FeoC from *Klebsiella pneumoniae* contains a [4Fe-4S] cluster

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Abbreviations used: Feo, ferrous iron transporter; GST, glutathione S-transferase; GDI, GDP dissociation inhibitor; EPR, electron paramagnetic resonance; NMR, nuclear magnetic resonance; EXAFS, extended X-ray absorption fine structure; XANES, X-ray absorption near-edge structure; DTT, dithiothreitol.

Abstract

Iron is essential for pathogen survival, virulence, and colonization. Feo is suggested to function as the ferrous iron (Fe²⁺) transporter. The enterobacterial Feo system is composed of 3 proteins: FeoB is the indispensable component and is a large membrane protein likely to function as a permease; FeoA is a small src-homology 3 (SH3) domain protein that interacts with FeoB; FeoC is a winged-helix protein...
containing 4 conserved Cys residues in a sequence suitable for harboring a putative iron-sulfur (Fe-S) cluster. The presence of an iron sulfur cluster on FeoC has never been shown experimentally. We report that under anaerobic conditions, the recombinant *Klebsiella pneumoniae* FeoC (*KpFeoC*) exhibited hyperfine-shifted NMR resonances and a UV-VIS absorbance spectra characteristic of a paramagnetic center. The EPR and EXAFS results were consistent only with the [4Fe-4S] clusters. Substituting the cysteinyl sulfur with oxygen resulted in significantly reduced cluster stability, establishing the roles of these cysteines as the ligands for the Fe-S cluster. When exposed to oxygen, the [4Fe-4S] cluster degraded to [3Fe-4S] and eventually disappeared. We propose that *KpFeoC* may regulate the function of the Feo transporter through the oxygen or iron sensitive coordination of the Fe-S cluster.

**INTRODUCTION**

Iron is an essential element for nearly all life forms (1-3). However, free cellular iron is toxic and the solubility of ferric iron is poor. Thus, bacteria tightly regulate cellular iron levels through multiple iron transport pathways to achieve effective homeostasis (3-5). Feo, the *ferrous* iron (Fe$^{2+}$) transport system, is likely a major route for transporting ferrous iron across the bacterial membrane under anaerobic or low pH conditions, such as those in the gastrointestinal tract (6). Several systems have demonstrated the importance of Feo. Feo is critical for both survival and virulence in *Helicobacter pylori* (7). The Feo system is critical for virulence in *Streptococcus suis* (8) and for colonization in *E. coli* and *Salmonella typhimurium* (9,10). Bacterial pathogens require the Feo system for enhanced colonization (1,3,6).

The *feo* operon was first identified in *E. coli* K-12 and its expression was shown to be under dual transcriptional control by the iron-sensing Fur (ferric uptake regulator) and oxygen-sensing FNR (fumarate nitrate reduction protein) regulators in response to different levels of iron and oxygen (6,11). The *feo* operon from *γ*-proteobacteria encodes for 3 proteins: FeoA, FeoB, and FeoC (11-13). FeoA,
present in 90% of the *feo* operons, is a small src-homology 3 (SH3) domain protein necessary for ferrous iron transport (6,12,14-16). Recent enzymatic assays have suggested that FeoA may not act as a GTPase-activating protein as originally proposed (14). FeoB, an indispensable component of the Feo system, is a large protein consisting of an intracellular amino-terminal domain (NFeoB) and a transmembrane carboxyl-terminal domain presumed to form the Fe$^{2+}$ pore function that functions as a permease. NFeoB consists of a small-GTPase (G) domain and a GDP dissociation inhibitor (GDI)-like helical domain. The crystal structures of several forms from *E. coli* and *Klebsiella pneumoniae* NFeoB assemble into a funnel-like trimer with a cytoplasmic pore that could facilitate gating and passage of un-hydrated ferrous irons (17,18). The G domain exhibits guanosine nucleotide-dependent conformational changes in Switch I, Switch II, and G5 motifs of the G domain, and changes in the distance between G and GDI-like domains. The conformational changes were suggested to trigger the opening (GTP bound) and closing (GDP bound) of the pore and to regulate Fe$^{2+}$ transport (17).

FeoC is a small hydrophilic protein that is present in only 15% of the *feo* operons (14). The solution structures of FeoC from *K. pneumoniae* and *E. coli* possess a winged-helix structure often associated with DNA binding (19). The long, disordered wing-loop-1 (W1) contains 4 conserved Cys residues in a sequence, Cx_{4}CxxCx_{5-8}C, suitable for harboring a putative iron-sulfur (Fe-S) cluster (6,19). Thus, FeoC was proposed to be an iron-sulfur cluster-dependent transcriptional regulator directly controlling the expression of the *feo* operon (transcriptional regulator model) (6). However, a recent study suggested that FeoC did not regulate the *feo* promoter in *Yersinia pestis* (20) and no report has confirmed the DNA binding activity of FeoC.

