Human Mitochondrial Chaperone (mtHSP70) and Cysteine Desulfurase (NFS1) Bind Preferentially to the Disordered Conformation, Whereas Co-chaperone (HSC20) Binds to the Structured Conformation of the Iron-Sulfur Cluster Scaffold Protein (ISCU)*

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Background: Iron-sulfur cluster biosynthesis involves a scaffold protein (ISCU), cysteine desulfurase (NFS1), chaperone (mtHSP70), and co-chaperone (HSC20). Results: Human mitochondrial ISCU populates structured (S) and disordered (D) conformational states. S interacts preferentially with NFS1 and mtHSP70; D interacts preferentially with HSC20. Conclusion: Shifts in the \( S \leftrightarrow D \) equilibrium reveal functional states.

Significance: The scaffold protein metamorphic property seen in *Escherichia coli* is conserved in humans.

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Fe-S clusters are ancient protein prosthetic groups that participate in a wide variety of biological processes, including electron transfer, substrate binding and activation, redox catalysis, DNA replication and repair, regulation of gene expression, and tRNA modification (1–3). In bacteria, three Fe-S cluster biogenic systems have been discovered: the NIF (nitrogen fixation) system involved in maturation of nitrogenase, the SUF (sulfur mobilization) system encoded by the *suf* operon and highly active under oxidative stress conditions, and the ISC2 (iron sulfur cluster) system, the general Fe-S cluster biosynthetic pathway (for review, see Refs. 4, 5, and 7–10). Among the three systems, the ISC system is the best-studied and is believed to be the “housekeeping” biosynthetic system (5). The *isc* operon of *Escherichia coli* codes for several proteins: a repressor (IscR), a cysteine desulfurase (IscS), a scaffold protein (IscU), a protein proposed to be an alternative scaffold (IscA), a DnaJ-type co-chaperone (HscB), a Dnak-type chaperone (HscA), a ferredoxin (Fdx), and a protein of uncertain function (IscX). By catalyzing the conversion of L-cysteine to L-alanine, the pyridoxal-5’-phosphate-dependent enzyme IscS generates \( S^0 \), which is transferred to Cys-328 to form a persulfide and then transferred to IscU (11, 12). Iron is added to form a [2Fe-2S] cluster. IscU-[2Fe-2S] then binds to HscB, which targets it to the HscA-ATP complex. In a reaction involving hydrolysis of ATP, the cluster is transferred to an acceptor protein such as apoferrredoxin (13, 14). We have shown by NMR studies that *E. coli* IscU populates two interconvertible conformational states: a more structured state (S) and a partially disordered state (D) (15). The two states play different roles in the cycle of Fe-S cluster assembly and transfer. The D-state is the substrate

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2 The abbreviations used are: ISC, iron-sulfur cluster; ISCU, human scaffold protein; IscS, iron-sulfur cluster; IscU, human mitochondrial DnaJ-type co-chaperone; HscA, *E. coli* Dnak-type chaperone; HscB, *E. coli* Dnak-type co-chaperone; HSC20, human mitochondrial DnaK-type co-chaperone; NFS1, human mitochondrial cysteine desulfurase; FMLR, fast metal affinity chromatography; TROSY, transverse relaxation optimized spectroscopy; IMAC, immobilized metal affinity chromatography.
for IscS (16); the S-state is the form that binds a [2Fe-2S] cluster (17) and binds preferentially to HscB (15, 18). Upon hydrolysis of ATP, HscA binds to the D-state of IscU, ensuing complete release of the cluster to an acceptor protein.

