Overlapping Binding Sites of the Frataxin Homologue Assembly Factor and the Heat Shock Protein 70 Transfer Factor on the Isu Iron-Sulfur Cluster Scaffold Protein*  

Received for publication, July 14, 2014, and in revised form, September 12, 2014  Published, JBC Papers in Press, September 16, 2014, DOI 10.1074/jbc.M114.596726  
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Background: Protein factors function in both FeS cluster assembly on and transfer from the Isu scaffold protein.  
Results: Common Isu residues are critical for binding both frataxin homologue (Yfh1) assembly factor and heat shock protein 70 (Hsp70) transfer factor.  
Conclusion: Yfh1 and Hsp70 binding is mutually exclusive.  
Significance: Mutual exclusive binding of assembly/transfer factors may be important in transitioning between cluster assembly and transfer.  

In mitochondria FeS clusters, prosthetic groups critical for the activity of many proteins, are first assembled on Isu, a 14-kDa scaffold protein, and then transferred to recipient apoproteins. The assembly process involves interaction of Isu with both Nfs1, the cysteine desulfurase serving as a sulfur donor, and the yeast frataxin homolog (Yfh1) serving as a regulator of desulfurase activity and/or iron donor. Here, based on the results of biochemical experiments with purified wild-type and variant proteins, we report that interaction of Yfh1 with both Nfs1 and Isu are required for formation of a stable tripartite assembly complex. Disruption of either Yfh1-Isu or Nfs1-Isu interactions destabilizes the complex. Cluster transfer to recipient apoprotein is known to require the interaction of Isu with the J-protein/Hsp70 molecular chaperone pair, Jac1 and Ssq1. Here we show that the Yfh1 interaction with Isu involves the PVK sequence motif, which is also the site key for the interaction of Isu with Hsp70 Ssq1. Coupled with our previous observation that Nfs1 and Jac1 binding to Isu is mutually exclusive due to partially overlapping binding sites, we propose that such mutual exclusivity of cluster assembly factor (Nfs1/Yfh1) and cluster transfer factor (Jac1/Ssq1) binding to Isu has functional consequences for the transition from the assembly process to the transfer process, and thus regulation of the biogenesis of FeS cluster proteins.

In all three domains of life proteins having iron-sulfur (FeS) cluster prosthetic groups play important roles in critical cellular processes such as redox reactions, catalysis, and sensing of stress (1, 2). In eukaryotic cells, mitochondria, which inherited their iron-sulfur cluster (ISC)4 biogenesis system from bacterial ancestors, play an essential role in maturation of FeS cluster containing proteins functioning in all subcellular compartments (3). The proteins involved in FeS clusters biogenesis are highly conserved. In eukaryotic cells, the majority of these proteins are essential for cell viability, underscoring the biological importance of FeS cluster proteins. In both mitochondrial and bacterial ISC systems, a highly conserved 14-kDa protein (called Isu in the yeast Saccharomyces cerevisiae) plays a central role in FeS cluster biogenesis, serving both as a scaffold for de novo cluster assembly and as a platform for cluster transfer onto recipient apoproteins (3). Both the assembly and transfer steps of FeS cluster biogenesis involve the interaction of Isu with protein factors (4). Because the ISC system of S. cerevisiae is the focus of this report, we use the gene/protein nomenclature for this organism throughout. In S. cerevisiae, Isu is encoded by the closely related and functionally exchangeable paralogous genes, ISU1 and ISU2 (5, 6). Isu1 plays the major functional role due to its higher level of expression.

De novo assembly of FeS clusters requires Isu interaction with two proteins, the cysteine desulfurase Nfs1, which releases sulfur from cysteine (7), and the yeast frataxin ortholog, Yfh1, which is a putative iron donor and/or regulator of desulfurase enzymatic activity (8–11). Yfh1 and Nfs1 interact both with each other and with Isu, thereby forming the FeS cluster assembly complex, which constitutes the structural and functional unit responsible for cluster synthesis within the Isu scaffold (9, 10, 12–16). In bacteria the cysteine desulfurase (IscS) functions as a single polypeptide. However, in eukaryotes, Nfs1 functions as a stable heterodimer in complex with a small accessory pro-

* This work was supported, in whole or in part, by National Institutes of Health Grant GM27870 (to E. A. C), Polish National Science Center Grant DEC-2012/06/A/N21/00002 (to J. M.), and Foundation for Polish Science International PhD. Projects Grant MPD/2010/5 co-financed by the European Union European Regional Development Fund, Operational Program Innovative Economy 2007–2013 (to J. M.).  
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tein Isd11 (referred to as Nfs1(Isd11) throughout) (17–21). Isd11 is proposed to both stabilize Nfs1 and regulate its catalytic activity (22). Although the role of Nfs1(Isd11) as a sulfur donor for FeS cluster synthesis is indisputable, the role of frataxin, which is associated with the neurodegenerative disease Friedreich ataxia (23) in this process is hotly debated (24, 25). Recent experimental data strongly support the view that the major biological function of frataxin is directly related to its role as a component of the FeS cluster assembly complex, but whether it serves as an iron donor, regulator of desulfurase activity, or both is unresolved (26–28).

