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Krisztina Z. Bencze Wayne State University

Kalyan C. Kondapalli Wayne State University

Jeremy D. Cook Wayne State University

Stephen McMahon Wayne State University

César Millán-Pacheco Universidad Autonoma del Estado de Morelos

 $See\ next\ page\ for\ additional\ authors$ 

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authors Krisztina Z. Bencze, Kalyan C. Kondapalli, Jeremy D. Cook, Stephen McMahon, César Millán-Pacheco Pastor, and Timothy L. Stemmler	o, Nina

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# The Structure and Function of Frataxin

Krisztina Z. Bencze\*, Kalyan C. Kondapalli\*, Jeremy D. Cook\*, Stephen McMahon\*, César Millán-Pacheco<sup>&</sup>, Nina Pastor<sup>&</sup> and Timothy L. Stemmler\*<sup>‡</sup>

\*Department of Biochemistry and Molecular Biology, Wayne State University, School of Medicine, Detroit, Michigan, USA; &Facultad de Ciencias, Universidad Autonoma del Estado de Morelos, Cuernavaca, Morelos, Mexico

<sup>‡</sup> Address correspondence to Timothy L. Stemmler, Department of Biochemistry and Molecular Biology, Wayne State University, School of Medicine, 540 E. Canfield Ave, Detroit, MI 48201, USA. E-mail: <a href="mailto:tstemmle@med.wayne.edu">tstemmle@med.wayne.edu</a>

# **Abstract**

Frataxin, a highly conserved protein found in prokaryotes and eukaryotes, is required for efficient regulation of cellular iron homeostasis. Humans with a frataxin deficiency have the cardio- and neurodegenerative disorder Friedreich's ataxia, commonly resulting from a GAA trinucleotide repeat expansion in the frataxin gene. While frataxin's specific function remains a point of controversy, a general consensus is the protein assists in controlling cellular iron homeostasis by directly binding iron. This review focuses on the structural and biochemical aspects of iron binding by the frataxin orthologs and outlines molecular attributes that may help explain the protein's role in different cellular pathways.

# Keywords

Friedreich's ataxia, heme biosynthesis, iron-sulfur cluster assembly, iron chaperone, Yfh1, CyaY, ferrochelatase, ISU, Aconitase Biological context

Frataxin, a conserved protein found ubiquitously in prokaryotes and eukaryotes, is required for the cellular regulation of iron homeostasis. Although of extreme interest to a large number of chemists and biologists alike, frataxin's exact role in helping cells regulate iron chemistry and metal availability remains controversial at best. Frataxin has been proposed to participate in at least 5 different capacities: as an iron chaperone during cellular heme and iron-sulfur (Fe-S) cluster production, as an iron-storage protein during conditions of iron overload, assisting in the repair of oxidatively damaged aconitase Fe-S clusters, as a factor that controls cellular oxidative stress by moderating the concentration of reactive oxygen species (ROS) and finally as an active participant in pathways involving energy conversion and oxidative phosphorylation. While these functions are not necessarily exclusive, it seems extraordinary that one protein could directly control so many biological pathways within cells. Disruption of cellular frataxin production does cause a general loss in the cellular control of iron bioavailability and reactivity, however phenotypes are complicated which further inhibits defining the proteins exact function. The goal of this manuscript is to take a structural approach to help understand frataxin's role during the regulation of cellular iron homeostasis. However, before we begin to draw structural correlations regarding frataxin's function, it is beneficial to first look in detail at the iron reaction chemistry the protein helps control.

Iron is extremely versatile in the chemistry it can perform, so it is no surprise that the metal is found ubiquitously in all living systems. Iron is found in the active sites of proteins involved in many diverse cellular functions ranging from oxygen transport, oxidative metabolism and electron transport to energy production. Iron's ability to cycle between the ferrous (Fe(II)) and ferric (Fe(III)) oxidation states allows the metal to be extremely useful when performing redox chemistry and "this chemical versatility is surely a reason why nature selected this element in so many life processes" (Cotton, et al., 1988). Unfortunately iron's flexibility towards performing redox chemistry means the metal, if left unregulated, can perform deleterious reactions that elevate ROS concentrations, and these reactive oxygen species can damage cells by oxidatively attacking cellular membranes, proteins and DNA alike. Ferrous iron will readily react with dioxygen to produce superoxide and ultimately hydroxyl radicals, as described by the Fenton reaction (Aisen, et al., 2001; Kosman, 2003):

$$Fe(II) + O_2 \rightarrow Fe(III) + O_2$$
 (Eq.1)

$$2 O_2^{-} + 2H^+ \rightarrow H_2O_2 + O_2$$
 (Eq. 2)

$$Fe(II) + H_2O_2 \rightarrow OH' + OH' + Fe(III)$$
 (Eq. 3)

It is therefore beneficial to control the oxidation of ferrous iron as a means to prevent the damaging effects of hydroxyl radical production. Under normal physiological conditions (pH > 2), ferric iron undergoes stepwise hydrolysis leading to the formation of insoluble neutral species:

$$[Fe(H_2O)_6]^{3+} \rightarrow [Fe(H_2O)_5(OH^-)]^{2+} + H_3O^+$$
 (Eq. 4)

$$[Fe(H_2O)_5(OH^-)]^{2+} \rightarrow [Fe(H_2O)_4(OH^-)_2]^+ + H_3O^+$$
 (Eq. 5)

$$[Fe(H_2O)_4(OH^-)_2]^+ \rightarrow [Fe(H_2O)_3(OH^-)_3] + H_3O^+$$
 (Eq. 6)

Organisms have therefore evolved protein-controlled mechanisms to modulate iron's redox chemistry. These mechanisms allow cells to quickly adapt to enhancement or suppression of iron import, the delivery of Fe for utilization and finally the storage of this important element for future use. In humans, iron abstracted from diet is reduced and acidified by the low pH environment of the stomach and a portion is eventually transported for use in the production of heme and Fe-S clusters. A small percentage of the metal is also stored in a manner that can be easily mobilized. More than two thirds of the iron found in the body is incorporated into hemoglobin in developing erythroid precursor cells or in mature red blood cells (Andrews, 1999), highlighting the importance of heme Fe-prosthetic groups as components that help control local O<sub>2</sub> concentration, availability and reactivity. Owing to their remarkable structural plasticity and versatile chemical/electronic features, Fe-S clusters are additional prosthetic groups that help control cell and organism viability by participating in electron transfer events, substrate binding and activation, iron and sulfur storage, the regulation of gene expression and in some cases enzyme activity (Johnson, et al., 2005). The balance of iron in biological systems is however fragile, as both iron deficiency and overload are destructive to cells. Diseases related to the disruption in cellular iron homeostasis are, as a result, among the most common diseases found in humans.

Friedreich's ataxia (FRDA), an autosomal recessive cardio- and neurodegenerative disorder that affects 1 in 50,000 humans, results from an inability to produce frataxin and protein deficiency leads to a disruption of cellular iron homeostasis (Delatycki, et al., 2000). The majority of FRDA patients (ca. 96%) have extensive trinucleotide repeat expansions in the first intron of the gene encoding frataxin, while a small percentage of patients have frataxin point mutations (Campuzano, et al., 1996). The expanded GAA repeat adopts abnormal DNA structures that impair frataxin transcription; the longer the repeat, the more profound the frataxin deficiency, the earlier the onset of the disorder and the greater the intensity of the disease (Bidichandani, et al., 1998; Ohshima, et al., 1998; Pandolfo, 2002). In humans, frataxin is a 210 amino acid protein that is nuclear encoded but has an N-terminal mitochondrial targeting sequence that gets removed during processing

(Campuzano, et al., 1996; Gibson, et al., 1996). Results from immunocytofluorescence and immunocytoelectron microscopy show processed frataxin is predominately located within the mitochondria and associated with mitochondrial membranes, crests and as free soluble protein within the matrix (Gibson, et al., 1996). Frataxin mRNA is predominantly expressed in tissues with a high metabolic rate, including the liver, kidney, brown fat neurons and heart (Koutnikova, et al., 1997). Phenotypes coupled with a frataxin deficiency include mitochondrial iron accumulation (Babcock, et al., 1997; Foury, et al., 1997), disruption in both heme and Fe-S cluster production (Rotig, et al., 1997; Stehling, et al., 2004) and a progressive breakdown in cellular iron homeostasis (Babcock, et al., 1997; Koutnikova, et al., 1997). Mitochondrial iron accumulation diverts metal away from other cellular compartments and frataxin deficient cells accommodate for what they perceive as a general cellular iron deficiency by elevating mitochondrial iron import (Babcock, et al., 1997). Boosting pathways that lead to increased mitochondrial iron import facilitates the problems associated with not being able to process the reactive iron that is already present, leading to a further elevation in ROS levels, gradual cell damage and finally the loss of cell viability (Delatycki, et al., 2000). FRDA patients therefore show a slow progression of muscular and neurological symptoms (loss of motor function, progressive limb and gait ataxia, etc.) linked to the disruption in proper iron regulation and the disorder is typically fatal due to complications resulting from cardiomyopathy (Orth, et al., 2001).

