of pseudorevertants that...
had been isolated from site-specific psaC mutants of *Synechocystis* sp. PCC 6803 (10). These pseudorevertants were able to suppress the inability of certain PsAC variants to assemble Fe/S clusters when non-Cys ligands to Fx and Fy were present in vivo. Characterization of these intergenic suppressors identified the sufr gene, whose product was subsequently shown to encode a negative transcriptional regulator of the suf regulon (11, 12). In many cyanobacterial genomes the major components of the SUF system are organized as an operon, sufBCDS (9). Reverse genetics functional studies of genes for the SUF and ISC systems in *Synechocystis* sp. PCC 6803 and *Synechococcus* sp. PCC 7002 have shown that cyanobacteria primarily rely on the SUF system rather than the ISC system for the biogenesis and maintenance of Fe/S clusters (9).

Studies on the insertion of metalloclusters into the nitrogenase apoproteins in *Azotobacter vinelandii* have established the molecular scaffold model for Fe/S cluster biogenesis (7, 8). Together with NifS, a cysteine desulfurase that provides sulfide for Fe/S cluster assembly, NifU is known to participate in the assembly of nitrogenase metalloclusters (13). *A. vinelandii* NifU is a homodimer that carries two stable [2Fe-2S] clusters (14), and this protein is thought to provide a molecular scaffold for maturation of the nitrogenase metalloclusters (15). Sequence comparisons have revealed three domains in NifU proteins (8). All NifU proteins have nine highly conserved Cys residues: three Cys residues in the N-terminal domain, four Cys residues in the central domain, and two Cys residues in the C-terminal domain. Genetic and biochemical experiments suggest that transient Fe/S clusters are formed in both the N- and C-terminal domains of NifU. NifU-dependent activation of the Fe protein of nitrogenase (NifH) is lost if any of the Cys residues in the N- and C-terminal domains is replaced by alanine (16).

The genomes of all nitrogen-fixing cyanobacteria encode a homolog of the *A. vinelandii* nifU gene, but all cyanobacterial genomes additionally have a gene, nfuA, that encodes a small protein with strong sequence similarity to the C-terminal domain of NifU (see supplemental Fig. S1; for reviews, see Refs. 9 and 17). IscU, a protein with sequence similarity to the C-terminal domain of NifU, has been proposed to function as the scaffold protein for the ISC system (7, 8, 17). Because of their labile Fe/S clusters, both IscA and SufA have been proposed as alternate molecular scaffold proteins for the ISC or SUF systems (18, 19). However, a sufA isca double mutant is viable and exhibits no detectable deficiencies in Fe/S proteins in cyanobacteria (20). In fact, the double deletion mutant grows at higher rates than does the wild-type under iron limitation and grows normally under growth conditions for which the wild-type undergoes chlorosis. Transcription analyses and physiological characterization of wild type and mutant strains lacking SufA, IscA, or both, suggest that SufA and IscA instead play regulatory roles in iron homeostasis, redox sensing, and sensing of the capacity of cells for Fe/S cluster biogenesis (20). In contrast, NfuA is absolutely required for cellular viability (20).

All cyanobacterial NfuA proteins contain a highly conserved CXXC motif that is presumably required for binding an Fe/S cluster. A previous study reported that NfuA (originally named SyNifU) from *Synechocystis* sp. PCC 6803 contained a labile [2Fe-2S] cluster and that this protein could act as scaffold protein in transferring its [2Fe-2S] cluster to a plant-type apoferrredoxin (PetF) (21). Five NfuA-like proteins have been found in the *Arabidopsis thaliana* genome, and three of these proteins (Nfu1, Nfu2, and Nfu3) have been localized to plastids (22). Nfu2 displays the highest sequence similarity to cyano bacterial NfuA proteins and binds a labile [2Fe-2S] cluster *in vitro* (23). A reverse genetics approach has shown that Nfu2 plays an important role in Fe/S cluster biogenesis in plastids (23). A human NfuA-like protein, HiRIP5, has also been reported to carry a [4Fe-4S] cluster (24).

The goals of the current study were to clarify the Fe/S cluster-binding properties of cyanobacterial NfuA and to determine whether this protein might participate in the biogenesis of PS I. We report here our results from the biochemical and spectroscopic characterization of the NfuA protein from *Synechococcus* sp. PCC 7002. Non-denaturing gel electrophoresis indicates that cyanobacterial NfuA forms dimers and tetramers. Biochemical, ERP, and Mössbauer measurements show that cyanobacterial NfuA binds one [4Fe-4S] cluster per dimer. Finally, we demonstrate that the [4Fe-4S] clusters of NfuA can be rapidly and efficiently transferred to the PsaC apoprotein *in vitro* in a reaction that occurs without the release of Fe/S clusters into solution.