In both mitochondria and bacteria, FeS cluster transfer from Isu onto recipient apoproteins requires a dedicated J-protein/ Hsp70 molecular chaperone pair (29). Similar to other members of the J-protein class (30), Jac1 binding to its client protein (i.e. Isu) targets it for binding to its Hsp70 partner (i.e. Ssq1) (31, 32). It is thought that conformational changes of the cluster-loaded holo-Isu induced upon Hsp70 binding triggers the release of the FeS cluster and its transfer onto a recipient apoprotein (33–35). In both bacteria and mitochondria, the C-terminal domain of Jac1 binds to a specific region on the surface of Isu (36–38), with the most important players of a large interaction interface being three hydrophobic residues of Jac1 (32) interacting with three hydrophobic residues of Isu1 (39). The Hsp70-Isu interaction is also highly sequence specific, interacting via a highly conserved Pro, Val, Lys (PVK) motif (40, 41), which is part of a flexible loop localized in proximity to the Hsp binding pocket (35).

Although the role of the assembly complex for de novo FeS cluster synthesis is well established (3, 26) and the importance of the Hsp70/Jac1 chaperones for FeS cluster transfer to recipient apoproteins has solid experimental support (29), relatively little is known about the specific interactions required for assembly complex formation and how the molecular architecture and protein-protein interactions involved in its formation affect transition to the chaperone-mediated transfer reaction. To address these issues, we have pursued structure/function analyses, using S. cerevisiae as a model system. Previously we showed that Nfs1 and Jac1 utilize overlapping binding sites on Isu and that their interactions with Isu are mutually exclusive (39). Here, we show that the stable association of Yfh1 within the tripartite FeS cluster assembly complex is dependent on both its Nfs1/Isd11- and Isu1-binding interfaces. Furthermore, we demonstrate that Yfh1 binding to Isu1 involves the PVK sequence motif, which is also the binding site for Ssq1 (41). Thus binding of assembly complex factors (Nfs1/Yfh1) to form the assembly complex and action of the transfer factors (Jac1/ Ssq1) are mutually exclusive. We hypothesize that this exclusivity likely has functional consequences for the transition from the cluster assembly step to the cluster transfer step in the biogenesis of FeS cluster proteins.

EXPERIMENTAL PROCEDURES

Yeast Strains, Plasmids, Media, and Chemicals—All strains were of the S. cerevisiae W303 background. Cells expressing Yfh1W131A were created by sporulation and dissection of a yfh1Δ/YFH1 strain (42) harboring a centromeric plasmid carrying the yfh1W131A gene under control of the YFH1 promoter and LEU2 marker. To assess the level of Nfs1 variants, strains having the WT gene at the normal chromosomal location under the control of the glucose-repressible GAL1–10 promoter were used (43). To construct an expression vector for the GST-Yfh1 fusion protein, the mature sequence of YFH1 (codons 52–174) was cloned into the glutathione S-transferase (GST)-expressing vector pGEX-KG (44) using BamHI and XhoI restriction sites. YFH1, ISU1, and NFS1 mutants were generated in pRS315- YFH1, pRS314-ISU1, and pRS316-NFS1 using the Stratagene QuikChange protocol (41), as were all mutants in Escherichia coli expression vectors. yfhΔ (42) and nfs1Δ (43) strains were described previously. Yeast were grown on YPD (1% yeast extract, 2% peptone, and 2% glucose) or synthetic media as described (45). All chemicals, unless stated otherwise, were purchased from Sigma.

Protein Purification—The sequences encoding the mature form of Yfh1 (residues 52–174) with a C-terminal polyhistidine tag was cloned into pET-3a (Novagen, Madison, WI). Protein overexpression was induced for 4 h at 30 °C in E. coli C41(DE3) by adding isopropyl 1-thio-D-galactopyranoside at a final concentration of 1 mM. Cells were harvested and lysed using a French press in buffer Y1 (50 mM Tris-HCl, pH 8.0, 50 mM NaCl, 1 mM phenylmethanesulfonyl fluoride (PMSF), 10% glycerol). The cell lysate was subjected to His-Bind Resin (Novagen) chromatography. After washing steps with buffer Y1, Y2 (Y1 + 0.5 M NaCl), and Y3 (Y1 + 20 mM imidazole), proteins were eluted with a linear 20–200 mM imidazole gradient in buffer Y1. Fractions containing pure Yfh1 protein, were collected, pooled, concentrated on Centriprep 10K (Millipore), dialyzed against buffer F (50 mM Tris-HCl, pH 8.0, 10% glycerol, and 50 mM NaCl), and stored at −70 °C.