Two studies have recently suggested that FeoC may function at the post-translational level. We showed that apo-*Kp*FeoC binds to the N-terminal domain of *Kp*FeoB (*Kp*NFeoB) with high affinity (21). In the crystal, apo-*Kp*FeoC binds to *Kp*NFeoB at a site encompassing the Switch II region of the G domain and the C-terminal GDI-like domain such that the flexible W1 loop is potentially capable of interacting with residues in the nucleotide-binding site. We proposed that FeoC might coordinate the
Fe-S cluster to regulate ferrous iron transport by modulating G-protein activity (G-protein modulator model). However, Kim et al. found that FeoC binds to FeoB and the presence of FeoC prevents FeoB from proteolytic degradation by FtsH in Salmonella enterica under low iron and low oxygen conditions (22). This results in an elevated level of FeoB that enables Salmonella to take up Fe (II) under anaerobic and low iron conditions (protease inhibitor model). The coordination of the Fe-S cluster on FeoC can play crucial roles in all 3 models. This study is the first to provide experimental evidence supporting the existence of an Fe-S cluster on FeoC. We present spectroscopic and mutational data proving its existence, the range of redox potentials, and the degradation of a [4Fe-4S] cluster on KpFeoC, with discussions regarding the models.

**MATERIALS AND METHODS**

**Chemicals, Bacterial Strains, and Vectors**

[99% \(^{15}\)N]H\(_4\)Cl, [99% \(^{13}\)C]-D-glucose, 99% D\(_2\)O, and sodium 2, 2-dimethyl-2-silapentane-5-sulfonate (DSS) were purchased from Cambridge Isotope Laboratories (Andover, MA). BME vitamins, redox reagents, and corresponding antibiotics were purchased from Sigma Aldrich (St. Louis, MO). Isopropyl \(\beta\)-D-thiogalactoside (IPTG) was purchased from MDBio Inc. (Taipei, Taiwan). Air-tight Hellma UV-VIS cuvettes (114B-QS) were purchased from Sigma Aldrich (St. Louis, MO). The E. coli strain BL21 (DE3) pLysS was purchased from Novagen (Madison, WI). The expression vector, pGEX-6p-1 carrying the cDNA encoding a chimera fusion protein of the glutathione S-transferase (GST) tag and the FeoC from K. pneumoniae (subsp. pneumoniae NTUH-K2044) (pGEX-6p-1 /GST-KpFeoC) was prepared in this lab. The fusion protein was first purified through the GST-affinity column. The enzyme was digested to remove the GST tag and further purified through the size-exclusion column (19). For preparing the holo-FeoC, the protein was purified through the GST column and concentrated aerobically, whereas the follow-up enzyme digestion was performed anaerobically (less than 5 ppm oxygen) in an anaerobic chamber from COY lab (Grass Lake, MI) and purified through a Superdex-75
10/300 column attached to an ÄKTA purifier (GE Healthcare) inside the anaerobic chamber. Samples were kept at low temperature in a labtop cooler from Nalgene/Sigma Aldrich (St. Louis, MO). All point mutants were cloned with the GST tag in this lab.

**Protein Expression and Labeling**

The *KpFeoC* protein was prepared by growing BL21 (DE3) pLysS cells carrying the pGEX-6p-1/GST-*KpFeoC* vector in LB medium without supplementing additional iron. Uniformly $^{15}$N labeled *KpFeoC* ([u-$^{15}$N]-*KpFeoC*) protein was prepared by growing cells in the M9 medium supplemented with 1 g/L $^{15}$NH$_4$Cl in the presence of 50 mg/L (w/v) FeCl$_3$ and 1% (v/v) BME vitamins, as described previously (18,19,21). For preparing $[U-^{13}$C, $^{15}$N]-*KpFeoC* protein, $[U-^{13}$C]-glucose was added in 2 aliquots; 2 g/L at the onset and 2 g/L when OD$_{600}$ reached 2.5 (immediately after IPTG induction).

Based on optical spectra analysis, the holo-*KpFeoC* expressed in the LB medium is identical to that expressed in the M9 media with iron. The addition of iron typically increases the yield and homogeneity.