**EXPERIMENTAL PROCEDURES**

Construction of the *nfuA* Expression Plasmid—The *nfuA* gene (SYNPC7002_A1413) was amplified by PCR from chromosomal DNA of *Synechococcus* sp. PCC 7002 using the primers 1nfuFN, 5’-TTCGTTCACAAAGACATATGGCATTAGC-3’ and 1nfuRB, 5’-CATGGCAACCCCTAGGGATCCTAAAGAAGT-3’. The purified PCR product was digested with restriction enzymes Ndel and BamHI (sites are underlined above), and cloned into pET16b-*nfuA*, which was verified by DNA sequencing (Penn State Nucleic Acid Facility). The product from pET16b-*nfuA* should produce a recombinant NfuA protein with a poly–His tag (His10) at its N terminus.

*NfuA Overproduction and Purification*—For overproduction of the NfuA protein, the pET16b-*nfuA* plasmid was transformed into *Escherichia coli* strain BL21(DE3). The overexpression of *nfuA* was induced by the addition of 0.1 mm isopropyl 1-thio-β-galactopyranoside to exponentially growing cells in LB medium at 37 °C. Cultures were grown for 4 h after addition of isopropyl 1-thio-β-galactopyranoside. Cells were harvested by centrifugation, and stored at −80 °C until required. For protein purification, cells were resuspended in 50 mm Tris-HCl buffer, pH 8.0, and lysed by passage through a chilled French pressure cell operated at 138 megapascal. The lysed cells were centrifuged at 12,000 × g. The soluble, cell-free extract was loaded onto a column packed with nickel-nitrilotriacetic acid resin (Qiagen, Valencia, CA), and washed with 5 column volumes of 50 mm Tris-HCl buffer at pH 8.0 containing 50 mm NaCl and 20 mm imidazole. NfuA was eluted with 50 mm Tris-HCl buffer at pH 8.0, containing 50 mm NaCl and 250 mm imidazole. The eluted NfuA fraction was immediately dialyzed against 50 mm Tris-HCl buffer, pH 8.0. The protein was concentrated by ultrafiltration with an Amicon concentrator.
equipped with a YM-3 membrane (membrane cut-off of 3 kDa) (Millipore, Billerica, MA).

Chemical Reconstitution of Fe/S Clusters in NfuA—Chemical reconstitution of Fe/S clusters in apo-NfuA was performed as described previously (5, 10, 25). Reconstitution reactions (50 ml) were routinely conducted at a protein concentration of 0.1 to 0.2 mg apo-NfuA ml⁻¹. To prepare a sample for Mössbauer measurement, 4.5 mM ⁵⁷FeCl₃ was used in the reconstitution protocol. Reconstituted holo-NfuA was concentrated by ultrafiltration to 2.5 ml, and loaded onto a PD-10 desalting column (Amersham Biosciences) to remove excess iron, sulfate, and other reagents.

Protein Biochemistry—Quantitative amino acid analysis (Molecular Analysis Facility, University of Iowa) of NfuA was performed in triplicate to obtain a correction factor (0.669) for protein concentrations determined by the Coomassie Plus protein assay (Pierce) with bovine serum albumin as the standard. The protein concentration of an NfuA solution determined by quantitative amino acid analysis (3.20 ± 0.04 mg ml⁻¹) was divided by the estimated protein concentration of the same protein solution from the dye-binding assay with bovine serum albumin as the standard (4.79 mg ml⁻¹) to produce the correction factor 0.669. This correction factor was subsequently applied to all calculations in which protein concentrations were estimated by the dye binding assay.

Quantitation of Fe was performed with ferene (5,5’-(3-(2-pyridyl)-1,2,4-triazine-5,6-diyl)-bis-2-furansulfonic acid) as the detection reagent as described previously (26). A freshly prepared FeCl₃ solution was used to produce a standard curve. The method for quantitation of sulfide has been described (27). A freshly prepared Na₂S solution was used to produce the standard curve. All incubations were performed in an anaerobic chamber with an atmosphere of H₂:CO₂:N₂ (10%:10%:80%, v/v) (Coy Laboratory Products, Grass Lake, MI).

Cross-linking and Alkylation—The methods for protein cross-linking with 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) (28) and protein alkylation with iodoacetamide (29) have been described previously. Cross-linking and alkylation of apo-NfuA and holo-NfuA were performed in the anaerobic chamber to avoid oxidation of Cys residues.