Expression of the mature form of GST-Yfh1 and GST protein alone were induced in the E. coli strain C41(DE3) carrying the pGEX-KG-YFH1 or pGEX-KG plasmid, respectively, by addition of 1 mM isopropyl 1-thio-D-galactopyranoside at A600 = 0.6. After 4 h growth at 30 °C, cells were harvested and lysed using a French press in buffer L (25 mM Tris-HCl, pH 8.0, 200 mM NaCl, 1 mM PMSF, 1 mM dithiothreitol (DTT), 10% glycerol, and 0.05% Triton X-100). After a clarifying spin, the supernatant was loaded onto a 1-ml glutathione-agarose column (Fluka) equilibrated with 10 volumes of buffer L. Next, the column was washed with 100 ml of buffer L without PMSF, 10 volumes of buffer L with 0.5 M NaCl, and 10 volumes of buffer L with 10 mM MgCl2 and 1 mM ATP. After a final wash with 10 volumes of buffer L, proteins were eluted with buffer E (25 mM Tris-HCl, pH 8.0, 200 mM NaCl, 1 mM DTT, 10% glycerol, 0.05% Triton X-100, and 50 mM reduced glutathione). Fractions containing GST-Yfh1 (or GST) were pooled, dialyzed against buffer F (50 mM Tris-HCl, pH 8.0, 10% glycerol, and 50 mM NaCl), and stored at −70 °C.

The mature form of Ssq1 (residues 19–655) with a C-terminal polyhistidine tag was purified from E. coli strain BL21(DE3) that co-expressed SSQ1 and HEPI, which encodes a small accessory protein, from the plasmid pRSFDuet1-SSQ1His/HEP1 (46). Hep1 stabilizes Ssq1 when expressed in the bacterial cells but, as Ssq1 does not form a stable complex with Hep1, it is not present in the Ssq1 final preparation. Expression was induced by addition of 1 mM isopropyl 1-thio-D-galactopyranos-
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side at $A_{600} = 0.6$ and cells were grown at 25 °C for 4 h. Cells were harvested, resuspended in C1 buffer (20 mM HEPES-KOH, pH 7.5, 150 mM KCl, 20 mM imidazole, 10% glycerol, 1 mM PMSF, 2 mM magnesium acetate), and lysed using a French press. After a clarifying spin, the supernatant was loaded onto His-Bind Resin (Novagen) equilibrated in buffer C1 and after washing with buffer C2 (C1 + 0.05% Triton X-100), C3 (C1 + 10 mM MgCl$_2$, 1 mM ATP), and C4 (C1 + 1 M KCl), proteins were eluted with a 40–250 mM imidazole gradient in buffer C1. Fractions containing pure Ssq1 were collected, pooled, and dialyzed to buffer C5 (20 mM HEPES-KOH, pH 8.0, 100 mM KCl, 10% glycerol, 5 mM β-mercaptoethanol). Ssq1 was aliquoted and stored at −70 °C.

Wild-type (WT) and mutant Nfs1(Isd11), possessing a polyhistidine tag at the N terminus, were purified as described previously (39). WT and mutant Jac1, with a polyhistidine tag at the C terminus, were purified as described previously (31), except E. coli strain C41(DE3) was used for expression. Recombinant Isu1 with a polyhistidine tag at the C terminus, and WT and mutant Isu1-GST fusion proteins were purified as described previously (Refs. 31 and 32, respectively). In all cases, the purity of WT and variant protein preparations, as judged by SDS-PAGE separation and Coomassie Blue staining, was determined using the Bradford (Bio-Rad) assay with 2-fold concentrated Laemmli sample buffer, and analyzed as described previously (39). WT and variant Nfs1(Isd11) variants in whole cell lysates a culture volume equivalent to 1 ml of $A_{600} = 1.0$ was harvested. The pellet was resuspended in 100 µl of 2-fold concentrated Laemmli sample buffer followed by addition of 15 µl of glass beads (0.5 mm). Cell lysates were prepared by bead-beating at 4 °C for 3 min in a cell disruptor. Proteins were solubilized by incubation at 100 °C for 5 min. Insoluble material was removed by centrifugation, and proteins in the supernatant were separated using SDS-PAGE gels. The resolved proteins were transferred electrophoretically to nitrocellulose. Yfh1 or Nfs1 variants were detected by the enhanced chemiluminescence method (48) using anti-Yfh1 or anti-Nfs1 polyclonal antiserum. When analyzing mitochondrial protein, mitochondria (containing 2.5 µg of protein) were solubilized in 2-fold concentrated Laemmli buffer, and analyzed as described above.