**NMR and EPR Sample Preparation**

Amicon (Millipore, Billerica, MA) tubes were used for protein concentration and buffer exchange following protein purification. Unless specified, the NMR (and EPR) buffers were 50 mM Tris, and 100 mM NaCl, in 9% D$_2$O at pH 7.8 for *KpFeoC*. We defined the native state of FeoC as freshly purified protein from *E. coli* in the anaerobic chamber. Unused samples were stored in bottles sealed with septa caps at -80°C. The DSS was added as the internal chemical shift standard. Excess molar ratios of sodium dithionite or dithiothreitol (DTT) were added to the reduced samples. Unless specified, the reduced state is the dithionite reduced state (for NMR and EPR). NMR samples in the reduced state were prepared in an anaerobic chamber and transferred to NMR tubes with a J. Young valve (Wilmad, Vineland, NJ). EPR samples were prepared and frozen in the anaerobic chamber. Spectra were taken aerobically as solids. The pH values were measured before entering the chamber. *Apo-KpFeoC* samples...
were prepared by dialyzing away irons from \textit{holo-KpFeoC}, which had been exposed to oxygen for more than 5 d.

**NMR Spectroscopy, data processing, and analysis**

NMR spectra were acquired at specified temperatures on Bruker Avance 500 or 600 MHz spectrometers equipped with triple resonance cryogenic probes, as described previously (19). To observe the fast-relaxing hyperfine shifts, the repetition times of superWEFT sequences were set to 0.1 s for protons and 0.2 s for carbons and fine-tuned before measurements. Spectra were signal averaged for 160,000 scans for carbons and 40,000 scans for protons for a total acquisition time of 8 h for carbon and 2 h for protons. Spectra were obtained by subtracting 2 spectra with broad and desired line broadening (typically 60 Hz for protons and 600 Hz for carbons) and were baseline corrected using spline protocol. Proton chemical shifts were referenced relative to internal DSS (taken as 0 ppm); carbon and nitrogen spectra were referenced indirectly by the canonical ratios (23). All spectra were processed by Topspin software (Bruker).

**EPR**

Unless specified, the EPR sample buffer conditions were identical to those used in NMR experiments. The EPR spectra were acquired at 9.5390 GHz (measured using a Hewlett-Packard 5246L electronic counter) and 4 Gauss modulations with 20 mW power at a 4096 point resolution and the average of 4 scans was reported. To avoid loss of EPR signals caused by exposure to air, the sodium dithionite reduced samples were transferred to the EPR tube and frozen within 10 s in the anaerobic chamber. To ensure full reduction, the reductant/protein molar ratios exceeded 4 and 8 for DTT and dithionite, respectively. Cavity signals were subtracted from the EPR spectra by exact g values using Origin (OriginLab, Northampton, MA). The simulated spectra were processed with SimFonia (Bruker BioSpin, Billerica, USA) and WINEPR (Bruker BioSpin, Billerica, USA).

**X-ray absorption spectra**
We conducted the measurements of the Fe K-edge X-ray absorption spectroscopy at the wiggler beamline (BL17C) with a beam size of $2 \times 2$ mm at the National Synchrotron Radiation Research Center in Taiwan. Beam energy was calibrated to iron foil standards. The sample cell at a volume of 120 $\mu$L was sealed with thin Fe-free kapton tapes. We performed sample loading in the anaerobic chamber and conducted data collection with the sleeping mode that halted the exposure by approximately 1-8 s after 2-24 s radiation. The Lytle detector collected signals ranging from 6912 to 7912 eV with florescence mode. Samples were maintained at 283 K throughout the experiments with an air-cooling device. Each scan ran for approximately 90 min. Based on collected spectra, the anaerobic samples were stable for 20 h indicated that the effects of the photo-reduction or radiation damage were negligible for the duration of the experiment. The data analysis and background subtraction were performed using the ATHENA program (24), a graphical interface in the IFEFFIT (version 1.2.12) (25). We used the average of 30 scans for the model fitting. Following baseline correction (with $R_{bkg} = 1$ using AUTOBK), the EXAFS data were analyzed and simulated using ARTEMIS to yield the fitting curves (24). Scattering paths of the [4Fe-4S] cluster were generated with ATOMS (26) and FEFF (25) programs, in which the initial distance of Fe-Fe was set as 2.7 Å and that of Fe-S as 2.2 Å, respectively, based on measurements from the synthetic (27) and experimental Fe-S clusters (28). Two-shell models centered on iron atoms surrounded by: (1) iron and sulfur atoms (such as [4Fe-4S]-(S-Cys) × 4 and [3Fe-4S]-(S-Cys) × 3); or (2) iron, sulfur, and oxygen ([4Fe-4S]-(O-Ser)×1-(S-Cys)×3) were used in data analysis and only atoms at distances within $1.8 \leq R \leq 3.3$ Å were considered.