Gel Electrophoresis—Methods for polyacrylamide gel electrophoresis in the presence of SDS-PAGE and immunoblotting have been described (5, 12). Native gel electrophoresis under anoxic conditions was performed in the anaerobic chamber. Non-denaturing gels (9% (w/v) acrylamide for the separating gel and 4% (w/v) acrylamide for the stacking gel) were prepared in the same way as for SDS-PAGE except that SDS was omitted from all buffers. All buffers were extensively sparged with N₂ gas to remove oxygen. Apo-NfuA and holo-NfuA samples were prepared under anoxic conditions in loading buffer without SDS and 2-mercaptoethanol. Gels were stained for iron as described previously (30). The iron-stained gel was scanned, after which the gel was stained with Coomassie Blue.

Low-temperature X-band EPR Spectroscopy of Holo-NfuA—The method for low-temperature, X-band EPR spectroscopy has been described (31). Measurements were performed using an ECS-106 X-band spectrometer (Bruker Biospin, Billerica, MA) equipped with an ESR900 liquid helium cryostat and an ITC-4 temperature controller (Oxford Instruments, Billerica, MA). The microwave frequency was determined with a Hewlett-Packard 5340A frequency counter. The spectrometer parameters were: temperature, 16 K; microwave frequency, 9.47 GHz; receiver gain, 2 × 10⁶; modulation amplitude, 10 gauss at 100 kHz. Spectra were obtained by averaging 10 scans. Holo-NfuA was prepared for EPR analysis by inclusion of 10 mM sodium dithionite, and 100 mM glycine at pH 10.0. The baseline was generated from a blank sample, which was prepared in the same manner as the NfuA samples except that buffer replaced the protein solution. The spectrum from the blank was subtracted from the spectrum of reduced NfuA.

Mössbauer Spectroscopy—The protocol for Mössbauer spectroscopy has been described (12). Spectra were recorded at a sample temperature of 4.2 K that was maintained by a liquid helium cryostat. The sample was kept inside an SVT-400 Dewar (Janis, Wilmington, MA) and a magnetic field of 53 millitesla was applied parallel to the γ-beam. The isomer shifts quoted are relative to the centroid of the spectrum of metallic Fe foil at room temperature. Data analysis was performed using the program WMOSS from WEB Research (Edina, MN).

PS I Reconstitution and Assay by Light-induced EPR Spectroscopy—The method for reconstitution of PS I complexes and the conditions for the light-induced EPR spectroscopy have been described (32). P700 F₅X-cores were prepared from purified PS I complexes of Synechococcus sp. PCC 7002 following the method described (25). Recombinant PsaC and PsaD were overproduced in E. coli and purified as described (33). For light-induced EPR spectroscopic studies of Fe/S clusters transfer from holo-NfuA to apo-PsaC, samples were prepared with 20 μM P700 F₅X-cores, 400 μM apo-PsaC, 400 μM PsaD, 840 μM holo-NfuA, 10 mM ascorbate, 10 μM 2,6-dichrophenolindophenol (DCPIP), 4 mM dithiothreitol (DTT), and 0.04% (w/v) n-dodecyl-β-D-maltoside (DM), and frozen under low light. The baseline was recorded in the dark, and spectra were recorded before and after illumination with an argon ion laser operated at 2.5 watts in all-lines mode with a 3-fold beam expander to fill the resonator grid.

PS I Reconstitution and Assay by Room Temperature Optical Kinetic Spectroscopy in the Near-IR Region—Transient absorbance changes in the near-IR were measured for reconstituted PS I complexes with a laboratory-built, double-beam spectrophotometer as described (6, 31), except that the cuvettes used were 2 × 10 mm instead of 5 × 10 mm. The samples were prepared under anoxic conditions, with the following final concentrations: 1 μM P700-F₅X cores, 20 μM apo-PsaC, 20 μM PsaD, 42 μM holo-NfuA, 10 mM DCPIP, 10 mM ascorbate, 4 mM DTT, and 0.04% (w/v) DM. The concentrations of apo-PsaC, PsaD, and the P700 F₅X-cores were determined according to the extinction coefficients described previously (32, 33). The concentration of holo-NfuA was calculated from the [4Fe-4S] cluster content as described (25).