A direct correlation between frataxin and cellular iron homeostasis is at present obvious, but early studies on the yeast frataxin homolog (Yfh1) provided the initial insight into what role(s) frataxin may play in helping cells maintain iron homeostasis (Babcock, et al., 1997). Deletion of the frataxin gene results in the accumulation of mitochondrial iron deposits (Babcock, et al., 1997; Foury, et al., 1997) coupled with both aconitase and general Fe-S cluster protein deficiencies (Foury, 1999; Rotig, et al., 1997). The presence of zinc suppresses iron accumulation phenotypes in yeast (Knight, et al., 1998). AYfh1 cells were shown to be hypersensitive to H<sub>2</sub>O<sub>2</sub>, iron and copper levels (Babcock, et al., 1997; Foury, et al., 1997). Based on the observation that reintroduction of Yfh1 expression under depleted conditions can promote the recovery and export of these mitochondrial iron deposits, frataxin was originally proposed to directly control mitochondrial iron efflux (Radisky, et al., 1999). Yfh1 was also shown to indirectly regulate mitochondrial iron uptake by interacting and partially controlling the activity of the yeast mitochondrial intermediate peptidase (YMIP), a metalloprotease required for maturation of ferrochelatase and other iron utilizing proteins (Branda, Yang, et al., 1999).

In eukaryotes, frataxin is proteolytically processed during the production of the mature soluble protein found within the mitochondrial matrix. During translation, a frataxin precursor is produced which contains a N-terminal targeting sequence that directs the protein to the mitochondria and eventually into the matrix (Babcock, et al., 1997; Knight, et al., 1998). Processing of the yeast frataxin precursor protein takes place uniquely in two steps (Branda, Cavadini, et al., 1999; Gordon, Wang, et al., 2001). Typically, proteins targeted to the mitochondria are processed in a single step (Branda, Cavadini, et al., 1999; Cavadini, Adamec, et al., 2000; Gordon, Kogan, et al., 2001). There are only a few additional proteins that show this variation and examples include the Rieske (Fe-S) containing proteins and Neurospora crassa ATPase subunit 9 (Branda, Cavadini, et al., 1999; Cavadini, Adamec, et al., 2000; Gordon, Kogan, et al., 2001). Even within this category, only frataxin and ATPase subunit 9 use a single enzyme for processing during both steps (Cavadini, Adamec, et al., 2000). In the first step, the mitochondrial processing peptidase removes the Nterminal 20 residues (domain I) of the protein generating a Yfh1 intermediate. This intermediate is then processed again by the peptidase to remove the next 31 amino acids (domain II), generating the 123 amino acid mature Yfh1 found within the matrix. The two processing steps are independent. Domain I is a typical mitochondrial targeting signal and it can be substituted by other mitochondrial targeting signals with similar cellular results. Domain II serves as a spacer. The first 20 amino acids of frataxin contain the 5 basic residues typically found in mitochondrial targeting signals. The N-terminal region of the mature protein is rich in acidic residues. A spacer separates basic residues in the targeting signal from acidic residues in the mature protein's Nterminus. This prevents unwanted interactions that might hinder recognition of the target signal by the import proteins. Both domains are required for import into the mitochondria, however protein with mutations blocking the first or second processing steps will still complement the Yfh1 deletion phenotype (Gordon, Kogan, et al., 2001).

N-terminal processing of human frataxin (HsFtx) is still a matter of debate. Cavadini *et al.* proposed a two-step processing of the N-terminus, whereas Gordon *et al.* could identify only a single processing event (Cavadini, Adamec, *et al.*, 2000; Gordon, Kogan, *et al.*, 2001). In the two-step processing proposal, the first cleavage occurs between residues 41 and 42, removing the first 41 residues of HsFtx to generate an intermediate. The second cleavage step occurs between residues 55-56, removing only the next 14 N-terminal residues. The kinetics of the first processing step was shown to be similar between the different species, however the second cleavage reaction rate is species specific. The second processing step (i.e., removing the 14 residues) is the rate-limiting step during HsFtx processing (Cavadini, Adamec, *et al.*, 2000). In contrast, Gordon *et al.* detected only a

single processing event generating the 18-kDa mature human frataxin and similar results were observed with rat or yeast mitochondrial peptidase (Gordon, Kogan, *et al.*, 2001).

# STRUCTURAL INSIGHTS INTO FRATAXIN'S FUNCTION

A detailed understanding of frataxin's structure provides some insight into how the protein may function within its different proposed roles. Specific questions relevant to the molecular structure of frataxin include: Is there a patch of residues that could accommodate iron binding if frataxin is acting as a metal chaperone? Is there a favorable interface that could promote contacts between frataxin and protein partners coupled with heme biosynthesis, Fe-S cluster assembly and aconitase Fe-S cluster repair? Are there molecular elements on frataxin's surface that might promote protein aggregation during an iron storage function? and finally, Does frataxin support an active site that could assist in the stabilization of the redox chemistry performed by the bound iron. We will attempt to address each question by looking closely at the structures of the different frataxin orthologs.

Solution and crystal structures have been reported for the yeast, human and bacterial (CyaY) frataxin orthologs (Cho, et al., 2000; Dhe-Paganon, et al., 2000; He, et al., 2004; Lee, et al., 2000; Musco, et al., 2000; Nair, et al., 2004). Frataxin has a unique fold that combines two terminal α-helices to make one plane, five antiparallel β-strands that construct the second plane of the protein and a sixth (and, in HsFtx, a seventh) β-strand that intersects the planes to give an overall planer  $\alpha$ - $\beta$  sandwich structural motif structure (Figure 1). The general shared frataxin topology is α1β1β2β3β4β5β6(β7)α2, strand seven is again only detected in HsFtx. In all three cases, the structures are extremely similar; RMSD's of HsFtx versus CyaY and Yfh1 in regions of secondary structure are 1.34 Å and 0.65 Å, respectively. The overall structural dimensions of the orthologs are:  $47 \times 32 \times 28 \text{ Å}^3$  for HsFtx,  $45 \times 30 \times 25 \text{ Å}^3$  for CyaY and  $47 \times 29 \times 23 \text{ Å}^3$  for Yfh1. Mature HsFtx spans amino acids 75-210 of the total expressed sequence, however auto-degradation and proteolysis at the protein's N-terminus prior to all residues have complicated structural studies of the fulllength mature protein. Therefore, the structures solved of HsFtx begin at residue 88 in the crystal structure and 90 in the solution structure (Dhe-Paganon, et al., 2000; Musco, et al., 2000). Circular dichroism data on full-length mature HsFtx indicate the N-terminal residues prior to residue 92 are predominately unstructured. Mature Yfh1 spans residues 52-174 of the expressed gene sequence. Unlike HsFtx, the 18 amino acids N-terminal to α1 in Yfh1 are stable and partially structured in a 3<sub>10</sub> helix observed between residues 65 and 68. Mature CyaY lacks any appreciable residues Nterminal to α1. In the case of HsFtx only, frataxin has a 17 amino acid C-terminal tail following α2.

This C-terminal tail of HsFtx adopts a random coil structure that is tethered to the protein's helical plane exposed surface.

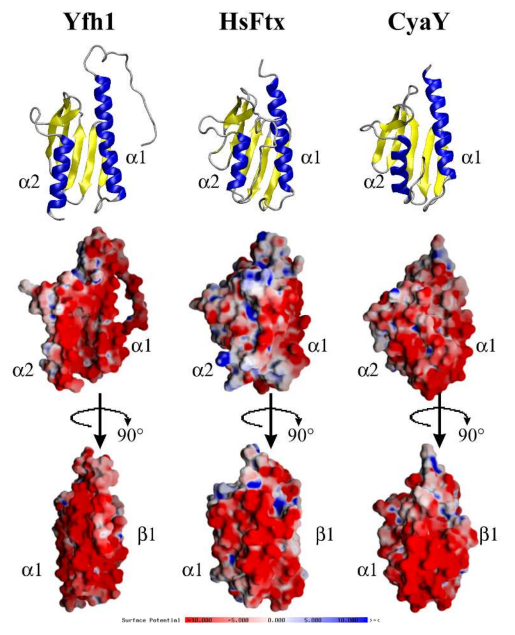


Figure 1. Top: ribbon diagram for yeast, human and bacterial frataxin. Middle: electropotential plots for proteins in same orientation. Bottom: electropotential plots for proteins rotated '90 degrees around the y-axis compared to top display. Structure figures made using solution structures of Yfh1 (PDB ID# 2GA5), HsFtx (PDB ID# 1LY7) and CyaY (PDB ID# 1SOY) frataxins.

The strong structural similarity between frataxin orthologs results from the fact that these proteins share an extremely high degree of amino acid sequence similarity. Sequence identity of Yfh1 versus both CyaY and HsFtx are 28.1% and 37.8%, respectively, while respective sequence

similarities are 59.8% and 65.0%. A large number of the highly conserved amino acids include a subset of Asp and Glu residues located in the N-terminal region of the protein. For Yfh1 this amounts to 9 of the first *ca.* 50 N-terminal residues in the mature protein constructed by conserved Asp and Glu residues (Figure 2).

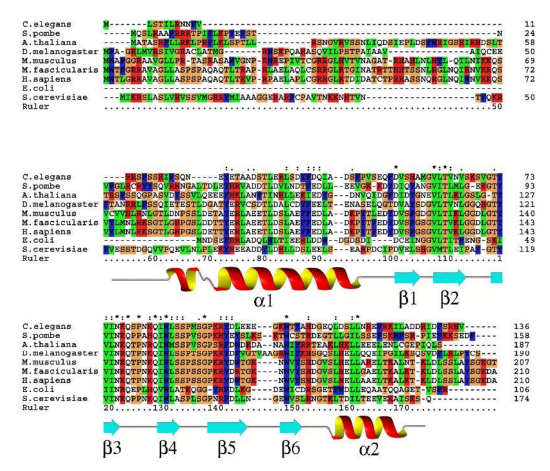


Figure 2. ClustalX alignment for a subset of characterized frataxin orthologs. Bottom three sequences represent structurally characterized frataxin orthologs. Secondary structural elements and ruler representing Yfh1 properties are given below the sequences.