RESULTS

NMR evidence supports a paramagnetic center in KpFeoC. Using relaxation-optimized sequences (see Methods), we observed a proton hyperfine shift of KpFeoC isolated directly from the cell culture (native state) at -14 ppm, indicating that KpFeoC possesses a paramagnetic center (Fig. 1A)
Upon addition of dithionite (reduced state), the resonance shifted upfield by 5 ppm to -19 ppm, indicating that the paramagnetic property of the paramagnetic center is redox-state-dependent. Hyperfine shifted $^{13}$C resonances were also observed from the native state [$U^{13}$C, $^{15}$N] -KpFeoC (Fig. 1B). Although the assignments of the hyperfine-shifted proton and carbon resonances are currently unknown, the detection of these resonances supports the presence of a paramagnetic Fe-S cluster in KpFeoC. Upon raising the temperature from 293 K to 300 K, the proton resonance of the reduced state KpFeoC shifted upfield by 0.5 ppm, characteristic of the anti-Curie (increasing paramagnetic shifts with increasing temperature) hyperfine-shifted resonance (data not shown). However, the $^{13}$C resonances exhibited both Curie (resonance 4) and anti-Curie (resonance 3) behaviors (Fig. 1B). In comparison, the [1Fe] system of rubredoxin exhibited only the Curie behavior, suggesting that holo-KpFeoC may possess a higher-order Fe-S cluster (30-32), consistent with the X-ray absorption results (see the following).

Spectrophotometric evidence suggests a [2Fe-2S] or [4Fe-4S] cluster in KpFeoC

UV-VIS spectra were diagnostic of the presence of iron sulfur clusters (33). Figure 2 shows the UV-VIS spectra of GST-tagged KpFeoC (GST-KpFeoC) and 4 cysteine-to-serine mutants (C56S, C61S, C64S, and C71S, Fig. 2A), and their spectra without the GST-tag (Fig. 2B). For comparison, we included the spectrum of Clostridium pasteurianum rubredoxin (CpRd), which contains a [1Fe] cluster resembling that of another winged helix protein, PF0610 (34). The color of the (concentrated) native state GST-KpFeoC was dark-red and the UV-VIS spectrum contained peaks at 417 nm, 450 nm, and 550 nm. Upon enzymatic removal of the GST tag, the wild-type KpFeoC still maintained a nearly identical UV-VIS spectrum, indicating that the absorption spectrum derived from KpFeoC (Fig. 2B). However, the peak intensity dropped 2-fold, likely because of cluster degradation during the prolonged enzyme digestion process (3 d). However, the UV-VIS spectrum of CpRd contained peaks at 384 nm, 490 nm, and 570 nm (Fig. 2C), suggesting that GST-KpFeoC likely contained a [2Fe-2S] (34, 35) or [4Fe-4S] (36-38) cluster, but not a [1Fe] cluster.
We further determined the fractional concentration of \( KpFeoC \) containing a \([4Fe-4S]\) cluster by inductively coupled plasma-mass spectrometry (ICP-MS) and spectrophotometry. For spectrophotometric measurements, we used the canonical extinction coefficient of 15,000 at 410 nm per \([4Fe-4S]\) cluster (39). The results from both methods showed that 10% of native \( KpFeoC \) contain the \([4Fe-4S]\) cluster.

**EPR evidence supports a \([4Fe-4S]\) cluster on \( KpFeoC \)**

Iron sulfur clusters exhibit the EPR spectra characteristic of the type of clusters (40–42). Thus, we employed EPR to assign the cluster type present in \( KpFeoC \). At high temperature (77 K), we detected no signal at approximately \( g = 2 \) from native or reduced states of \( KpFeoC \) (data not shown). Upon lowering the temperature to 14 K, we observed 2 signals at \( g = 2.060 \) and 2.007 from the native \( KpFeoC \), characteristic of the \([4Fe-4S]\)^{3+} state (Fig. 3A). Reducing \( KpFeoC \) by using DTT significantly reduced the resonance intensity, and we observed only residual resonances at \( g = 2.05 \) and 2.008 (Fig. 3B), indicating that most of the protein had been reduced to the EPR-silent diamagnetic \([4Fe-4S]\)^{2+} state. After further reducing \( KpFeoC \) with sodium dithionite, a stronger reducing agent, the EPR signals reappeared at \( g = 2.038 \) and 1.937, reminiscent of those of the \([4Fe-4S]\)^{1+} cluster (Fig. 3C) (36,37). Thus, the redox potential of \([4Fe-4S]\)^{2+/1+} lies between that of DTT and dithionite (43, 44). We also observed EPR signals at \( g = 4.3 \) for both native and DTT-reduced states (data not shown). We attribute this signal to free iron or non-specifically bound iron on the protein (45). Supplemental Table S1 presents a summary of the EPR signal observed for \( KpFeoC \) under various conditions.