RESULTS

Overproduction and Purification of NfuA—The nfuA gene from Synechococcus sp. PCC 7002 was cloned into pET11b and overexpressed to produce a recombinant protein with a His₁₀ tag at its N terminus. As determined by SDS-PAGE, purified
NfuA had an apparent mass of ~11 kDa (Fig. 1A, lane 3), which matched the predicted mass of 11.1 kDa (NfuA (8558 Da) + His tag (2538 Da) = 11.1 kDa). Apo-NfuA was purified from the soluble fraction of the whole cell lysate, which had a light brown color; this suggested that Fe/S clusters might be present in the NfuA protein. However, based upon the loss of the brownish color and a decrease in absorbance at about 420 nm, these Fe/S clusters were labile and were lost during the purification procedure. Therefore, it was necessary to reconstitute the Fe/S clusters of NfuA in vitro with inorganic reagents under anoxic conditions.

The absorption spectrum of chemically reconstituted holo-NfuA (Fig. 2, solid line) showed broad absorbance bands between 300 and 600 nm that are typical of sulfur → iron charge transfer bands of Fe/S proteins. The absence of maxima in the 450- and 550-nm regions suggested that holo-NfuA harbored [4Fe-4S] clusters. The stability of the Fe/S clusters in holo-NfuA was examined by exposing holo-NfuA to oxygen or 10 mM EDTA (data not shown). Exposure of holo-NfuA to air levels of oxygen led to a 50% decrease in absorbance at about 420 nm, these Fe/S clusters were labile and were lost during the purification procedure. Therefore, it was necessary to reconstitute the Fe/S clusters of NfuA in vitro with inorganic reagents under anoxic conditions.

EPR Spectroscopy of Reconstituted NfuA—A low-temperature, X-band EPR spectrum of dithionite-reduced holo-NfuA is presented in Fig. 3. At 16 K and a microwave power of 40 milliwatts, NfuA had an axial spectrum with g values of 2.037 and 1.903. The inset depicts the temperature dependence of the EPR signal. At constant microwave power, the intensity of the EPR signal was maximal at 10 to 16 K and decreased dramatically above 20 K. These spectroscopic properties suggested that chemically reconstituted holo-NfuA harbored [4Fe-4S]1+ cluster(s).

Mössbauer Spectroscopy of the Holo-NfuA—To obtain further evidence for the presence of [4Fe-4S] cluster(s) in holo-NfuA, the protein was reconstituted with 57Fe2+ and studied by Mössbauer spectroscopy. The spectrum of 57Fe-holo-NfuA was recorded at 4.2 K in a 53-millitesla external magnetic field oriented parallel to the γ-beam. The vertical bars in Fig. 4 are the Mössbauer spectrum of oxidized holo-NfuA, and the overlaid
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solid line shows a simulation of the experimental data. This spectrum was deconvoluted to assess the contributions from iron in different forms to the overall spectrum (Fig. 4, lines A–D). Line A represents the quadrupole doublet arising from [4Fe-4S]2+ clusters. Line B represents the iron quadrupole doublet attributed to [2Fe-2S]2+ clusters; it had δ = 0.28 mm/s and ΔE_Q = 0.64 mm/s, and accounted for 11% of the iron in the reconstituted sample. The broad line at ~2–3 mm/s is typical of high-spin Fe(II), and the broadness suggests the presence of several distinct Fe2+ species. Lines C and D are quadrupole doublet simulations with parameters typical of six-coordinate, N/O-ligated Fe2+ and four-coordinate, sulfur-ligated Fe2+ species; these account for 17 and 11% of the total iron, respectively. No signals were detected by low-temperature EPR spectroscopy of the identical oxidized holo-NfuA sample that was studied by Mössbauer spectroscopy (data not shown). These data indicate that most of the Fe/S clusters in holo-NfuA were [4Fe-4S] clusters, although the data suggests the presence of a small fraction of [2Fe-2S] clusters.

NfuA Forms Oligomers That Carry One [4Fe-4S]2+ Cluster Per Dimer—To examine the oligomerization state of NfuA, apo-Nfu and reconstituted holo-NfuA were analyzed by native PAGE under anoxic conditions. After electrophoresis, two forms of holo-NfuA were detected by iron staining (Fig. 1B, lane 1); however, a third form of the protein was detected after staining with Coomassie Blue (Fig. 1B, lane 2). By comparing the electrophoretic migration of NfuA as a function of acrylamide concentration (data not shown), it was concluded that the three observed forms of NfuA were most likely monomeric apo-NfuA and dimeric and tetrameric holo-NfuA. As can been seen in Fig. 1B, the approximate ratio of these three NfuA forms was about 5:20:75, respectively. For apo-NfuA, two major bands, corresponding to monomeric (major) and dimeric (minor) forms, were resolved after native gel electrophoresis (supplemental Fig. S2A). Only a very faint band with the apparent migration of a tetramer was observed for apo-NfuA (supplemental Fig. S2A).