Conserved N-terminal acidic residues are located in the  $\alpha 1$  and  $\beta 1$  secondary structural regions of frataxin. Electrostatic potential plots show these conserved acidic residues line the exposed surface of the  $\alpha 1$  and  $\beta 1$  interfaces, generating a general negatively charged surface that covers roughly a quarter of frataxin's total accessible surface (Dhe-Paganon, *et al.*, 2000). Carboxylate side chains from Asp and Glu residues often serve as ligands for bounds metal in many iron-binding proteins, suggesting the possibility of these regions participating in metal binding. While HsFtx does not contain any cysteines, CyaY and Yfh1 contain poorly conserved cysteines at positions distant from the acidic patches.

Multiple factors have been shown to influence the stability of frataxin's fold. The Pastore laboratory (MRC, London) performed extensive characterization studies on the three in vitro isolated and structurally characterized frataxins with the goal of defining factors that help stabilize each protein's structure (Adinolfi, Nair, et al., 2004; Adinolfi, et al., 2002). Apo-bacterial, yeast and human frataxin are predominately stable as monomers at 1 mM protein concentrations, as confirmed by proton linewidths observed in <sup>15</sup>N labeled protein <sup>15</sup>N-HSQC specta (He, et al., 2004; Mori, 1995; Musco, et al., 1999; Musco, et al., 2000; Nair, et al., 2003; Nair, et al., 2004). Apo-Yfh1 is however stable as a monomer for only up to two weeks after isolation, at which point it begins to self aggregate (Cook, et al., 2006). Although CyaY, Yfh1 and HsFtx share a high degree of sequence homology and fold, the bacterial and human orthologs have melting points in the respective range of 54 and 60°C (Adinolfi, Nair, et al., 2004), while the yeast protein is less stable, with a melting point at ca. 39°C ((Adinolfi, Nair, et al., 2004) and unpublished results from our laboratory). The type of buffer (organic vs. phosphate) has minimal effect on protein stability for these three frataxin orthologs, however the presence of salt generally increases the stability of frataxin's fold (most pronounced in Yfh1). Lowering the pH raised the stability of Yfh1 but this had no effect on the helical fold of CyaY and HsFtx within the pH range of 6-8. The presence of iron at high metal to protein stoichiometries further increased the stability of all frataxins orthologs. Residues Cterminal to α2 in the protein's structure appear to dramatically increase the general stability of frataxin. C-terminal CyaY truncation mutants matching the shorter Yfh1 construct showed a substantial (14°C) drop in the protein's melting temperature. A C-terminal extension to Yfh1 caused a ca. 7°C increase in the protein's melting temperature. Structural studies show the Cterminal extension in CyaY and HsFtx are unstructured but make surface contacts with  $\alpha 1$  and  $\alpha 2$ amino acids, most likely providing additional contacts that help stabilize the protein's fold. Finally, the non-conserved N-terminal region of full-length HsFtx appears to be important for the partial apo-protein aggregation often found in the *in vitro* expressed/isolated protein (O'Neill, Gakh, et al., 2005).

Close inspection of frataxin's helical plane shows interesting structural attributes that may contribute to a better understanding of how the protein binds metal and also in mutations that cause FRDA. On Yfh1,  $\alpha$ 1 is lined with exposed surface acidic residues Glu71, Asp78, Asp79, Asp82, Asp86, Glu89, Glu90 and Glu93 (Figure 3A). The orientation and proximity of these conserved residues generates a conserved acidic patch on the protein in this region and these residues would be in a perfect position to contribute to cation binding. Frataxin helices  $\alpha$ 1 and  $\alpha$ 2 are parallel and orientated at a average adjacent backbone distance of ca. 11 Å, predominately due

to interactions between the long side chains of hydrophobic and aromatic residues within this plane (in Yfh1,  $\alpha$ 1 residues in this region include Leu84, Leu88, Leu91 and  $\alpha$ 2 residues include Leu162, Val166 and Ile170). Additional interactions between hydrophobic core amino acids on the  $\beta$ -sheet surface of the protein help secure the spatial orientation of  $\alpha$ 1 to  $\alpha$ 2.

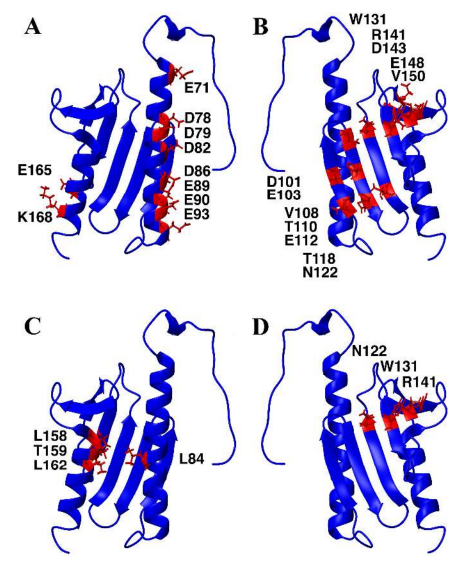


Figure 3. Yfh1 residues that are highly conserved on the helical (A) and β-sheet (B) planes of the protein. Identity of HsFtx FRDA point mutations on the helical (C) and β-sheet (D) planes of Yfh1. Structure figures made using Yfh1 solution structure (PDB ID# 2GA5).

In case of the human frataxin, the C-terminal tail of HsFtx, including residues Thr196, Leu198 and Leu200, appear to further contribute to the stabilization of these inter-helical residue interactions (Cho, et al., 2000; Dhe-Paganon, et al., 2000; Lee, et al., 2000; Musco, et al., 2000; Nair, et al., 2004). Surface exposed residues on the α2 surface of frataxin are less conserved; in Yfh1,

Glu165 and Lys168 are fairly well conserved however the remaining surface residues are mixed. Yfh1 surface residues in the β6-loop (or in HsFtx β6-β7) region share highly conserved polar Arg153 and Asn154 residues. Disease causing HsFtx helical plane point mutations are identified on the conserved Yfh1 sequence as α1 residue Leu84 mutation (to a Ser) and on α2, Leu158 mutation (to a His), Thr159 mutation (to an Arg) and Leu162 mutation (to an Arg) (Figure 3C) (Bartolo, et al., 1998; Cossee, et al., 1999; Filla, 1996; Zuhlke, et al., 2004). Since these are all buried in the globular domain of the protein and play an important role in the packing of the two helices, mutating these residues most likely causes a disruption of elements that stabilize frataxin's fold.

Numerous surface exposed residues on the β-sheet plane of frataxin are also highly conserved, and close inspection of this surface aids in the understanding of mutations that cause FRDA. Surface exposed residues on the Yfh1 β-sheet plane include Asp101, Glu103, Val108, Thr110, Glu112, Thr118, Asn122, Trp131, Arg141, Asp143, Glu148 and Val150, of which Asp101, Thr110, Asn122, Trp131 and Asp143 are fully conserved (Figure 3B). HsFtx amino acids, complementary in position to Yfh1 residues Asn122, Trp131 and Arg141, are surface exposed and mutated in a small set of FRDA patients (Figure 3D) (Forrest, et al., 1998; Labuda, et al., 1999; Zuhlke, et al., 2004). Yfh1 β-sheet residues with side chains directed towards the hydrophobic core of frataxin include the conserved residues Ile121, Leu132 and Trp149. Along the β-sheet surface of HsFtx, Yfh1 residues at complementary positions of Ile130, Leu132 and Trp149 were found as point mutations in some FRDA patients, and mutating these residues would surely alter frataxin's fold (Campuzano, et al., 1996; Cossee, et al., 1999). In addition, Gly107 in the loop region between Yfh1's β1 and β2 is an additional point mutation found in a subset of FRDA patients (Bidichandani, et al., 1997). Yfh1 residue Ile99 is the final HsFtx point mutation seen in a subset of FRDA patients, and this residue is located on the  $\beta$ -sheet plane of frataxin, connecting  $\alpha 1$  to  $\beta 1$  (Cossee, et *al.*, 1999).

Although frataxin's fold is fairly unique, a  $\alpha$ - $\beta$  sandwich motif structure is seen in a surprising number of proteins that share similar functions to those proposed for frataxin. The  $\alpha$ - $\beta$  sandwich motif can be easily recognized in the ferredoxin-like fold of some cytosolic copper chaperones. Although human HAH1 (Anastassopoulou, *et al.*, 2004), yeast ATX1 (Arnesano, *et al.*, 2001) and BsCopZ (Banci, *et al.*, 2003) (PDB ID's: 1TL5, 1FES and 1P8G) are smaller in their overall size compared to frataxin (ca. 18Å x 22Å x 30Å), and their helices are tilted at 45 degrees to each other, the helices and beta strands still form a general two layer structure that is similar to the  $\alpha$ - $\beta$  sandwich motif found in frataxin. The copper binding domains of these proteins involve a conserved CXXC motif formed in a loop region; this binding site is surrounded by hydrophobic

residues that are essential for binding to partner proteins but distinct in position from the acidic patch on frataxin's  $\alpha 1/\beta 1$  surface (i.e., the proposed metal binding region of frataxin). Recent crystallographic data on complex I of the *Thermus thermophilus* mitochondrial respiratory chain revealed a previously unknown protein Ngo15 that is also a member of this superfamily (Sazanov, et al., 2006). Ngo15 binds in close contact with 4 other domains to help stabilize the larger complex. This protein is constructed by 129 amino acids, held together in an  $\alpha-\beta$  sandwich motif fold. Despite a low sequence homology (average of 12%), the crystal structures superimpose well with RMSD values of 2.5 Å and 3.3 Å for CyaY and HsFtx, respectively. This domain binds to the complex through its β-sheet surface in a manner that creates a hydrophilic channel connecting the solvent with the N3 subdomain. The N-terminal  $\beta1$  and loop connecting  $\beta1$  and  $\beta2$  contain several surface exposed histidine residues. Based on the fold similarity between frataxin and Nqo15, and given that a recent publication from the Cowan laboratory (Ohio State University) that proposed frataxin's β-sheet plane may be an essential interface when interacting with ISU proteins during Fe-S cluster assembly (Yoon, et al., 2003), the authors of the Ngo15 structure proposed that the conserved β-sheet plane histidine residues could serve as an iron binding site during iron storage and delivery by the protein when reconstitution of the Fe-S cluster in subdomain N3 is required.