**Evolutionary Analysis**—Amino acid sequences of orthologs of the scaffold protein (ISU1), cysteine desulfurase (NFS1), and frataxin (YFH1) were obtained by retrieving sequences from InterPro (49) database entries: IPR011339 for ISU1, IPR010240 for NFS1, and IPR002908 for YFH1. The Proteobacterial dataset included all sequences listed under the taxon Proteobacteria in InterPro except sequences listed under Alphaproteobacteria, which constituted their own dataset. The Eukaryotic dataset included sequences listed under Eukaryota. The retrieved sequences were filtered to include only sequences from species for which orthologs of all three genes were present in each dataset. Sequences were aligned using MAFFT version 7.123b with default options (50) and amino acid frequencies were determined at positions homologous to Isu1, Nfs1, and Yfh1 from *S. cerevisiae* (complete data available on request from J. M.).

**RESULTS AND DISCUSSION**

*Stable Yfh1 Binding Requires the Presence of both Nfs1(Isd11) and Isu1*—As a first step in defining the molecular mechanism of the interaction of Yfh1 with its binding partners Nfs1(Isd11) and Isu1, we developed a semiquantitative pulldown assay that allowed us to measure protein-protein interactions in vitro. Encouraged by the earlier finding that a GST-Yfh1 fusion protein is fully functional in vivo (8), we developed a pulldown assay. GST-Yfh1 (2.5 µM) was preincubated with an excess of both Nfs1(Isd11) and Isu1 (15 µM each). Both Nfs1(Isd11) and Isu1 were efficiently pulled down along with GST-Yfh1 (Fig. 1A). In contrast, if Isu1 was left out of the reaction, no complex between GST-Yfh1 and Nfs1(Isd11) was detected above background levels. Similarly, we did not observe GST-Yfh1 interaction with Isu1 if Nfs1(Isd11) was left out of the reaction. Next, we varied the concentration of Nfs1(Isd11) or Isu1 independently, keeping the other in excess (15 µM compared with 2.5 µM GST-Yfh1). When GST-Yfh1 and Isu1 were incubated with...
increasing concentrations of wt Nfs1(Isd11) from 1 to 15 μM (Fig. 1B), Nfs1(Isd11) bound to GST-Yfh1 in a concentration-dependent manner. The opposite was also the case when the concentration of Isu1 was varied. These results indicate that Yfh1 formed a stable complex with Isu1 and Nfs1(Isd11), but only when both binding partners were present in the reaction mixture.

**Disruption of Isu1-Nfs1 Interaction Prevents Stable Binding of Yfh1 to Either Nfs1(Isd11) or Isu1**—The results described above are consistent with the idea that the physical interaction between Isu1 and Nfs1(Isd11) is required for stable interaction of Yfh1. To test this idea directly, we took advantage of an Isu1 variant, Isu1L63A/V72A/F94A, which we previously demonstrated to be defective in interacting with Nfs1(Isd11) (39). Indeed, in the presence of Isu1L63A/V72A/F94A, neither Nfs1(Isd11) nor Isu1L63A/V72A/F94A were pulled down with GST-Yfh1 (Fig. 1C and D), indicating that formation of a stable Isu1-Nfs1(Isd11) complex is required for Yfh1 binding to form the tripartite assembly complex.

It had been previously demonstrated with mammalian orthologs that Yfh1 stimulates the enzymatic activity of cysteine desulfurase, but only in the presence of Isu, whereas Isu, in the absence of Yfh1, inhibits desulfurase activity (2, 9, 10, 51). We decided to take advantage of this functional test of the interaction between Yfh1 and its binding partners. First, we carried out controls to determine whether the yeast proteins behaved similarly to their mammalian orthologs. We observed results similar to those reported for the mammalian proteins. When Yfh1 was added to Nfs1(Isd11) no effect on desulfurase activity was observed. However, when Isu1 was added to Nfs1(Isd11) cysteine desulfurase activity was reduced by ~70% (Fig. 1E). Upon addition of Yfh1 and Isu1 together desulfurase activity was stimulated ~1.7-fold. In contrast, when Isu1 WT was replaced by Isu1L63A/V72A/F94A, the variant defective in interaction with Nfs1, no inhibitory effect on Nfs1 desulfurase activity was observed. Moreover, addition of Yfh1 in the presence of Isu1L63A/V72A/F94A had no stimulatory effect on the enzymatic activity of Nfs1.