The oligomerization of apo-NfuA and holo-NfuA was studied further by cross-linking with the zero-length cross-linking reagent EDC and by using the alkylation reagent iodoacetamide, which was employed to prevent the formation of intermolecular disulfide bonds. As shown in Fig. 1A, lane 3, NfuA monomers, dimers, and trimers and/or tetramers were sometimes observed after SDS-PAGE of NfuA samples that had not been alkylated prior to SDS-PAGE. However, alkylation of the Cys residues with iodoacetamide prior to SDS-PAGE largely prevented the formation of these oligomers (Fig. 1C, lane 3). When holo-NfuA was subjected to cross-linking with EDC followed by alkylation, monomers, dimers, and tetramers were observed after SDS-PAGE (Fig. 1C, lane 4). NfuA dimers and tetramers were more abundant after EDC treatment of the holo-NfuA sample, but apo-Nfu dimers were also observed after EDC treatment (supplemental Fig. S2B). Based upon the results from native PAGE under anoxic conditions and from cross-linking of holo-NfuA with EDC, we concluded that both apo-NfuA and holo-NfuA could form dimers. However, in the presence of Fe/S clusters holo-NfuA occurred in solution as a mixture of dimers (minor form) and tetramers (major form).

Chemical analyses of holo-NfuA for non-heme iron and sulfide yielded ~2.39 iron and ~1.87 sulfides per NfuA monomer. Each NfuA polypeptide has only two conserved Cys residues that could serve as ligands to Fe/S clusters (supplemental Fig. S1); this suggests that a NfuA dimer is required to ligate an Fe/S cluster. Combining the chemical analyses with the results from EPR and Mössbauer spectroscopy, chemically reconstituted holo-NfuA contained ~0.73 [4Fe-4S]2+ and ~0.12 [2Fe-2S]2+ clusters per dimer. In summary, the data showed that holo-NfuA formed dimers and tetramers that predominantly harbored one [4Fe-4S]2+ cluster per homodimer.

Reconstitution of PS I Complexes by Cluster Transfer from Holo-NfuA to Apo-PsaC—Previous mutagenesis studies have shown that NfuA is essential in cyanobacteria, and other studies have suggested that NfuA might play an essential role in the assembly of Fe/S clusters (20). We next tested whether NfuA could transfer Fe/S clusters to apo-PsaC, which ligates two [4Fe-4S] clusters (1). In the presence of PsaD, Fe/S cluster transfer to apo-PsaC can be monitored by reconstitution of long-lived electron transfer in P700-FX cores (34). Optical kinetic spectroscopy can be used to monitor electron transfer in the reconstituted PS I complexes (35). In these measurements, a single turnover laser flash induces an absorbance increase at 820 nm, which arises from the P700+ Chl cation. The decay
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kinetics of this absorbance provides information about the electron acceptor from which the charge recombination occurs (35). Because binding of holo-PsaC to P700-Fx cores is rapid, the recovery of long-lived charge separation on a laser flash provides a convenient and accurate assay for the insertion of the [4Fe-4S] clusters into PsaC.

Fig. 5 compares the decay kinetics of the P700+ cation absorbance for P700-Fx core complexes incubated with various combinations of PsaC, PsaD, and NfuA. As shown in Fig. 5A, trace 1, and supplemental Fig. S3A, four kinetic phases are observed in the decay of the 820-nm absorbance after P700-Fx cores had been incubated with apo-PsaC and PsaD. The 51-μs phase was assigned to the backreaction of A1+ with P700+; these represent PS I cores in which Fx+ has been inadvertently denatured. The 643-μs and 4.3-ms phases were assigned to charge recombination between Fx- and P700+. The remaining low-amplitude phase, which had a time constant of >100 ms, was sensitive to DCPIP concentration, allowing this phase to be assigned to electron donation from DCPIP to P700+ (data not shown). In PS I complexes showing this latter kinetic phase, electrons had been transferred to oxygen and thus were no longer available for charge recombination with P700+. Because PsaD is required for stable binding of PsaC to the stromal surface of PS I (33), very similar results were obtained when P700-Fx cores were incubated with holo-PsaC in the absence of PsaD (Fig. 5A, trace 2).

When P700-Fx cores were incubated with holo-PsaC and PsaD, electron transfer to Fx+ and Fx_ was restored and long-lived charge separation occurred (Fig. 5A, trace 4). The reconstituted PS I complexes also exhibited four kinetic phases, but a major new kinetic phase, corresponding to charge recombination between [FA/Fb]+ and P700+, was observed at ~59 ms (Fig. 5A, trace 4; supplemental Fig. S3B). The two slowest kinetic phases (59 ms and 844 ms) accounted for ~60% of the total absorbance change and were assigned to PS I complexes in which electrons had been transferred to FA/Fb prior to charge recombination with P700+ or reduction of oxygen.