# FRATAXIN'S IRON BINDING ABILITY

Given the spatial arrangement of conserved acidic residues on the α1/β1 regions of frataxin (Figure 1), and the fact that Asp and Glu carboxylate side chains are chemically suited as ligands for binding metals, it is not surprising to learn that frataxin is an iron binding protein. However, the Isaya laboratory (Mayo Clinic) were the first to identify that frataxin directly binds iron; this iron binding ability was first determined in the yeast system (Adamec, et al., 2000). The iron binding ability of additional frataxin orthologs has since been confirmed for both the human and bacterial proteins (Adinolfi, et al., 2002; Bou-Abdallah, et al., 2004; Cavadini, et al., 2002; Cook, et al., 2006; Yoon, et al., 2003; Yoon, et al., 2004). While it was initially recognized that iron binding in the yeast system induces frataxin to oligomerize under unique solution conditions, numerous subsequent reports suggest that oligomerization is not essential when the protein functions in an iron chaperone capacity (i.e., during heme and Fe-S cluster assembly) but it may be important when the protein participates in helping control ROS production under iron overloading conditions. In an attempt to deconvolute the published reports concerning the iron binding ability of frataxin in its different oligomeric states and how these relate to the different proposed functions of frataxin, we will selectively discuss, in the following text, issues concerning: first the iron binding ability of

monomeric frataxin, secondly frataxin's ability to form iron loaded oligomers and finally the protein's ability to regulate the redox chemistry performed by iron bound to the protein. Each of these points is summarized below.

# Frataxin will bind iron as a stable protein monomer.

Metal to protein stoichiometries, measured for monomeric bacterial, yeast and N-terminally truncated human frataxin show these proteins will tightly bind 2, 2 and 7 Fe(II) atoms, respectively (Bou-Abdallah, et al., 2004; Cook, et al., 2006; Yoon, et al., 2003). Iron dissociation constants, measured for CyaY, Yfh1 and N-terminally truncated HsFtx give average K<sub>d</sub> values of 3.8, 3.0 and 55.0 μM, respectively(Bou-Abdallah, et al., 2004; Cook, et al., 2006; Yoon, et al., 2003). The presence of magnesium or calcium salts, at physiologically relevant concentrations found within the mitochondria, stabilize CyaY and Yfh1 in the iron bound monomeric state against oligomerization, while a large portion of bacterially overexpressed HsFtx is stable as an iron loaded monomer regardless of the solution conditions (Adinolfi, et al., 2002; Cook, et al., 2006). N-terminally truncated HsFtx was shown to bind ca. 6 Fe(III) atoms at a K<sub>d</sub> of 10.2 μM (Yoon, et al., 2003). In the presence of oxidants, monomeric CyaY can bind up to six Fe(III) atoms and will interact with up to 25 Fe(III) atoms/monomer (Bou-Abdallah, et al., 2004). Ferric iron binding studies for Yfh1 are linked only to protein homooligomerization (Adamec, et al., 2000).

NMR spectroscopy provided a powerful tool for identifying frataxin amino acids directly affected by the presence of iron. Backbone amide resonances for the three characterized frataxins are generally well dispersed, hence providing a fingerprint for the majority of protein amino acids (He, et al., 2004; Musco, et al., 1999; Nair, et al., 2003). Subtle perturbations in the amide resonances and general signal intensity due to the addition of iron therefore serve as markers for residues affected by the presence of paramagnetic iron. Ferrous iron titrations performed anaerobically with the <sup>15</sup>N labeled frataxin orthologs show distinct alterations in α1 and β1 amino acid NMR resonances and, in the case of the CyaY and HsFtx, these perturbations are dependent on metal to protein stoichiometry. General trends in amide resonance perturbations upon addition of iron can be divided into two categories: amide resonances that were significantly line broadened (often beyond detection) and amide chemical shifts that were shifted in the presence of metal. Bacterial amide resonances broadened upon addition of a single Fe(II) atom include α1 residues Arg20, Leu21, Asp22 and Asp23 (Table 1), while human resonances include α1 residues Asp112, Leu113, Asp115 and Val125 (Nair, et al., 2004).

Table 1. Comparison of the iron interaction with Yfh1, HsFtx and CyaY residues in  $\alpha 1$  thru

**B2** secondary structural regions

β2 secondary structural regions									
<u>Yfh1</u>		<u>HsFtx</u>		<u>CyaY</u>		<b>Structure</b>			
Sequence	Fe Effect	Sequence	Fe Effect	Sequence	Fe Effect	Consensus			
Glu71		Thr93		Ser4	br2				
Lys72		Thr94		Glu5	br2				
Tyr73		Tyr95		Phe6	br2	α1			
His74		Glu96		His7	br2	α1			
Glu75	sh2	Arg97		Arg8		α1			
Glu76	sh2	Leu98		Leu9		α1			
Ala77		Ala99		Ala10		α1			
Asp78	sh2	Glu100		Asp11		α1			
Asp79	br2	Glu101		Gln12		α1			
Tyr80		Thr102		Leu13		α1			
Leu81		Leu103		Trp14		α1			
Asp82	br2	Asp104	br2	Leu15		α1			
His83	br2	Ser105		Thr16		$\alpha 1$			
Leu84		Leu106		Ile17		α1			
Leu85		Ala107	br2	Glu18		α1			
Asp86	br2	Glu108	br2	Glu19	br2	α1			
Ser87	sh2	Phe109		Arg20	br1	$\alpha 1$			
Leu88		Phe110	br2	Leu21	br1	α1			
Glu89	sh2	Glu111		Asp22	br1	α1			
Glu90	sh2	Asp112	br1	Asp23	br1	α1			
Leu91		Leu113	br1	Trp24	br2				
Ser92	sh2	Ala114	br2	Asp25					
Glu93	sh2	Asp115	br1	Gly26					
Ala94		Lys116							
		Pro117							
His95		Tyr118							
Pro96		Thr119	br2						
Asp97		Phe120	br2	Asp27					
Cys98		Glu121		Ser28					
Ile99		Asp122	br2	Asp29					
Pro100		Tyr123		Ile30					
Asp101		Asp124	br2	Asp31	br2	β1			
Val102		Val125	br1	Cys32	br2	β1			
Glu103	br2	Ser126		Glu33		β1			
Leu104		Phe127		Ile34		β1			
Ser105	sh2	Gly128		Asn35		β1			
His106		Ser129		Gly36					
Gly107		Gly130		Gly37					
Val108		Val131		Val38		β2			
Met109		Leu132		Leu39		β2			
Thr110	1.0	Thr133		Thr40		β2			
Leu111	sh2	Val134		Ile41		β2			
Glu112	br2	Lys135		Thr42		β2			
Ile1113		Leu136		Phe43		β2			

**Table 1.** The table represents a stoichiometry of between 1 to 2 Fe(II) per protein monomer. Abbreviations include: br1 and br2 (broadened and shifted at 1:1 and 2:1 respective Fe(II) to protein ratios) and sh2 (shifted at a 2:1 Fe(II) to protein). The final column contains the ortholog consensus secondary structure. The dashed line represents gaps in sequence homology (see Figure 2 for clarification). Conserved residues are shaded.

Additional bacterial resonances broadened upon the addition of a second Fe(II) atom include α1 residues Ser4, Glu5, Phe6, His7, Glu19 and Trp24, and β1 residues Asp31 and Cys32; human resonances include those from α1 residues Asp104, Ala107, Glu108, Phe110, Ala114, Thr119, Phe120, Asp122 and Asp124. Addition of up to six Fe(II) atoms to CyaY and HsFtx caused a general broadening of amide resonances throughout the α1 and β1 regions. A number of additional CyaY and HsFtx resonances in the α1 and β1 regions underwent chemical shift perturbations upon the addition of ferrous metal. Yfh1 resonances broadened under similar buffer conditions by addition of up to 2 iron atoms include α1 residues Asp79, Asp82, His83 and Asp86, the β1 residue Glu103 and the β2 residue Glu112, while amide resonances for additional α1, β1 and β2 residues undergo chemical shift perturbations in the presence of iron (Cook, et al., 2006). Ferric iron titrations were performed on CyaY with similar results to those obtained from ferrous iron binding. CyaY residues significantly broadened at a 1:1 Fe(III) to protein stoichiometry include α1 residues Arg20, Asp22 and Asp23. In the presence of a 2:1 ration, additional broadened resonances include α1 residue Asp29, as well as β1 residues Ile30, Asp31, Cys32, Glu33 and Ile34. Again, addition of up to six iron atoms causes a general broadening of  $\alpha 1$  and  $\beta 1$  resonances. Finally binding studies performed on CyaY with the non-paramagnetic Ca(II) ions showed general chemical shift perturbations in only the α1 and β1 regions (Nair, et al., 2004). In summary, these NMR titrations paint a picture of frataxin's metal binding residues located predominately in the conserved acidic α1 and β1 regions of frataxin (Table 1).