Together, the results described above strongly support the idea that Yfh1 only stably interacts with Isu and/or Nfs1(Isd11) when they are in complex. Formation of stable Nfs1(Isd11)-Isu orthologous complexes was demonstrated previously by biochemical experiments utilizing purified proteins (9, 14, 16, 39, 52) or mitochondrial and cellular extracts (10, 15). It was also postulated (15) that Yfh1 binds the pre-formed Nfs1(Isd11)-Isu complex. Here we demonstrate that the simultaneous presence of Yfh1 binding partners Isu1 and Nfs1(Isd11) is not sufficient for the tripartite assembly complex formation, as when they are unable to interact with each other due to substitutions that prevent Isu1 interaction with Nfs1(Isd11), stable Yfh1 binding does not occur.
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Prediction of Amino Acid Residues Involved in Yfh1 Interactions with Isu1 and Nfs1(Isd11)—The biochemical analysis described above suggests that Yfh1 stably interacts with a preformed Isu1-Nfs1(Isd11) complex. Mechanistically, this requirement could be interpreted in two different ways. (i) Simultaneous interactions of Yfh1 with both Isu1 and Nfs1(Isd11) are required for tripartite complex stability, due to the additive effect of two “low affinity” interactions. (ii) Yfh1 interacts with a single component (i.e. either Nfs1 or Isu1), but the “high affinity” interaction site is only present in the Isu1-Nfs1(Isd11) complex, not the individual proteins, due to a conformational change caused upon interaction of Isu1 and Nfs1(Isd11).

To understand the critical interactions needed for formation of the tripartite assembly complex, we decided to dissect the Yfh1-Isu1 and Yfh1-Nfs1(Isd11) interactions, with a goal of isolating variants that severely affected each individual interaction. As no structural data are available for the mitochondrial FeS cluster assembly complex, we took advantage of a structural model of the homologous bacterial complex (13). In this model, cysteine desulfurase (IscS) and Isu scaffold (IscU) interact with each other and with a monomer of bacterial frataxin (CyaY), which inserts deeply into the cleft formed between IscU and IscS (Fig. 2). Thus, CyaY could form binding interfaces with both IscU and IscS. Using this model (13), together with published experimental data from a variety of systems (8–12, 15, 53–55), we predicted amino acid residues critical for the Yfh1 interaction with each binding partner. Our predictions are depicted in Fig. 2. Below, we describe in more detail our rationale behind each prediction and present its experimental validation.

Trp131 Is Critical for Yfh1 Interaction with Isu1—Several lines of evidence indicate that the evolutionarily conserved Trp131 is critical for interaction of Yfh1 with Isu1 (2, 15, 20, 55, 56). Using our in vitro assay, we tested whether GST-Yfh1 with Trp131 replaced with alanine was able to pull down Isu1 and Nfs1(Isd11) (Fig. 3A). The ability of GST-Yfh1W131A to form the assembly complex, as monitored by Isu1 binding, was strongly reduced. Even at the highest Isu1 concentration (15 µM) binding was reduced by ~90%. Moreover, consistent with the reduced binding ability, Yfh1W131A did not stimulate desulfurase activity when mixed with Nfs1(Isd11) and Isu1 (Fig. 3B). To assess the in vivo effects of the Trp131→Ala alteration, we constructed a yfh1W131A strain. yfh1W131A cells grew slowly (Fig. 3C), even though the level of expression of Yfh1W131A was similar to that of WT as determined using immunoblot analysis (Fig. 3D). Together, these results suggest that Trp131 plays a critical role in Yfh1 binding to Isu1 to form the tripartite assembly complex.

Isu1 Residues Pro134, Val135, Lys136, Which Are Critical for Ssa4 Binding, Are Also Important for Yfh1 Interaction with Isu1-Nfs1(Isd11) Complex—In analyzing the bacterial model (Fig. 2) to predict residues of Isu1 potentially involved in interaction with Yfh1 we noticed that a flexible loop of IscU having the evolutionary invariant Pro, Val, Lys (PVK) motif (29, 35) is in close proximity to bacterial frataxin Trp61, a site homologous to Yfh1. This juxtaposition was very intriguing, as this

![Figure 2. Structural model of bacterial FeS cluster assembly complex.](Image)