When P700-Fx cores were incubated with apo-PsaC, PsaD, and holo-NfuA, electron transfer to FA/Fb was also restored, resulting in long-lived charge separation (Fig. 5A, trace 5). These reconstituted PS I complexes exhibited four kinetic phases, and their decay kinetics were very similar to those of P700-Fx cores that had been reconstituted with holo-PsaC and PsaD (compare Fig. 5A, traces 4 and 5, and supplemental Fig. S3C). The two slowest kinetic phases together accounted for ~53% of the total absorbance change and were assigned to PS I complexes in which the electron had been transferred to FA/Fb prior to charge recombination with P700+ or reduction of oxygen. In other similar experiments, these slow kinetic phases accounted for as much as 60–70% of the total absorbance change. These results clearly showed that the [4Fe-4S] clusters could be transferred from holo-NfuA to the FA and FB sites in apo-PsaC. In these experiments, the molar ratio of [4Fe-4S] clusters to PsaC was 2.1:1, and much higher molar ratios (5-, 10-, and 20-fold) of [4Fe-4S] clusters to PsaC did not significantly improve the reconstitution as measured by charge recombination from [FA/Fb]+ to P700+ (data not shown).

Thus, cluster transfer from holo-NfuA to apo-PsaC appeared to be nearly stoichiometric. As shown in Fig. 5B (trace 1), the ability of holo-NfuA to transfer Fe/S clusters to PsaC was completely abolished when holo-NfuA was denatured by boiling for 1 min. However, when the reagents for chemical reconstitution of Fe/S proteins were boiled prior to mixing with apo-PsaC, PsaD, and P700-Fx cores, reconstitution of the PS I complexes still occurred (Fig. 5B, trace 2). This experiment clearly suggested that interactions between holo-NfuA and apo-PsaC...
were required during transfer of Fe/S clusters from holo-NfuA to apo-PsaC.

The standard reconstitution buffer for cluster transfer from holo-NfuA to apo-PsaC contained 4 mM DTT. DTT could be acting as a reductant, a thiol reagent, or both in the cluster-transfer reaction. When DTT was omitted from the reaction mixture, very little (<12%) charge recombination occurred from electron carriers beyond F_X (data not shown); thus, the Fe/S clusters were not transferred from holo-NfuA to apo-PsaC. As described above, the charge recombination between \([F_A/F_B]\) and P700\(^+\) accounted for >55% of the total absorption change in the presence of 4 mM DTT. However, DTT could be replaced by the non-thiol reductant, \(tris(2\text{-carboxyethyl})\)phosphine. When the cluster transfer from holo-NfuA to apo-PsaC was performed in the presence of 1 mM \(tris(2\text{-carboxyethyl})\)phosphine, the 121-ms phase, which was assigned to charge recombination between \([F_A/F_B]^-\) and P700\(^+\), accounted for ∼44% of the total absorbance change (data not shown). Presently it is not clear whether DTT and \(tris(2\text{-carboxyethyl})\)phosphine increase the efficiency of the reconstitution reaction by acting as reducing agents for the Fe/S clusters or whether these reducing agents improve the efficiency of cluster transfer by reducing disulfide bonds in apo-PsaC that interfere with cluster transfer, or both. However, because cluster transfer occurs in the presence of the non-thiol reductant, \(tris(2\text{-carboxyethyl})\)phosphine, thiol reagents are clearly not required for cluster transfer from holo-NfuA to apo-PsaC.

**Holo-NfuA Rapidly and Efficiently Transfers Fe/S Clusters Directly to Apo-PsaC Without Cluster Release**—The relative efficiencies of Fe/S cluster insertion into apo-PsaC by holo-NfuA and by chemical reconstitution were compared in the following experiment. As shown in Fig. 6 (trace 2), PS I complexes were reconstituted with P700-F_X cores, PsaD, apo-PsaC, and holo-NfuA. The decay kinetics of P700\(^+\) were measured at 820 nm after an incubation of only 10 min, which was approximately the minimal time to mix the reconstitution reagents in the glove box and transfer the reaction mixture to the spectrophotometer. In this experiment, the ∼30-ms phase corresponding to electron transfer from \([F_A/F_B]\) to P700\(^+\) accounted for ∼56% of the total absorbance change. This result showed that holo-NfuA rapidly transferred \([4\text{Fe-4S}]\) clusters to apo-PsaC. Increasing the incubation time (30 min, 1 h, 2 h, and 16 h) did not lead to a significant increase in cluster transfer (data not shown).