**Fe K-edge X-ray absorption structures support the \([4Fe-4S]\) cluster**

We further investigated *holo*-\( KpFeoC \) using Fe K-edge X-ray absorption spectra. We conducted our initial attempt under air and the Fe K-edge X-ray absorption near-edge structure (XANES) features of the rising-edge appeared at approximately 7125.9 eV (Fig. 4A) suggesting oxygen degradation, which was confirmed by extended X-ray absorption fine structure (EXAFS, data not shown). To
protect the cluster from oxygen degradation, we repeated the experiment with samples prepared in
anaerobic conditions (Fig. 4A). Consequently, the near-edge down-shifted to 7118.7 eV and the pre-
edge (corresponding to 1s to 3d transitions) peak down-shifted to 7112.7 eV, suggesting that anoxic
samples have fewer positive charges to the irons because of a reduced oxidation state (46). We then
collected the EXAFS to verify the cluster and ligands (Fig. 4B). We chose the EXAFS of k ranging
from 3.12–12.00 Å\textsuperscript{-1} for analysis (Fig. 4B inset) and applied the Fourier transform to yield the
distances of the ligating atoms (iron and sulfur, Fig. 4B). We selected three models for simulation:

\begin{itemize}
\item [4Fe-4S]-{(S-Cys)} \times 4,
\item [3Fe-4S]-{(S-Cys)} \times 3 (degradation intermediate), and
\item [4Fe-4S]-{(O-Ser)} \times 1-{(S-Cys)} \times 3 (serine replacement).
\end{itemize}

Among them, [4Fe-4S]-{(S-Cys)} \times 4 was the most optimal model, which yielded the lowest R\textsubscript{f} value of 0.07 %. According to the model, the average Fe-S distances were
reported as 2.26 ± 0.05 Å and the Fe-Fe distance was 2.71 ± 0.09 Å, consistent with canonical [4Fe-
4S] clusters (27, 28). The EXAFS results were in good agreement with the EPR results, and a higher
coordinated number of Fe-Fe bonds excluded the [2Fe-2S], supporting the temperature-dependence
EPR results. Therefore, both the EPR and EXAFS results suggest that the [4Fe-4S] is the native cluster
in FeoC.

**The [4Fe-4S] cluster of KpFeoC is sensitive to oxygen**

Similar to other iron-sulfur proteins, the holo-KpFeoC was oxygen sensitive (40). When exposed to
oxygen at 4 °C, native KpFeoC gradually lost its characteristic absorption peaks (Fig. 5). The change in
peak height was fitted to a single exponential decay function of OD = c + A*0.5\textsuperscript{\tau/2}, and the results
yielded a half-life of \tau_{1/2} = 16.0 ± 1.0, 15.8 ± 1.0, and 18.3 ± 1.3 h for absorbance at 417 nm, 450 nm,
and 550 nm, respectively, with an average half-life of 17 h. We monitored the protein using SDS-
PAGE analysis, which indicated that the proteins were non-degraded (data not shown). Our data
suggested that the [4Fe-4S] cluster in KpFeoC was oxygen labile. Proteins purified from a size-
exclusion column in air for 2 h at 4°C were depleted of the Fe-S cluster, suggesting that the degradation
rate is likely less than 1 h under the chromatography conditions. The limited oxygen availability in the 1 mm × 10 mm × 10 mm cuvette without stirring might slow the degradation rate measured by the spectrophotometric experiments.

To identify the degraded products, we initially treated the freshly prepared native \( Kp \text{FeoC} \) sample with 1 mM DTT anaerobically and then exposed it to air at 4 °C. We used a series of EPR spectra at 14 K at various exposure times (supplemental Fig. S1). Because DTT-reduced \( Kp \text{FeoC} \) was diamagnetic, the initial spectrum showed only a signal from the residual non-reduced protein. Further exposure to oxygen resulted in the appearance of a peak at \( g = 2.010 \), which gradually increased in intensity and peaked at 20 h. Further exposure to oxygen resulted in a gradual loss of the EPR signal. The time course of the oxidation process is consistent with the initial buildup of the \([3\text{Fe}-4\text{S}]^{1+}\) cluster, the only iron-sulfur cluster with an isotropic \( g \) value at 2.01. The end product of the oxidation process is the loss of the iron-sulfur cluster and the generation of \( \text{apo-FeoC} \), thus the loss of the EPR signal.

To assess the final state of the oxidation product, we added DTT or dithionite to reduce the cluster after 25 h of exposure to oxygen. DTT did not produce any change in the EPR spectrum (Fig. S1D), whereas dithionite greatly reduced the EPR signal (Fig. S1E). These results suggested that 25-h \( \text{O}_2 \)-exposed \( Kp \text{FeoC} \) did not contain a sufficient concentration of \([2\text{Fe}-2\text{S}]^{2+}\) (EPR silent) because dithionite should produce the \([2\text{Fe}-2\text{S}]^{1+}\) (EPR active). The absence of a \( g = 1.96 \) was indicative of the depleted \([4\text{Fe}-4\text{S}]^{1+}\) cluster (the product of the dithionite reduction of the \([4\text{Fe}-4\text{S}] \) cluster) in the final oxidation product, indicating that \([4\text{Fe}-4\text{S}] \) was completely degraded. We concluded that \([3\text{Fe}-4\text{S}] \) is likely the intermediate oxidative degradation process of the \([4\text{Fe}-4\text{S}] \) cluster in \( Kp \text{FeoC} \).