![Figure 3. Replacement of Yfh1 Trp131 by alanine results in defective interaction with Isu1 of the Isu1-Nfs1(Isd11) complex.](Image)
PVK motif is the only known binding site of the Hsp70s necessary for cluster transfer in the bacterial and mitochondrial systems (29, 41). To test the idea that the PVK motif might be critical for Yfh1, as well as Hsp70, binding, we tested an Isu1 variant having alanines substituted for Pro134, Val135, and Lys136 (Isu1PVK/AAA) in our GST-Yfh1 pulldown assay. The amount of Isu1PVK/AAA pulled down by GST-Yfh1 in the presence of excess Nfs1(Isd11) was at the background level (Fig. 4A), indicating the importance of the PVK motif. Next we tested single alanine substitutions of the PVK motif (Fig. 4A). Altering Lys136 had the most severe effect on binding, with alteration of Pro134 or Val135 only moderately reducing binding. At the highest Isu1K136A concentration used (15 μM) the amount of Isu1K136A pulled down by GST-Yfh1 was reduced by ~80% (Fig. 4A). As a control, we also tested whether the PVK alteration affected interaction with Nfs1(Isd11), as stable Yfh1 binding to Isu1 requires the Isu1-Nfs1(Isd11) interaction (see Fig. 1). We used our previously published Isu1-GST pulldown assay (39). Each of the Isu1PVK/AAA variants pulled down Nfs1(Isd11) as efficiently as the WT control (Fig. 4B), indicating that substitutions within the PVK motif did not affect the ability of Isu1 to form a stable complex with Nfs1(Isd11). We conclude that in the context of assembly complex formation, the PVK motif of Isu1 is directly involved in Yfh1 binding.

We also compared the relative importance of the three residues of the PVK motif for interaction with Yfh1 and Hsp70 Ssq1. We incubated Isu1PVK/AAA-GST variants (2.5 μM) with excess purified Ssq1 protein (4 μM). We also included substoichiometric amounts of Jac1 co-chaperone, as it serves to target Isu1-GST for Ssq1 binding, thus promoting formation of a stable Ssq1-Isu1-GST complex (31, 41) (Fig. 4D and E). When the Isu1PVK/AAA-GST variant was used, the amount of Ssq1 pulled down was reduced to background levels, even at the highest Jac1 concentration tested (1 μM) (Fig. 4D and E). However, individual residues of the PVK motif had different contributions to Ssq1 binding compared with Yfh1 binding. For example, whereas the Lys136→Ala substitution had the strongest effect on Yfh1 binding (Fig. 4A) its effect on interaction of Ssq1 was the weakest (Fig. 4E). In contrast, the strongest defect for...
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Ssq1 binding was observed with the Pro$^{134}$ → Ala substitution, which had only a moderate effect on Yfh1 binding. We conclude that the PVK motif plays a critical role for Isu1 interaction with both Yfh1 and Ssq1, but individual residues contribute differently to the two interactions. However, the dependence of both on the PVK motif for binding, suggests that Yfh1 and Ssq1 interactions with Isu1 are mutually exclusive.

Because of the potential importance of such mutual exclusivity, we also tested the effect of the alanine replacements within the PVK motif on the cysteine desulfurase activity of Nfs1(Isd11) (Fig. 4C). In the presence of the Isu1PVK/AAA variants, Yfh1 displayed a reduced ability to stimulate desulfurase activity. These defects are consistent with the reduced ability of Yfh1 to bind Nfs1(Isd11)-Isu1 complexes having substitutions within the PVK motif.

Interestingly, addition of either Isu1PVK/AAA or Isu1K136A to Nfs1(Isd11) revealed an unexpected behavior. These two Isu1 variants did not inhibit desulfurase activity, despite the fact that both bind Nfs1(Isd11) as efficiently as WT protein (Fig. 4B). In contrast, the Isu1P134A and Isu1V135A variants significantly inhibited desulfurase activity as well as WT Isu1. These results point to the interpretation that the PVK motif, and more specifically, the Lys$^{136}$ residue may play a significant role in the inhibition of the desulfurase enzymatic activity. However, the mechanistic basis of this inhibition and its biological importance are not well understood.

**Charged Residues of Yfh1 and Nfs1 Are Critical for Their Interaction in the Context of the Assembly Complex Formation**—Structural and experimental data available for bacterial homologs (Fig. 2) (13) and experimental results published for mitochondrial proteins (24, 53–55, 57) suggested negatively charged Asp$^{86}$ and Glu$^{89}$ of Yfh1 as potential interacting partners with positively charged Arg$^{313}$, Arg$^{316}$, and Arg$^{318}$ of Nfs1. Reasoning that replacement by alanines may have a lesser effect compared with charge reversal, we purified variants of Yfh1 and Nfs1 having substitutions of the residues to either alanine (Yfh1DE/AA and Nfs1RRR/AAA) or to a residue of the opposite charge (Yfh1DE/KK and Nfs1RRR/EEE). Indeed, although Yfh1DE/AA had a markedly reduced ability to pull down Nfs1(Isd11) and Isu1, no Yfh1DE/KK binding above background levels was observed (Fig. 5A). The Yfh1DE/KK variant was also unable to stimulate Nfs1(Isd11) desulfurase activity in the presence of Isu1 (Fig. 5B). Moreover, cells expressing the DE/KK variant as the only copy of Yfh1, displayed a slow growth phenotype at all temperatures tested (Fig. 5C and data not shown), despite the fact that the Yfh1DE/KK was expressed at levels comparable with WT (Fig. 5D). These results are consistent with Asp$^{86}$ and Glu$^{89}$ playing a significant role in Yfh1 interaction with Nfs1(Isd11) to form the assembly complex, as well as verifying the observation that a stable association of Yfh1 with both Nfs1(Isd11) and Isu1 requires simultaneous interactions with both binding partners.