Dos Santos et al. (16) compared the activation of the nitroge

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**FIGURE 6. Flash-induced absorption changes for P700\(^+\) reduction at 820 nm.** Decay kinetics of 1 μM P700-F_X cores, 20 μM apo-PsaC, 20 μM PsaD, 4 mM DTT, 0.04% (v/v) DM, 84 μM Fe\(^{2+}\), 84 μM sulfide, 10 μM DCPIP, and 10 mM ascorbate following a 1-h incubation; 2 μM P700-F_X cores, 42 μM holo-Nfu, 20 μM apo-PsaC, 20 μM PsaD, 0.04% (v/v) DM, and 4 mM DTT. After a 10-min incubation, 10 μM DCPIP and 10 mM ascorbate were added and flash-induced kinetics at 820 nm were measured. The discontinuity in the noise level is a consequence of fusing two kinetic traces taken at different time bases on the same laser flash.
concentration was \( \approx 33 \text{ mM} \). This experiment implied that cluster transfer from holo-NfuA to apo-PsaC required a direct interaction between the two proteins.

**Light-induced EPR Spectra of PS I Complexes Reconstituted with Holo-NfuA**—The \( F_A \) and \( F_B \) clusters of PsaC have characteristic EPR resonances that can easily be detected when PS I complexes are illuminated at low temperature (36). As shown in Fig. 8 (trace 1), P700-\( F_X \) cores that were frozen at low light intensity exhibited only weak EPR signals after illumination at 16 K. However, P700-\( F_X \) cores that had been reconstituted with holo-NfuA, apo-PsaC, and PsaD had the characteristic EPR resonances of reduced \( F_A \) and \( F_B \) when the complexes were illuminated at low temperature (Fig. 8, trace 2). Consistent with the results from optical kinetic spectroscopy, this result showed that holo-NfuA was able to transfer [4Fe-4S] clusters to apo-PsaC and that both of these clusters could be photoreduced.

**DISCUSSION**

The NifU/Nfu Family in Cyanobacteria—Through studies in various organisms from all three kingdoms of life, three major systems for Fe/S cluster biogenesis are now recognized: the NIF, ISC, and SUF systems (7–9). Although these systems may have different biochemical mechanisms for Fe/S cluster assembly and are differently regulated, each system apparently requires a cysteine desulfurase and the participation of a “scaffold” protein for the assembly and transfer of Fe/S clusters. The NifU protein family is generally believed to play a central role in Fe/S cluster formation and transfer in the NIF system, which functions in the maturation of nitrogenase (7, 8, 16). It has been proposed that this scaffold protein provides an intermediate site for the assembly of Fe/S clusters or Fe/S cluster precursors, and there is strong biochemical and genetic evidence that NifU can transfer Fe/S clusters to NifH, the iron-protein of nitrogenase (7, 8).

Although all cyanobacteria perform oxygenic photosynthesis, the cyanobacteria can further be divided into two major groups on the basis of their ability to perform nitrogen fixation. The genomes of all nitrogen-fixing cyanobacteria encode a nifU gene that is clustered together with other nif genes. Cyanobacterial NifU proteins are highly similar (~49% identity and 63% similarity) to *A. vinelandii* NifU, and they form a distinctive
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clade of sequences within the NifU/NfuA sequences (supplemental Fig. S4).

All cyanobacterial genomes contain a single nfuA gene, and NfuA proteins, which have strong sequence similarity to the C terminus of NifU, also form a discrete clade in phylogenetic analyses (supplemental Fig. S4). Because efforts to inactivate nfuA inevitably led to the generation of meridiploids, we previously reported that NfuA is essential for viability in cyanobacteria (20). Because nfuA was essential even in cells for which the psaAB genes had been deleted, it is clear that NfuA is required for functions that extend beyond the assembly of PS I (see below). NfuA-like proteins have recently been identified in several non-photosynthetic organisms, including E. coli (37), A. vinelandii (38), Saccharomyces cerevisiae (39), and humans (24). In contrast to the situation in cyanobacteria, an S. cerevisiae nfu1 mutant had no obvious growth defect (39). In oxygenic photosynthetic cyanobacteria and the chloroplasts of higher plants, the SUF system is the major machinery for Fe/S cluster assembly (9). NfuA-like proteins have been suggested to be involved in Fe/S cluster biogenesis in the chloroplasts of Arabidopsis thaliana (22). Because several proteins of the ISC system are not essential for Fe/S biogenesis in cyanobacteria, it is reasonable to conclude that NfuA functions together with the SUF machinery in Fe/S cluster assembly. Interestingly, wild type A. vinelandii can grow at an oxygen concentration twice that of the atmosphere, but an nfuA mutant was unable to grow under these conditions (38). This observation is consistent with the idea that NfuA might participate in an Fe/S cluster biogenesis pathway that is relatively insensitive to oxygen.