Structural studies directed at characterizing the metal-ligand coordination geometry and electronic properties of bound iron provide additional insight into the metal binding ability of the monomeric frataxin orthologs. X-ray absorption spectroscopy (XAS) provides a powerful tool for characterizing structural and electronic properties of metals bound to proteins in solution (Teo, 1986). This technique has been used to characterize iron bound to Yfh1 and HsFtx monomers. Iron bound to monomeric Yfh1 and HsFtx is stable in the ferrous state when samples are prepared anaerobically or in the presence of the reducing agent dithionite (Bencze, et al., 2006; Cook, et al., 2006). Analysis of the iron 1s $\rightarrow$ 3d transitions in the x-ray absorption near edge structure (XANES) portion of the XAS spectrum are consistent with ferrous iron existing in the high-spin state and

coordinated in a highly centrosymmetric metal-ligand coordination geometry when bound to monomeric HsFtx and Yfh1 (Bencze, et al., 2006; Cook, et al., 2006). Structural analysis of the Fe(II)-ligand coordination geometry for monomeric Yfh1 and HsFtx from the extended x-ray absorption fine structure (EXAFS) portion of the XAS spectrum are consistent with iron bound in a highly symmetric 6 coordinate ligand environment. Ligands coordinating Fe(II) are exclusively oxygen and/or nitrogen based, in agreement with α1 and β1 Asp, Glu and His residues identified in the multiple NMR titrations as residues that possibly interact with iron. Additional ligands to the protein bound metal surely also come from water or hydroxide ions. There is no evidence for any metal • • • metal interaction for the ferrous iron bound to monomeric frataxin, even when up to two metals are bound in both Yfh1 and HsFtx.

# Frataxin can form stable iron loaded homooligomers

Frataxin ability to homooligomerize, forming aggregates with metal binding abilities similar to that of ferritin, suggests frataxin may act in an iron storage protein. Under elevated metal to protein stoichiometries, the presence of oxygen and the absence of salt, Yfh1 will form a 48-multimeric homooligomer that can bind up to 50 iron atoms per protein monomer (Gakh, et al., 2002). CyaY aggregates of a similar size and iron loading capacity will also form under low salt, aerobic and iron-overloaded solution conditions (Adinolfi, et al., 2002). Assembly in both systems is predominately averted under high concentrations of other divalent metal ions or in the presence of physiological salt concentrations (Adamec, et al., 2000; Adinolfi, et al., 2002; Cook, et al., 2006), indicating CyaY and Yfh1 assembly is iron specific and will proceed under low salt but molecular oxygen (O<sub>2</sub>) dependent conditions. Monomeric HsFtx does not self-assemble at high iron concentrations, however a ca. 59-mer HsFtx homooligomer is isolated as part of the E. coli overexpressed protein and this oligomer can bind approximately 10 iron atoms under aerobic and salt free conditions (Cavadini, et al., 2002). The bulk iron bound in Yfh1 and HsFtx assemblies is predominately in the high-spin Fe(III) state (Nichol, et al., 2003).

Yfh1 homooligomers have a unique morphology that highly resemble those seen from the iron storage protein ferritin, suggesting iron storage may be an important function of frataxin (Adamec, et al., 2000; Gakh, et al., 2002). Unlike ferritin, which forms spherical aggregates in the absence of metal (Theil, 1973), Yfh1 will only self assemble in the presence of Fe, O<sub>2</sub> and low salt (Adinolfi, et al., 2002; Park, et al., 2002). Stoichiometric and higher Fe(II):protein ratios drive Yfh1 to progressively self-assemble into larger homooligomers beginning from a protein trimer repeating unit, directing next to a hexamer, then to a 12-mer, 24-mer and finally resulting in the 48-mer

spherical protein aggregate that resembles ferritin (Adamec, et al., 2000). The 48-mer Yfh1 aggregate stores bulk metal, of greater than 2000 iron atoms, predominately as ferrihydrite (Nichol, et al., 2003). The ca. 59-mer HsFtx homooligomers, isolated from a portion of the protein overexpressed in E. coli, will store ca. 600 iron atoms as ferrihydrite in a rod-shaped polymeric protein structural morphology (Cavadini, et al., 2002). Non-covalent protein subunit interactions, mediated by the non-conserved N-terminal region of human frataxin, were shown to be important for assembly (O'Neill, Gakh, et al., 2005). Iron does not induce the assembly of the monomeric portion of overexpressed HsFtx isolated from E. coli (Adinolfi, et al., 2002). CyaY can also form aggregates in the presence of iron and like Yfh1, aggregation is closely controlled by the presence of salt (Adinolfi, et al., 2002). Analytical ultracentrifugation studies show CyaY tetramers can be formed when Fe(II) is added anaerobically; different protein aggregates are formed upon oxidation of the bound Fe(II) (Bou-Abdallah, et al., 2004). These in vitro assembly results are intriguing as they suggest protein aggregation may be an important property of the protein, in that it may help regulate cellular iron homeostasis under iron overloaded cellular conditions.

Residues in the α1/β1 region are essential for controlling frataxin homooligomerization. Specific conserved acidic residues in the α1/β1 region of Yfh1 and CyaY are required for assembly. Mutating CyaY α1 residues Glu18, Glu19 and Asp22, along with β1 residue Glu33 abolishes the iron induced oligomerization behavior (Adinolfi, et al., 2002). Mutating conserved Yfh1 acidic residues Asp86, Glu90 and Asp93 abolishes the iron induced oligomerization behavior of the protein, surprisingly with no real *in vivo* phenotypes under normal growth conditions (Aloria, et al., 2004). Additional Yfh1 mutational studies indicate Asp93 is critical for protein self-assembly (Gakh, et al., 2006). Unlike CyaY and Yfh1, a portion of the full length mature HsFtx (residues 56-210) will self-assemble in the absence of metal; N-terminally truncated HsFtx lacking the initial 22 residues does not self assemble in the absence of iron (Cavadini, et al., 2002; O'Neill, Gakh, et al., 2005).

XAS results show iron bound to Yfh1 and HsFtx spherical aggregates is predominately stable in the ferric state when samples are prepared aerobically at high metal to protein ratios (Nichol, *et al.*, 2003). Analysis of the 1s→3d transitions in the XANES spectra indicate ferric iron is coordinated in a highly centrosymmetric metal-ligand structural environment. Structural analysis of ferric iron coordinated to Yfh1 and HsFtx aggregates are consistent with bulk metal being bound as ferrihydrite, a biomineral composed of ferric oxide/hydroxide octahedra. The ligands coordinating the metal come predominately from complexed mono- and bidentate oxygen ligands that hold the multinuclear Fe cluster together and not from protein based ligands. These structural studies were essential for the early identification of how frataxin oligomers store bulk metal, hence participating

as an iron storage capacity, maintaining metal in an inert form to shield the cell against unwanted iron redox chemistry.

Functional correlations between frataxin oligomers and ferritin are obvious. Mitochondrial ferritin (MtF) is an additional iron storage protein found in eukaryotes. Mature MtF is a 22kDa protein encoded by an intronless gene on 5q23.1 chromosome in humans. MtF shares a 79% sequence identity with H-ferritin and the proteins possess ferroxidase activity. Its expression is restricted to tissues with high number of mitochondria (i.e., the testis) unlike frataxin, which is ubiquitous in its expression. MtF expression is not correlated to tissues like liver, which are involved in iron storage. MtF expression is observed in iron-loaded erythroblasts from patients with sideroblastic anemia, but not in normal erythroblasts, suggesting it may be induced only under conditions of stress in some cells. Consistent with these data, reports show that MtF expression in normal cells results in cytoplasmic and mitochondrial iron deprivation and a decrease in enzymatic activity of Fe-S cluster containing enzymes like aconitase (Levi, et al., 2004). Correlating these findings with what is known about iron storage role of frataxin, it is possible that frataxin may be the iron storage molecule under low salt but normal iron level conditions whereas both frataxin and mitochondrial ferritin scavenge iron to protect the cell from oxidative damage under iron overloading conditions.

# Frataxin controls iron's redox chemistry

As previously outlined, iron is extremely reactive towards oxygen based redox chemistry, so it is not surprising that a breakdown in cellular iron homeostasis resulting from a frataxin deficiency causes elevated oxidative stress to affected cells. Controlling the ability of iron to perform redox chemistry appears to be an additional function of frataxin, suggesting this protein may directly participate in controlling cellular oxidative stress by reducing ROS production. *In vivo* reports indicate frataxin deficiency leads to oxidative damage in humans (Emond, *et al.*, 2000; Schultz, *et al.*, 2000), mice (Ristow, *et al.*, 2003; Thierbach, *et al.*, 2005), yeast (Karthikeyan, *et al.*, 2003) and *Caenorhabditis elegans* (Vazquez-Manrique, *et al.*, 2006). Mitochondrial iron overload, resulting from a frataxin deficiency, leads to oxidative damage in mitochondrial and nuclear DNA as well as to Fe-S clusters in mitochondrial aconitase and other respiratory enzymes (Babcock, *et al.*, 1997; Cavadini, Gellera, *et al.*, 2000; Foury, 1999; Karthikeyan, *et al.*, 2002). Interestingly, frataxin affords protection for DNA against iron-induced oxidative damage (Gakh, *et al.*, 2006; O'Neill, Gakh, *et al.*, 2005), presumably by directly binding iron and modulating its oxidation chemistry (Bou-Abdallah, *et al.*, 2004).