**Cysteines in the W1 loop are the ligands for the \([4\text{Fe}-4\text{S}] \) cluster**

\( Kp \text{FeoC} \) contains 4 conserved cysteine residues, which are all located in the W1 loop. To assess the roles of these cysteines in cluster formation, we generated 4 single-site Cys to Ser mutants (C56S, C61S, C64S, and C71S) and analyzed them with EPR (Fig. 6). Optical spectra (Fig. 2) of all GST-
tagged mutants were similar to those of the wild-type *KpFeoC*. However, upon enzymatic removal of
the GST-tag, only the C61S mutant maintained the characteristic absorption spectrum of the [4Fe-4S]
cluster (Fig. 2) and EPR signals similar to those of the wild-type *KpFeoC* (Fig. 6B, right panel). The
intensity of [4Fe-4S]$^{1+}$ of C61S was approximately 30% that of the wild type at a similar concentration,
indicating a less stable cluster. These results indicated that Cys56, Cys64, and Cys71 are crucial for the
formation and stability of the iron-sulfur cluster, whereas Cys61 also plays a role in stabilizing the
cluster, but is less essential. We suggest that Cys61 is the fourth ligand with the support from EXAFS,
but it is possible to substitute Cys61 with other nearby glutamic acid or serine residues in the C61S
mutant. In certain cases, serine can also serve as a ligand for the iron-sulfur cluster (35).

**DISCUSSION**

*KpFeoC* forms an oxygen sensitive [4Fe-4S] cluster

The Fe-S proteins contain inorganic iron and sulfur as cofactors (40,47,48). Iron and sulfur are redox
active and the cluster can undergo redox reactions in physiological potential ranges. Studies have
documented several types of biological Fe-S clusters, including the simplest [1Fe] cluster in
rubredoxin, to the complex [8Fe-7S] cluster in nitrogenase (42,49). A change in cluster oxidation states
is associated with characteristic magnetic properties, thus EPR spectra are effective fingerprints of the
cluster type and redox state of the Fe-S cluster (40,41). For the [4Fe-4S] cluster at near-liquid helium
temperature, the [4Fe-4S]$^{3+}$ and [4Fe-4S]$^{1+}$ states are paramagnetic and thus can be distinguished from
the diamagnetic [4Fe-4S]$^{2+}$ state (41,50). The 2 paramagnetic states can be distinguished from the g
factors in the EPR spectra, which can be detected only at near-liquid helium temperature because of
fast relaxation at ambient temperature (50). However, in NMR spectroscopy, the paramagnetic effect
from [2Fe-2S]$^{2+}$ and [4Fe-4S]$^{2+}$ clusters causes a hyperfine shift in the NMR resonances, which can be
detected (32,51-53). A summary of the EPR results in the literature, combined with our results of the
*holo-KpFeoC*, is presented in the supplemental Table S1 for comparison.
Based on UV-VIS spectrophotometry, NMR, and EPR evidence, we demonstrated that *KpFeoC* forms an Fe-S cluster. The characteristic UV-VIS spectrum and NMR spectra suggested that the iron-sulfur cluster is not the simple [1Fe] type. The presence of the EPR signal at \( g = 1.937 \) in the dithionite reduced state ruled out the possibility of the [3Fe-4S]\(^{1+/-0}\) cluster because the reduced state of [3Fe-4S]\(^{0+}\) is diamagnetic (41, 50). As suggested by Cammack et al., [2Fe-2S] and [4Fe-4S] clusters can be distinguished by temperature-dependent EPR intensities (32,50,54). The EPR signals of [2Fe-2S] are observable above 77 K, whereas the EPR signal of the [4Fe-4S]\(^{1+}\) cluster is observable only at a temperature below 30 K. This was the case in *KpFeoC* (resonances disappeared above 37 K, data not shown), confirming the identity of the [4Fe-4S] cluster in *KpFeoC*. The EXAFS results further support the conclusion of [4Fe-4S], excluding the possibility of [2Fe-2S], and confirm that the [4Fe-4S] is ligated by 4 cysteines. Protein exposure to oxygen generated the [3Fe-4S] intermediate, further supporting the [4Fe-4S] identity in native *KpFeoC*. In summary, UV-VIS, NMR, EPR, and EXAFS provide unequivocal evidence of the presence of the [4Fe-4S] cluster on *KpFeoC*.