The genomes of most nitrogen-fixing cyanobacteria also contain a third member of the NifU/Nfu family of proteins, which we designate here as NfuB (supplemental Fig. S4). Members of the NfuB clade have sequence similarity to the C-terminal domain of NifU, have Rieske domains at their C termini, and contain a third member of the NifU/Nfu family of proteins, which have strong sequence similarity to the C terminus of NifU, also form a discrete clade in phylogenetic analyses (supplemental Fig. S4). Because efforts to inactivate nfuA inevitably led to the generation of meridiploids, we previously reported that NfuA is essential for viability in cyanobacteria (20). Because nfuA was essential even in cells for which the psaAB genes had been deleted, it is clear that NfuA is required for functions that extend beyond the assembly of PS I (see below). NfuA-like proteins have recently been identified in several non-photosynthetic organisms, including E. coli (37), A. vinelandii (38), Saccharomyces cerevisiae (39), and humans (24). In contrast to the situation in cyanobacteria, an S. cerevisiae nfu1 mutant had no obvious growth defect (39). In oxygenic photosynthetic cyanobacteria and the chloroplasts of higher plants, the SUF system is the major machinery for Fe/S cluster assembly (9). NfuA-like proteins have been suggested to be involved in Fe/S cluster biogenesis in the chloroplasts of Arabidopsis thaliana (22). Because several proteins of the ISC system are not essential for Fe/S biogenesis in cyanobacteria, it is reasonable to conclude that NfuA functions together with the SUF machinery in Fe/S cluster assembly. Interestingly, wild type A. vinelandii can grow at an oxygen concentration twice that of the atmosphere, but an nfuA mutant was unable to grow under these conditions (38). This observation is consistent with the idea that NfuA might participate in an Fe/S cluster biogenesis pathway that is relatively insensitive to oxygen.

The genomes of most nitrogen-fixing cyanobacteria also contain a third member of the NifU/Nfu family of proteins, which we designate here as NfuB (supplemental Fig. S4). Members of the NfuB clade have sequence similarity to the C-terminal domain of NifU, have Rieske domains at their C termini, and contain an additional N-terminal domain, which we designate here as NfuB (supplemental Fig. S4). Mem-

which resembles an A-type scaffold domain, and, unlike the results reported here, the C-terminal domain alone was not active in cluster transfer (37). *E. coli* nfuA null mutants had oxidative stress and iron starvation phenotypes, which suggested that NfuA participates in both ISC- and SUF-dependent Fe/S cluster biogenesis. The nfuA gene is found in all cyanobacterial genomes, is essential for viability, and cannot be inactivated in strains in which the psaAB genes have been deleted (20). Thus, it is highly likely that NfuA is involved in Fe/S cluster transfer to proteins that are not involved in photosynthesis. Moreover, its product can transfer [2Fe-2S] clusters to apo- (PetF)-ferredoxin (21, 23) and [4Fe-4S] clusters to PsacC. Therefore, we propose that NfuA plays a central and ubiquitous role in Fe/S cluster biogenesis and iron homeostasis in cyanobacteria, and related proteins probably play similar roles in *E. coli* and many other organisms.

It is possible that redundant pathways for Fe/S cluster insertion into cyanochemical proteins exist. Picciocchi et al. (43) recently showed that cyanobacteria have a monothiol glutaredoxin, which ligates one intrasubunit [2Fe-2S] cluster per dimer by single cysteines on each subunit in combination with two glutathione ligands. The SyGx3p-like protein of *Synechocystis* sp. PCC 6803, the product of open reading frame 7184, can complement the Fe/S cluster biogenesis defect of a yeast mitochondrial mutant lacking ScGx3p (44). Another recent study suggested that two chloroplast-targeted monothiol glutaredoxins, GrxS14 and GrxS16, transfer their [2Fe-2S] clusters to chloroplast apo-ferredoxin (45). Finally, recent studies suggest that IscU can harbor either two [2Fe-2S] clusters or one [4Fe-4S] cluster but only the latter can be transferred to apo-aconitase to activate this enzyme (46, 47). Given the wide range of Fe/S proteins that exist in biological systems and the importance of these proteins to cellular metabolism and physiology, it is not surprising that nature has apparently devised multiple pathways to produce these essential prothetic groups.

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