In numerous biological systems, protein controlled ferroxidase centers direct the controlled oxidation of ferrous iron. These centers are common among proteins within different metabolic pathways that involve iron. One of many examples of ferroxidase centers is found in the iron storage protein ferritin (Theil, 1987; Theil, 2003). Mammals contain two types of ferritin subunits, the heavy (H) chain and the light (L) chain, which differ in size. Ferritin binds ferrous iron *via* Glu and His residues on the H chain and metal is ultimately oxidized at the protein's ferroxidase center to form oxygen bridged diferric center. Metal is then moved from the ferroxidase center to the nucleation site for storage as inert ferric oxide cores.

Recent reports have shown frataxin will also perform ferroxidase chemistry (Park, et al., 2002). In Yfh1 at sub-stoichiometric metal to protein levels, coupled with ferrous iron oxidation, H<sub>2</sub>O<sub>2</sub> is most likely generated through O<sub>2</sub> consumption during the ferroxidase reaction. The presence of hydrogen peroxide is only minimally detected under these conditions and therefore it has been proposed that H<sub>2</sub>O<sub>2</sub> immediately reacts with protein to attenuate all Fenton chemistry. Ferroxidation is progressively overcome by slower autooxidation at stoichiometric and higher iron to protein concentrations in Yfh1, and metal oxidation leads to progressive assembly of higher order protein oligomers. Therefore, it has been suggested that frataxin's ability to initially perform ferroxidase chemistry, retaining a portion of bound metal in a bioavailable form, and eventually storing iron in a less accessible form allows this protein to serve both as an iron chaperone and an iron storage protein (Park, et al., 2003). Formation of the ferroxidase center in yeast frataxin is correlated to all residues Asp79 and Asp82; yeast mutants targeting these residues that have reduced ferroxidase activity and assemble at a slower rate than wild type protein (Gakh, et al., 2006). A series of frataxin  $\alpha 1/\beta 1$  residues were implicated as participating in the iron mineralization chemistry observed under iron overloaded conditions, possibly by acting as iron binding amino acids in the yeast frataxin oligomers. Yfh1 residues α1 Asp86, Glu89, Glu90 and Glu93, and \$1 residues Asp101 and Glu103 are generally important in promoting the iron binding and metal sequestering properties of frataxin oligomers (Gakh, et al., 2006).

Ferroxidase centers were also detected in HsFtx and CyaY aggregates (O'Neill, Gakh, *et al.*, 2005). The oligomeric form of iron loaded HsFtx showed distinct protection of DNA against oxidative damage in the presence of H<sub>2</sub>O<sub>2</sub> (O'Neill, Gakh, *et al.*, 2005). Iron oxidation studies on CyaY monomers showed no apparent ferroxidase activity in the presence of molecular oxygen; rather, O<sub>2</sub> acted as a rather poor oxidant to Fe(II) bound to the protein (Bou-Abdallah, *et al.*, 2004). The oxidation of Fe(II) bound to CyaY was accelerated in the presence of H<sub>2</sub>O<sub>2</sub> and as a result there was an increase in the protein's ability to bind additional metals (Bou-Abdallah, *et al.*, 2004).

Unpublished results from our laboratory for Yfh1 monomers showed similar results to those published for CyaY monomers, with  $O_2$  induced Fe(II) oxidation rates on the minutes time scale (50% conversion in ca. 10 minutes at 100  $\mu$ m protein/iron concentrations). In these published studies, during the reduction of  $O_2$  to  $2H_2O$ , the presence of highly reactive  $H_2O_2$  or hydroxyl radicals was minimal.

# FRATAXIN'S ROLE IN CELLULAR HEME BIOSYNTHESIS

Cells often utilize heme prosthetic groups when helping control normal cellular pathways. Given that frataxin is required for the production of heme, it is not a far stretch to suggest this protein could directly participate in heme bioassembly. Heme, a ubiquitous iron-containing tetrapyrrole ring system, is involved in multiple aspects of cellular metabolism. The Fe-prosthetic groups are used by globins, cytochromes and several additional enzymes (Ponka, 1999) that play key roles in the sensing and/or utilization of molecular oxygen in all living organisms (Andrew, et al., 1990; Padmanaban, et al., 1989; Zhu, et al., 1999). Heme centers help drive cellular energy generation during respiration by storing and transporting oxygen (Atamna, 2004). Heme groups can also participate in both catalytic and regulatory functions within cells (Zhu, et al., 2002). This prosthetic group can function as an effector center that regulates several biological processes encompassing transcription, translation, protein translocation and erythroid differentiation (Padmanaban, et al., 1989). They function as ligands for transcription factors within prokaryotes (Monson, et al., 1992) and in yeast (Creusot, et al., 1988; Fytlovich, et al., 1993; Pfeifer, et al., 1989; Zhang, et al., 1998). In higher eukaryotes, heme helps control the activity of specific transcription factors (Ogawa, et al., 2001; Sassa, et al., 1996) and proteins in several biochemical pathways by binding to the short protein sequence labeled the "heme regulatory motif" (Zhang, et al., 1995). Heme regulation is essential during erythroid differentiation. In mammalian erythroid cells, heme initiates changes in key factors controlling numerous activities ranging from cell cycle and Ras signaling to chromatin structure, splicing and protein folding (Zhu, et al., 1999). Thus the multifunctional roles of heme suggests fluctuations in its concentration help control several key aspects of cellular metabolism (Atamna, 2004). In order to better understand the role frataxin plays in heme bioassembly, we present a brief description of the assembly pathway as it applies to our protein.

The heme biosynthetic pathway was outlined in detail during the 1950's and 60's (Labbe, et al., 1999). Cellular heme biosynthesis occurs in eight sequential steps, four of which (step 1 and steps 6-8) occur within the mitochondria, while the other steps occur within the cytoplasm (Tait, 1978). The enzyme ferrochelatase catalyses the final step in the heme biosynthetic pathway.

Ferrous iron is inserted into porphyrin by ferrochelatase to produce a functional heme prosthetic group (Taketani, 2005). Ferrochelatase is associated with the inner mitochondrial membrane in eukaryotes; in prokaryotes, ferrochelatase is found within the cytoplasm (Bacillus subtillus) or associated to the cytoplasmic membrane (Dailey, et al., 2000). To this date, structures of apo- and metal-loaded bacterial (Bacillus subtilis), yeast (Saccharomyces cerevisiae) and human ferrochelatases have been solved (Al-Karadaghi, et al., 1997; Karlberg, et al., 2002; Wu, et al., 2001). While the functional bacterial protein is monomeric, human and yeast ferrochelatase are homodimers (Karlberg, et al., 2002). Mammalian, yeast (Schizosaccharomyces pombe) and some bacterial ferrochelatases are metalloenzymes that contain [2Fe-2S] prosthetic groups which are required for activity (Ferreira, 1999). All three orthologs have a conserved overall fold of two Rossmann-type domains with a four-stranded parallel  $\beta$ -sheet that is flanked by  $\alpha$ -helices. These domains contribute to the formation of the porphyrin-binding cleft. The differences between the eukaryotic and prokaryotic structures are mostly limited to the region in the eukaryotic ferrochelatase suggested to be involved in interactions with membranes and to the C-terminus, which contains the Fe-S cluster in the human but not the S. cerevisiae and B. subtilis enzymes (Karlberg, et al., 2002). The localization of ferrochelatase to the matrix side of the inner mitochondrial membrane assures uptake of the poorly soluble porphyrin and for heme release (Ferreira, 1999; Taketani, 2005). The hydrophobic exterior of ferrochelatase is in stark contrast to the hydrophilic interior of the active sites that are lined with conserved charged residues well positioned to receive the positively charged iron. The porphyrin-binding cleft of ferrochelatase exhibits a high degree of conservation from prokaryotes to higher eukaryotes (Al-Karadaghi, et al., 2006).

Ferrochelatase can chelate various divalent metal ions besides Fe<sup>2+</sup>, most prominent being Zn<sup>2+</sup> (Dailey, 2003). A general mechanism for the metallation of the tetrapyrrole, proposed in 1974, suggested that deformation of the porphyrin ring after an outer-sphere complex formation between the metal ion and porphyrin would generate an appropriate configuration for metal insertion (Hambright, et al., 1974). More recent studies have looked into mechanisms that endow ferrochelatase assisted ferrous ion metallation of porphyrin. Quantum and molecular mechanical calculations showed ferrochelatases induce a thermodynamically favorable distortion of free base protoporphyrin IX (Sigfridsson, et al., 2003). Al-Karadaghi et al. recently proposed that the degree of distortion imposed by ferrochelatase modulates the type of metal ion that is inserted and this in turn is determined by subtle structural changes in the highly conserved active sites in the porphyrin binding cleft of ferrochelatase (Al-Karadaghi, et al., 2006).