Analysis of the Nfs1RRR variants revealed that either substitution to alanine or to glutamic acid caused a severe reduction in binding. No detectable pulldown by GST-Yfh1 was observed in either case, indicating that Arg$^{313}$, Arg$^{316}$, and Arg$^{318}$ are critical for Yfh1 binding. To ensure the validity of our interpretation, we tested whether Nfs1RRR/AAA and Nfs1RRR/EEE were able to form a stable complex with Isu1, using the Isu1-GST pull-down assay as described above. We found that both Nfs1 variants formed the Nfs1(Isd11)-Isu1-GST complex as efficiently as WT protein (Fig. 6B). We also found that Yfh1 was unable to stimulate the desulfurase activity of either Nfs1RRR/AAA(Isd11) or Nfs1RRR/EEE(Isd11), despite the fact that the basal enzymatic activity of these variants was similar to the WT control (Fig. 6C).

We also tested the effect of these alterations on in vivo function. Cells expressing either the RRR/AAA or RRR/EEE variant as the only copy of Nfs1 were inviable (Fig. 6D). To ensure that the null phenotypes were due to altered protein function, not low expression, we used a yeast strain in which Nfs1 WT expression was driven by the GAL-10 promoter and, thus, repressed upon glucose addition. After transformation of these cells with a plasmid harboring a mutant NFS1 allele under control of the native NFS1 promoter, cultures were shifted from galactose- to glucose-based medium. WT Nfs1 was depleted...
below the level of immunodetection, whereas the levels of both Nfs1\textsuperscript{RRR/AAA} and Nfs1\textsuperscript{RRR/EEE} were similar to that of Nfs1 in a WT strain (Fig. 6E). We conclude that, despite normal expression levels, neither the RRR/AAA nor the RRR/EEE Nfs1 variants are able to support cell growth.

These \textit{in vivo} results were unexpected, as replacement of the residues forming the apparently complementary binding interface of Yfh1 resulted in a slow growth, not a null, phenotype (see Fig. 5C). One possible explanation is that our model-based prediction of residues forming the Yfh1-Nfs1 interface is not precise. The three residues substituted on the Nfs1 side could be contributing more to the interaction than the two residues substituted on the Yfh1 side. This interpretation is consistent with our observation that Yfh1\textsuperscript{DE/AA} had markedly reduced, but measurable, binding capacities in the pulldown assay (Fig. 5A), whereas no Nfs1\textsuperscript{RRR/AAA} binding was detected (Fig. 6A). Alternatively, this site of Nfs1 may be involved in more than one protein-protein interaction. Data published recently for the bacterial system (58, 59), indicates that the homologous RRR residues of the Nfs1 ortholog are not only involved in binding of bacterial frataxin, but are also critical for interaction with ferredoxin, a protein whose activity is needed for the reduction of sulfate during the process of FeS cluster assembly (3, 59). As ferredoxin is an essential component of the FeS cluster biogenesis system, reduced binding by the Nfs1\textsuperscript{RRR/AAA} variant could well explain the null phenotype. In addition, these RRR residues of Nfs1 were also identified as part of a highly conserved RRRPR motif of Nucleotide Localization Sequences (NLS) (60, 61). Substitutions within the NLS may potentially prevent targeting of Nfs1 to the nucleus and thus affect its nuclear function(s). Additional studies are needed to test these possibilities.