**Redox states and oxygen-induced degradation of holo-FeoC**

Cysteine typically coordinates each tetrahedral Fe site in the form of thiolate (RS\(^-\)). However, other residues, such as aspartate (RCO\(\cdot\)), histidine (N\(=\)), and serine (R-O\(\cdot\)) are occasionally encountered in clusters and these ligands were shown to modify redox potential (55). The protein environment also affects the redox potential of an iron-sulfur cluster. We showed that the [4Fe-4S] cluster on *KpFeoC* is coordinated to the cysteine residues in the W1 loop. We estimated the redox potential of the [4Fe-4S] in the 3 oxidation states by examining the effect of DTT or dithionite on the native *KpFeoC* sample. We detected that most *KpFeoC* protein in the [4Fe-4S]\(^{3+}\) state reduced to the [4Fe-4S]\(^{2+}\) state by DTT and further reduced to the [4Fe-4S]\(^{1+}\) state by dithionite. Because the redox potential of DTT is -0.33 V and that of dithionite is -0.66 V (43,44), the results indicated that the redox potential of the [4Fe-4S]\(^{3+/-2+}\) of FeoC is higher than -0.33 V and that of [4Fe-4S]\(^{2+/1+}\) is between -0.33 V and -0.66 V. However, we did not determine the precise redox potential of holo-*KpFeoC*. 

13
The loss and gain of the iron-sulfur cluster is a common sensing mechanism for the Fe-S proteins to exhibit their biological activity. The cluster assembling machinery assembles the iron-sulfur cluster, and degradation by oxidative agents removes the cluster. The ICP-MS results showed that 20% of the freshly prepared GST-KpFeoC contained the Fe-S cluster, indicating that the Fe-S cluster assembling machinery is capable of assembling at least 20% of the over-expressed GST-KpFeoC. We examined the degradation of holo-KpFeoC by exposing the protein to oxygen and detected the presence of a [3Fe-4S]^{2+} intermediate. To assess whether the [2Fe-2S] state is the final cluster degradation product of holo-KpFeoC, we added dithionite to the final product to reduce the diamagnetic [2Fe-2S]^{2+} state, if present, to the paramagnetic [2Fe-2S]^{1+} state. We did not detect the EPR signal near g = 2.0, suggesting that the [2Fe-2S]^{2+} state is not the final product. The pathway is similar to that of FNR, but the rates differ, suggesting various sensory/regulatory mechanisms (15,56-59).

**Biological implications**

Iron-sulfur proteins play key roles in catalytic reactions, in electron transfer in both oxidative phosphorylation and photosynthesis, and in gene regulation (41,42,48,60,61). The roles of bacterial iron-sulfur regulatory proteins as sensors/switches have been extensively reviewed (42,61-63). Iron-sulfur clusters sense environmental changes by interacting with small molecules to exhibit rich chemistries and regulate cellular events. Iron-sulfur proteins acting as transcriptional regulators, such as the *E. coli* fumarate-nitrate reduction regulator protein (EcFNR), can alter binding affinities to specific DNA sequences by various cluster states. EcFNR is activated only when the O2-labile [4Fe-4S] cluster is assembled; holo-FNR recognizes specific binding sites in excess of 100 promoters (15,64,65). The iron-sulfur cluster regulator (IscR) exhibits different DNA-binding properties in its apo- and holo states, controlling different subsets of gene expressions (66-70). The cytoplasmic aconitase also regulates gene expression through differential binding affinities of the apo- and holo protein to the iron regulatory elements within the mRNA of genes related to iron metabolism (71,72). The NreB contains a [4Fe-4S]^{2+} cluster, but it does not bind to nucleotides, acting as a transcriptional activator by
interacting with the response regulator, NreC, to regulate the expression of the nreABC operon (73,74). Thus, the iron-sulfur proteins regulate transcription through direct binding to the DNA or RNA. They also exhibit transcriptional activity indirectly by affecting the activity of other proteins that interact with DNA activators or repressors.

In the literature, 3 models have been proposed for FeoC function: the transcriptional regulator model (6), the G-protein modulator model (21), and the protease inhibitor model (22). Available data does not currently support the transcriptional regulator model and we could not detect KpFeoC binding to DNA using the gel-shift or SELEX experiments (20,22); evidence supporting the G-protein modulator model is lacking. The protease inhibitor model appeared to be the only model supported by in vivo data. However, it is likely too early to discount any of these models and it is conceivable that FeoC may possess dual functions, or function differently in various systems. The presence of the Fe-S cluster on FeoC can have a substantial effect on its function, regardless of its role as a transcriptional regulator, a G-protein modulator, or a protease inhibitor. Our present work confirming the existence of the Fe-S cluster on FeoC should facilitate future studies in defining the roles and function of the Fe-S cluster. Future works are necessary to confirm their biological role. The low yield and oxygen sensitivity of holo-KpFeoC hampers current studies focused on clarifying the role of the Fe-S cluster. Advanced understanding requires the development of methods for generating high-yield holo-KpFeoC. However, the oxygen sensitivity of [Fe-S] cluster on FeoC in vitro does not necessarily indicate that the cluster functions as an oxygen sensor in vivo.