Ferrochelatase requires ferrous iron as a substrate to complete heme assembly (Taketani, 2005) and the cellular mechanism that makes ferrous iron available, while also limiting the toxic effects of the metal, most likely involve the presence of a ferrous iron chaperone. Recent reports indicate frataxin is the elusive iron chaperone that delivers the ferrous iron to ferrochelatase, hence promoting heme biosynthesis in a direct manner. In frataxin deleted yeast strains, Foury et. al. observed a decrease in cytochrome c oxidase activity (Foury, et al., 1997). Later Dancis and coworkers, working with frataxin null yeast strains, observed a severe deficiency of cytochromes b, c and (a+a3) (Lesuisse, et al., 2003). Using a yeast strain with a single copy of Yfh1, placed under the control of a regulatable promoter to prevent rho minus conversion and secondary nuclear mutations that might mask the defect, the Dancis laboratory (Univ. of Penn., Philadelphia) observed the recovery of cytochromes in general and cytochrome c in particular upon re-induction of frataxin expression (Lesuisse, et al., 2003). Although frataxin deleted cells have reduced levels of ferrochelatase, it was shown that cellular heme production is modulated by the insertion of zinc instead of iron into the porphyrin ring (Lesuisse, et al., 2003). Furthermore, increasing ferrochelatase levels using a multicopy plasmid did not correct the heme synthesis defect (Lesuisse, et al., 2003). These experiments proved that mitochondrial iron located in yeast frataxin deficient cells is not available for heme synthesis. Surface plasmon resonance studies showed, in the absence of iron, recombinant Yfh1 interacts with yeast ferrochelatase with a high affinity ( $K_d = 40 \text{ nM}$ ) (Lesuisse, et al., 2003). Working towards obtaining a global result of frataxin deficiency, Schoenfeld et. al. found frataxin deficiency leads to the down regulation of mitochondrial transcripts and a kinetic inhibition of the heme pathway. An additional significant observation from their studies was a large increase in zinc chelatase activity of ferrochelatase (Schoenfeld, et al., 2005), consistent with earlier studies by Lesuisse et. al. (Lesuisse, et al., 2003). This growing evidence implicates frataxin in determining the specificity of metal for the ferrochelatase by directly binding to the protein partner.

The presence of frataxin *in vitro* has been shown to stimulate heme development under controlled conditions. *In vitro* ferrochelatase activity assays, in the presence of citrate, showed monomeric human frataxin can deliver the ferrous iron required for heme synthesis; an optimal ferrochelatase activity was observed at a stoichiometric ratio of one frataxin monomer per ferrochelatase dimer (Yoon, *et al.*, 2004). The Cowan laboratory, using isothermal titration calorimetry and fluorescence quenching experiments, were able to quantitatively investigate complex formation between human frataxin and ferrochelatase. The  $K_d$  for the interaction between HsFtx and human ferrochelatase in the presence of iron was 17 nM; no interaction was detected for the human proteins in the absence

of iron. In the yeast system, the Isaya laboratory showed ferrous iron associated with Yfh1 oligomers was available to yeast ferrochelatase for stimulation of *in vitro* heme synthesis (Park, *et al.*, 2003). Metal transfer was shown to occur in the presence of Fe(II) specific chelators, suggesting again a strong intermolecular interaction is formed at the ferrous iron site in the frataxinferrochelatase protein complex (Park, *et al.*, 2003).

Spectroscopic analysis of the yeast frataxin/ferrochelatase complex has been useful in helping to identify where and how the proteins interact. Our laboratory has shown that solution titrations, adding unlabeled yeast ferrochelatase into <sup>15</sup>N-labeled Yfh1, allowed us to probe by NMR which frataxin amino acids underwent amide chemical shift perturbations as a result of complex formation (He, et al., 2004). Frataxin chemical shift perturbations were localized predominately on the helical plane of the protein. Specific Yfh1 residues affected by ferrochelatase binding included surface exposed α1 residues His83, Glu89, Glu90, Glu93 and His95, in the metal binding region of the protein (of which all Glu's are conserved), and β6-loop-α2 residues Val150, Asn154, Thr159, Asp160, Thr163, Glu164, Ser171 and Lys172 (of which Val150, Asn154, Thr163 and Glu164 are conserved in their physical properties). Yfh1 residues in the  $\Box$ 6-loop region provide a hydrophobic patch on the protein and thus it is tempting to speculate that this domain might help target and facilitate the binding of frataxin to the hydrophobic exterior of ferrochelatase (He, et al., 2004). Complementary experiments were also performed on the human proteins by our laboratory in collaboration with the Cowan laboratory, with similar results identifying the helical residues perturbed upon complex formation (Bencze, et al., 2006). These results suggest the helical plane of frataxin, including the □1 iron-binding residues, is responsible for generating a favorable interface when frataxin binds to ferrochelatase.

Recently we performed modeling studies to predict how frataxin may interact with ferrochelatase. Using the Yfh1 solution structure (He, et al., 2004) and the Co2+ bound yeast ferrochelatase crystal structure (Karlberg, et al., 2002), we were able to generate a plausible model for the interaction between these two proteins (Figure 4) in an orientation that may suggest how iron is delivered to ferrochelatase during heme biosynthesis. We generated a large collection of complexes using the automatic docking program (ZDOCK), however we applied strict inclusion criteria that match published biophysical data when selecting appropriate docking interactions. The simulation selection criteria utilized were: a) Yfh1 residues identified in our NMR titration as perturbed by the presence of ferrochelatase, b) the location of the metal binding sites from both the human and yeast metal loaded ferrochelatase structures and c) excluding the region of ferrochelatase that docks to the membrane (Wu, et al., 2001). In our docking model, the Yfh1 monomer interacts with the yeast

ferrochelatase dimer through highly conserved frataxin helices  $\Box 1/\Box 2$  and the highly conserved ferrochelatase metal docking site including  $\Box 4$ -loop- $\Box 6$ ,  $\Box 8$  and  $\Box 11$  amino acids. Human ferrochelatase binds cobalt at this docking site using residues His231 and Asp383, located opposite to the membrane binding loops near a groove lined by arginines and lysines (Wu, *et al.*, 2001). These Lys and Arg residues could provide complementary electrostatic interactions to the acidic helices found in frataxin (Figure 1).

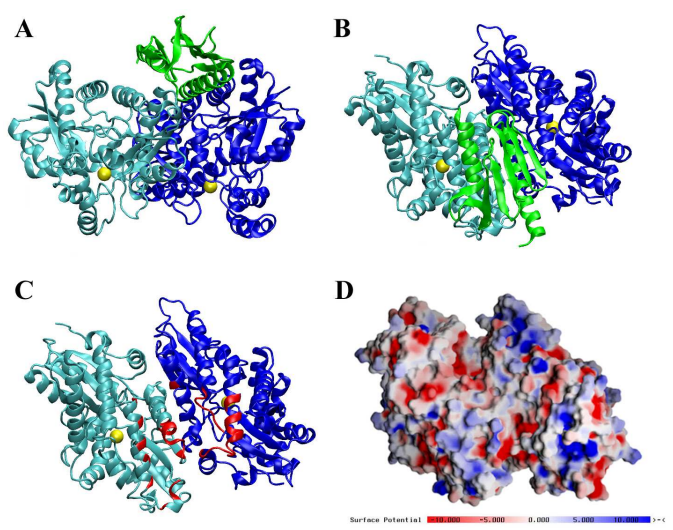


Figure 4. Lowest energy simulation of Yfh1 monomer docked to the metal loaded yeast ferrochelatase. (A) Side view of single Yfh1 monomer (green) docked to yeast ferrochelatase dimer (dark and light blue). Co<sup>2+</sup> is bound in the yeast ferrochelatase structure in the assembly active site close to the four membrane attachment lips at the bottom of the figure. (B) Side view (90° - Horizontal rotation of Figure 4A) showing monomeric Yfh1 interacts with both units in the ferrochelatase dimer. (C) Ferrochelatase side view with Yfh1 structure removed to show the residues that directly interact with frataxin (in red). (D) Electrostatic potential plots of ferrochelatase side view. Dynamics simulations were performed by VMD (Humphry, et al., 1996) while

docking was performed using ZDock (Chen, et al., 2003) using the Yfh1 structure (PDB ID# 2GA5) and Co<sup>2+</sup> loaded yeast ferrochelatase structure (PDB ID# 1L8X).

In our docking model, the Yfh1 iron-binding α1 unit docks in close proximity to the ferrochelatase cobalt binding site found in the human ortholog; in yeast, two glutamate side chains are at this site and these could function during iron binding by the ferrochelatase. Iron bound to ferrochelatase would then be free to translocate towards the heme assembly active site (i.e., the position of bound Co<sup>2+</sup> in the *yeast* ferrochelatase structure), as has been proposed by Wu *et al.* (Wu, *et al.*, 2001). These data are also consistent with the published ratio of 1:1 frataxin monomer/ferrochelatase dimer binding interactions seen in both the yeast and human systems (Lesuisse, et al., 2003; Yoon, et al., 2004). A recent report postulated protoporphyrin might help modulate the frataxin-ferrochelatase interaction, with porphyrin stabilizing the interaction (Al-Karadaghi, et al., 2006). Porphyrin was however not added to our docking simulation. Given that the entry site for porphyrin is at the ferrochelatase surface opposite to the frataxin binding site, one would expect no direct effect of porphyrin binding on frataxin-ferrochelatase affinity. However, the lowest energy normal mode explored by free ferrochelatase, and therefore the most likely to be physiologically relevant, shows that opening the porphyrin binding site closes up partially the frataxin binding site, suggesting a possible indirect effect of porphyrin on frataxin binding by the ferrochelatase and vice versa. Current studies are underway in our laboratory to determine if iron or porphyrin alters the binding affinity between the wild type yeast frataxin and ferrochelatase proteins, and if mutating surface contact residues between the partners affects complex formation and metal transfer.