Residues Involved in Yfh1 Interaction with Isu1 and Nfs1(Isd11) in the Context of the FeS Cluster Assembly Complex Formation Are Evolutionary Conserved—To place our experimental findings in a wider evolutionary context, we identified residues homologous to those involved in the Yfh1-Isu1 and Yfh1-Nfs1 interactions across phylogeny. First, we identified orthologs of Yfh1, Isu1, and Nfs1 in fully sequenced genomes of 422 bacterial and 199 eukaryotic species. Our analyses revealed that residues involved in Yfh1 interaction with Isu1 are extremely evolutionarily conserved, as both Trp\textsuperscript{131} of Yfh1 and Pro\textsuperscript{134}, Val\textsuperscript{135}, and Lys\textsuperscript{136} of Isu1 are invariant across all analyzed orthologous sequences (Fig. 7). In the case of residues involved in Yfh1 interaction with Nfs1, we observed that sites occupied by Asp\textsuperscript{86} and Glu\textsuperscript{89} of Yfh1 are occupied mostly by acidic residues in both bacterial and eukaryotic orthologs, indicating that the chemical nature of the Yfh1 interaction with Nfs1 is evolutionary conserved. An even higher level of conservation was observed on the Nfs1 side of the interaction. In most orthologs, sites occupied by Arg\textsuperscript{313}, Arg\textsuperscript{316}, and Arg\textsuperscript{318} of Nfs1 were invariant across phylogeny. In \alpha-proteobacteria, Arg\textsuperscript{313} was replaced by Lys in most orthologs, which maintains the chemical nature of the site involved in Yfh1 interaction. Overall, the phylogenetic analyses revealed a high conservation of the residues involved in interaction of Yfh1 with Isu1 and Nfs1, supporting the idea that that the molecular mechanism of the interaction of these proteins in the context of the FeS assembly complex formation, is also highly evolutionarily conserved.

\textit{Conclusions}—Our results indicate that formation of a stable tripartite assembly complex requires the interaction of Yfh1 with both its binding partners, Nfs1(Isd11) and Isu1. In the alternative hypothesis, that the interaction between Isu1 and
Nfs1(Isd11) caused a conformational change in one of the partners resulting in formation of a high affinity binding site. If this were true, we would have expected to find one of the interactions to be of predominant importance. This was not the case, as similar effects of individual binding site replacements were observed, regardless of whether formation of the tripartite assembly complex was detected by the GST based pulldown technique or by measuring Nfs1(Isd11) desulfurase activity. Overall, an ordered maturation of the FeS cluster assembly complex, with Isu1 and Nfs1(Isd11) interacting first and Yfh1 binding to the preformed Nfs1(Isd11)-Isu1 heterodimer via interactions with both proteins, is the most likely scenario in vivo. The Isu-Nfs1(Isd11) heterodimer is readily detectable under a variety of conditions both in vivo and in vitro (9, 10, 14–16, 39, 52). Although interaction between Yfh1 (or its orthologs) and Isu and Nfs1 individually have been detected under some conditions (12, 13, 62), individual interactions among pairs of components of the tripartite complex are possible (Nfs1(Isd11)-Yfh1 or Yfh1-Isu1), yet, an ordered mechanism is likely dominant, due to the high stability of the Nfs1(Isd11)-Isu1 heterodimer.

We also conclude from our results that the PVK motif of Isu is involved in both critical stages of FeS cluster biogenesis, cluster assembly, and cluster transfer, as it is necessary for binding of Yfh1 as well as Hsp70 Ssq1. We previously demonstrated that two other proteins involved in cluster assembly (Nfs1) and cluster transfer (Jac1) also share overlapping, mutually exclusive binding sites on the surface of Isu1 (Fig. 8A) (39). Together these results suggest that exclusivity of binding could have important functional consequences, initiating and directing the ordered transition from the FeS assembly complex to the FeS cluster transfer complex.
transfer reaction catalyzed by the Hsp70 chaperone system. How might such exclusivity help orchestrate such a progression? We proposed previously (39) that Jac1 co-chaperone facilitates displacement of holo-Isu1 from the FeS assembly complex by interfering with the Nfs1-Isu1 interaction. Results presented here indicate that disruption of the Isu1-Nfs1 (Isd11) interaction also destabilizes the Isu1-Yfh1 interaction, as the latter depends on Nfs1 (Isd11)-Isu1 binding. Thus, holo-Isu1 released from Nfs1 (Isd11) by the action of Jac1 will also result in Yfh1 release, thereby exposing the PVK site for Hsp70 interaction (Fig. 8B). Once Ssq1 binds the PVK motif, Isu1 is “protected” against Yfh1 rebinding, even if Nfs1 (Isd11) rebinds to Isu1 after dissociation of Jac1. Thus the chaperones may play a regulatory role by controlling the flow of FeS clusters from the assembly complex to the recipient proteins. Further work comparing cluster-bound and cluster-free forms of the Isu scaffold and the interacting proteins is needed to verify the models discussed above.

Acknowledgments—We thank Annalisa Pastore (King’s College London) for providing PDB files of the bacterial FeS assembly complex and Roland Lill (Philips-Universität Marburg, Marburg, Germany) and Patrick D’Silva (Indian Institute of Science, Bangalore, India) for sharing plasmids.

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