In summary, by using spectrophotometric, NMR, EPR, and X-ray absorption methods, we showed that KpFeoC contains a [4Fe-4S] cluster that can be degraded by oxygen. Using single-site mutation and EXAFS techniques, we identified the crucial cysteine residues in the W1 loop as the ligands of the Fe-S cluster. Detection of the oxygen-sensitive Fe-S cluster in FeoC raises the possibility that the Fe-S cluster might play a role in regulating Feo activity.
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REFERENCES


FIGURE CAPTIONS:

**Figure 1.** Hyperfine shifted NMR resonance of 1 mM unlabeled-\(KpFeoC\) detected by superWEFT pulse sequence. (A) \(^1\)H spectra of native state (~ mostly oxidized, top trace) and the dithionite-reduced state (bottom trace). (B) \(^{13}\)C spectra of 1.3 mM \([U-^{13}\text{C},^{15}\text{N}]\)-\(KpFeoC\) at native state at 283K (top trace) and 178K (bottom trace). The recycle delay times for the superWEFT sequence were ~0.1 and ~0.2 seconds for proton and carbon spectra, respectively.

**Figure 2.** The UV-VIS spectra of various freshly prepared \(KpFeoC\) samples (0.5-3 mM) in native state. (A) The UV-VIS spectrum of GST-tagged \(KpFeoC\) (GST-\(KpFeoC\)) and four cysteine to serine single site \(KpFeoC\) mutants at concentrations of 2.5, 1.5, 2.6, 2.6 and 0.9 mM for C56S, C61S, C64S, C71S and wild type GST-FeoC, respectively. (B) The UV-VIS spectra of GST-tag free \(KpFeoC\) and its single site mutants at concentrations of 0.5, 0.9, 0.8, 0.5 and 0.15 mM for C56S, C61S, C64S, C71S and wild type \(FeoC\), respectively. The 300 nm – 700 nm region, which exhibits the characteristic absorption of iron-sulfur clusters is shown on the insert, shifted on the Y-axis direction for clarity. For clarity the Y-axis (C) Comparison of the UV-VIS spectrum of 3 mM \(KpFeoC\) (red) and 1.5 mM \(CpRd\) (black) which possesses the [1Fe].

**Figure 3.** EPR spectra (first derivatives) of \(KpFeoC\) at different redox states at 14 K. (A) 2.1 mM \(KpFeoC\) at the native state. (B) 2.7 mM \(KpFeoC\) at DTT reduced state. (C) 2.7 mM \(KpFeoC\) fully reduced by sodium dithionite. A SimFonia simulated spectrum was overlaid (thin solid line). The assigned redox states of the [4Fe-4S] clusters were labeled on the right. EPR properties were summarized in Table S1.

**Figure 4.** X-ray absorption spectra. (A) The normalized XANES of \(holo\)-FeoC at the Fe K-edge. Both experiments done in anaerobic and aerobic conditions were overlaid. The pre-edge region was enlarged with absorption peaks labeled with black arrow at 7113 eV and red arrow at 7115 eV. (B) The EXAFS data of \(holo\)-FeoC. The both experimental data (open circles) and simulated spectra (solid lines)
according to model [4Fe-4S]-(S-Cys) × 4 were overlaid. The $k^3$-weighted Fourier transform over $k$ range of 3.12–12 Å$^{-1}$, respectively. The raw data of EXAFS ($k^3\chi$) used for the analysis is shown in the inset.

**Figure 5.** Kinetics of the O$_2$-induced degradation of the [4Fe-4S] cluster in *Kp*FeoC (0.1 mM). The optical absorbance at 417 nm (filled square), 450 nm (empty square) and 550 nm (filled circle) were monitored at various time points of O$_2$ exposure. 100 μL of sample was kept at 4°C in 50 mM Tris, 100 mM NaCl, pH 7.8 without stirring. Fittings of the curves (solid lines) yielded half-lives of 16.0 ± 1.0, 15.8 ± 1.0, and 18.3 ± 1.3 (hrs) as monitored at 417 nm, 450 nm, and 550 nm, respectively.

**Figure 6.** EPR spectra (first derivatives) of four *Kp*FeoC mutant proteins at 14 K at native (left panel) and dithionite reduced state (right panel). (A) C56S, 1.5 mM; (B) C61S, 2.9 mM; (C) C64S, 2.4 mM; (D) C71S, 1.1 mM.