# FRATAXIN AND Fe-S CLUSTER ASSEMBLY

Mounting evidence implicates frataxin also plays a direct role in Fe-S cluster biosynthesis. Fe-S clusters are among the most complex and ancient prosthetic groups found in biology. They play essential roles in cellular processes ranging from electron transport, catalysis, gene regulation and iron uptake (Chen, et al., 2004). As an illustration of their importance, there are three separate pathways in bacteria for the production of Fe-S clusters. The first pathway identified involves the nitrogen fixation machinery that assembles the Fe-S clusters required for the maturation of nitrogenase (Frazzon, et al., 2002; Rees, et al., 2000). For most cellular needs, Fe-S clusters are provided by the second pathway, involving the ISC-assembly machinery (Kispal, et al., 1999; Zheng, et al., 1998). In some bacteria, a minor contributor under stressed conditions is the third pathway,

the sulfur mobilization (SUF) machinery (Takahashi, et al., 2002). Many bacteria contain more than one of these pathways and interestingly they appear to be somewhat interchangeable (Ali, et al., 2004; Takahashi, et al., 2002). The mitochondria of eukaryotes however utilize only ISC orthologs for Fe-S cluster assembly and numerous reports outline the different characteristics of this highly important pathway (Johnson, et al., 2005; Lill, et al., 2005; Mansy, et al., 2004). A brief outline of the proteins within the yeast ISC-assembly machinery is given below as a basis for introducing frataxin's role in Fe-S cluster assembly.

Within the ISC-assembly pathway, yeast Fe-S cluster biosynthesis begins with the liberation of sulfur by the cysteine desulfurase, Nfs1 (Kispal, et al., 1999). A disulfide bond is formed between Nfs1 and free mitochondrial cysteine, leading to bond cleavage and alanine release (Lill, et al., 2005). Arh1, a ferredoxin reductase, and Yah1, a ferredoxin, are both required for efficient ISC assembly (Muhlenhoff, et al., 2003), and it has been proposed they provide the electrons necessary for sulfur liberation (Lill, et al., 2005). Next, sulfur is transferred to the cluster assembly scaffold proteins, a ~28 kDa protein dimer constructed of Isu1 (and/or) 2, usually via heterotetrameric complexes of the scaffold and the desulfurase (Garland, et al., 1999; Gerber, et al., 2003; Lill, et al., 2005; Schilke, 1999). The recently identified protein Isd11 is required to both stabilize and act as an adaptor between Nfs1 and the scaffold proteins and this helps promote sulfur release (Adam, et al., 2006; Muhlenhoff, et al., 2003; Wiedemann, et al., 2006).

A general property of the scaffold proteins is that they are dynamic in nature, hence providing a flexible platform on which to construct Fe-S clusters while retaining the ability to release the prosthetic group for transfer to future acceptor proteins (Adinolfi, Rizzo, *et al.*, 2004; Bertini, *et al.*, 2003; Mansy, *et al.*, 2004). While the structure of the yeast ortholog has not yet been solved, structures for Isu orthologs are consistent with having a β-sheet surface connected to a globular helical protein arrangement (Bertini, *et al.*, 2003; Liu, *et al.*, 2005; Ramelot, *et al.*, 2004). The structure of Zn<sup>2+</sup> bound to the *Streptococcus pyogenes* and *Haemophilus influenzae* orthologs shows divalent metal is ligated by conserved cysteine residues in a surface exposed region of the protein monomer (Liu, *et al.*, 2005; Ramelot, *et al.*, 2004).

Frataxin is required for the *in vivo* productions of Fe-S clusters and is believed to play a direct role in their assembly. A Fe-S cluster deficiency was identified in FRDA patients early in the recognition of the disorder, suggesting the need for frataxin in Fe-S cluster assembly (Rotig, *et al.*, 1997). This idea was supported by mouse knockouts showing similar disease phenotypes (Puccio, *et al.*, 2001) and more recently in the bacterial system (Vivas, *et al.*, 2006). The majority of the direct frataxin *in vivo* Fe-S cluster correlations come again however from studies in the yeast system

(Chen, et al., 2002; Duby, et al., 2002; Muhlenhoff, Richhardt, Ristow, et al., 2002). Suppressing Yfh1 expression results in respiratory deficiency, mitochondrial iron accumulation and reduced Fe-S enzyme activity while non Fe-S containing enzymes remained active (Chen, et al., 2002). Additional studies suggest frataxin may not be essential for Fe-S cluster assembly, but it does improve the efficiency of the assembly process (Duby, et al., 2002). A direct interaction between frataxin and the assembly apparatus proteins has been detected (Gerber, et al., 2003; Muhlenhoff, et al., 2003; Ramazzotti, et al., 2004) and the requirement of frataxin for the maturation of Fe-S clusters in yeast has been confirmed (Stehling, et al., 2004). Taken together, these data suggest frataxin plays a direct role in the cellular assembly of Fe-S clusters.

Recent *in vitro* studies show frataxin binds to the ISU scaffold protein with high affinity and stimulates the production of Fe-S clusters, indicating frataxin may be acting as the iron chaperone that delivers the Fe(II) required for bioassembly. Binding between *in vitro* human frataxin and the ISU dimer has been reported to occur with an affinity in the nanomolar range (Yoon, *et al.*, 2003). Binding of monomeric HsFtx with the human ISU dimer was dependent on the presence of iron, in close correlation with yeast studies showing that the interaction between Yfh1 and Isu1/Nfs1 is also iron dependent (Gerber, *et al.*, 2003). *In vitro* activity assays for the yeast and human protein systems show the presence of frataxin stimulates Fe-S cluster assembly (Muhlenhoff, Richhardt, Gerber, *et al.*, 2002; Yoon, *et al.*, 2003). Based on the fact that frataxin is an iron-binding protein, that it binds tightly to the ISU complex and finally that it stimulates Fe-S cluster assembly, it seems highly probable that frataxin can act as the iron chaperone during Fe-S clusters bioassembly.

# FRATAXIN ROLE IN ADDITIONAL PATHWAYS

Numerous reports have linked frataxin deficiency with an enhanced reduction in aconitase activity (Chen, et al., 2002; Foury, 1999; Rotig, et al., 1997). Additional reports indicate frataxin may also deliver Fe(II) to aconitase for the use in repairing oxidatively damaged [4Fe-4S]<sup>2+</sup> clusters converted to the inactive [3Fe-4S]<sup>+</sup> form (Bulteau, et al., 2004). Damage to aconitase, a Krebs-cycle enzyme that converts citrate to isocitrate (Beinert, et al., 1996), has shown to be a marker for cellular oxidative damage (Bulteau, et al., 2003). Reduced aconitase activity leads to increased transcription of the iron-import mechanism and this leads to further mitochondrial iron overload in FRDA patients (Chen, et al., 2004). Frataxin interacts in a citrate dependent manner with oxidatively damaged aconitase, promoting aconitase enzyme reactivation in a manner that suggests frataxin delivers the Fe(II) required to reactivate the damaged iron-deficient Fe-S center (Bulteau, et al., 2004). Furthermore, aconitase appears to directly associate with frataxin under high concentrations

of ROS, possibly as a means to further protect aconitase against [4Fe-4S]<sup>2+</sup> cluster disassembly, irreversible inactivation and potential degradation (Bulteau, *et al.*, 2005). ΔYfh1 yeast cells have low manganese superoxide-dismutase activity that can be recovered by supplementing Mn or limiting Fe in the media, suggesting a correlation between cellular iron overload and reduced SOD activity as a cause of oxidative stress in FRDA cells (Irazusta, *et al.*, 2006). Finally, the overexpression of frataxin in human cell lines leads to increased mitochondrial oxidative metabolism, as shown by an increased aconitase activity, mitochondrial membrane potential, cellular respiration and ATP content (Schulz, *et al.*, 2006). These data suggest frataxin provides a direct line of defense against cellular oxidative stress by controlling aconitase activity and iron import, as well as a possible additional indirect correlation between the partial regulation of the enzymatic activity of other proteins that control ROS activity.

# **OPEN QUESTIONS REGARDING FRATAXIN**

Frataxin has been shown to play an important role in regulating cellular iron homeostasis, although the exact function(s) of the protein continues to remain controversial. While numerous groups have implicated frataxin as a protein that directs the utility of bioavailable iron, many key questions remain unanswered. If frataxin is an iron chaperone, what drives this chaperone to deliver iron specifically to multiple different protein partners (i.e., ferrochelatase, ISU scaffold or aconitase) when members of the copper chaperone family are highly specific with regard to their protein partners (Rosenzweig, et al., 2000)? Certainly additional structural data of frataxin in complex with each protein partner will help provide a better understanding of how and why these proteins interact within the different pathways. How can frataxin participate in two counterproductive roles, providing bioavailable and labile Fe(II,) when acting as an iron chaperone, while at other times retaining iron as inert Fe(III) when acting in an iron storage capacity, to help prevent toxicity? Structural data regarding iron-loaded frataxin oligomers will surely help address this issue of how and why these assemblies form and identifying conditions when these assemblies can be observed *in vivo* will provide additional insight into their physiological relevance. Finally, what is the dominant function of frataxin? Frataxin has recently been shown to interact with the succinate dehydrogenase complex subunits, implying an additional possible role for frataxin in

mitochondrial electron transport (Gonzalez-Cabo, et al., 2005). Frataxin has also been shown to be a key activator of mitochondrial energy conversion and oxidative phosphorylation, suggesting yet another additional role for the protein in energy production (Ristow, et al., 2000). Finally, CyaY has recently been shown to interact with the cysteine desulfurase IscS to promote both ferric iron reduction and Fe-S cluster assembly, suggesting iron delivery by frataxin may only occur when coupled with iron reduction (Layer, et al., 2006). In vivo mutational studies that split the individual phenotypes for each pathway will provide biological relevance to help understand frataxin's role in each of the many functions proposed for this important protein. Addressing these and other equally important issues regarding how frataxin functions will surely help drive research directed at developing improved treatment strategies that better help control iron regulation in disorders like Friedreich's ataxia.

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