ISCU-type proteins are highly conserved throughout living systems (Fig. 1A). Mitochondria contain an ISC-type Fe-S cluster assembly and transfer system, parts of which are homologous to the ISC system of prokaryotes (6). In humans it has been proposed that this system involves a scaffold protein (ISCU) (19), a cysteine desulfurase consisting of two subunits, NFS1 (homologous to IscS) and ISD11 (no bacterial homologue found), a DnaJ-type co-chaperone (HSC20), and a DnaK-like chaperone (mtHSP70) (6, 19). These mitochondrial proteins are synthesized in the cytoplasm with N-terminal extensions that facilitate mitochondrial entry and are cleaved off upon protein maturation in mitochondria (20). The mature form of

**FIGURE 1.** Alignment of sequences of IscU homologues and structural representation of ISCU. A, alignment of sequences of IscU homologues. Analysis (15) of a much larger set of aligned sequences than those represented here showed that the residues highlighted in black were identically conserved, those in blue were conserved, and those in yellow were semi-conserved. The conserved cysteine residues are marked with red arrows, and the conserved LPPVK motif recognized by chaperone proteins is boxed in magenta. Mitochondrial proteins contain an N-terminal sequence that targets ISCU to cross the inner mitochondrial membrane. Excluding this region (boxed in red), human ISCU and *E. coli* IscU share 77% sequence identity. The numbering systems for mature ISCU and *E. coli* IscU are identical. Abbreviations used are: *Av*, *A. vinelandii*; *Ec*, *Escherichia coli*; *Hi*, *H. influenzae*; *Aa*, *A. aeolicus*; *Hs*, *Homo sapiens*; *Mm*, *M. musculus*; *Sc*, *S. cerevisiae*. B, solution structure of Zn$^{2+}$/H11001 bound *M. musculus* ISCU (PDB code 1WFZ) (47), which shares ~98% sequence identity with human ISCU, provides insights into the structure of human ISCU. Similar to other IscU homologues, *M. musculus* ISCU consists of three $\beta$-strands and four $\alpha$-helices. Residues mutated in this study (Asp-39, Asn-90, and His-105) are shown in red stick format. Asp-39 and His-105 are close to the Zn$^{2+}$/H11001 binding site, and Asn-90 is a solvent-exposed polar residue located in a hydrophobic region. All three residues are highly conserved among IscU homologues, and amino acid substitutions at these sites in *E. coli* IscU have been shown to perturb the position of the D $\rightleftharpoons$ S conformational equilibrium (35). Another important residue Trp-76 is shown in red stick format.
human ISCU, which shares 77% sequence identity with \textit{E. coli} IscU (Fig. 1A), has been shown to play an important role in cellular iron homeostasis (21). A tissue-specific splicing mutation of human ISCU has been associated with the disease ISCU myopathy (22). It has been proposed that the small protein ISD11 stabilizes the cysteine desulfurase NFS1 (23) and is important for Fe-S cluster assembly and cellular iron homeostasis (24, 25). Although it has been shown in an \textit{in vitro} assay that the ISCU-NFS1-ISD11 complex can assemble Fe-S clusters (26), little is known about the interactions among these three proteins. Human mtHSP70 (also termed Mortalin, PBP74, GRP75, or HSPA9) appears to be a multifunctional DnaK-type chaperone (27) (unlike \textit{E. coli} HscA, which is involved only in Fe-S cluster biogenesis). Human mtHSP70 is known to perform other cellular functions, including protein folding, intracellular trafficking, antigen processing, and aging (28, 29). It has been shown that mtHSP70 binds a variety of substrates, including cancer suppressor protein p53 and Parkinson disease-related protein DJ-1, and mtHSP70 has been associated with Alzheimer disease (30, 31). Because mtHSP70 is the only Hsp70 chaperone present in human mitochondria, it has been suggested that it plays a role in the human ISC machinery similar to that of \textit{E. coli} HscA (32). This hypothesis is supported by the finding that the mtHSP70 homolog in \textit{Saccharomyces cerevisiae} (Ssc1) participates in Fe-S cluster biosynthesis when the specialized Hsp70 chaperone (Ssq1) is knocked out (33). HSC20 recently has also been found to be involved in human mitochondrial Fe-S cluster biogenesis (34).

Given the sequence similarities of the bacterial and human proteins, it was of interest to determine, whether the human scaffold protein (ISCU) shares the conformational duality of the \textit{E. coli} ortholog (IscU) and, if so, whether the two states exhibit differential interaction with the cysteine desulfurase, co-chaperone, and chaperone proteins. We report here that the human scaffold protein for Fe-S cluster biogenesis (ISCU), like its \textit{E. coli} counterpart (IscU), populates two conformational states, a more structured state (S) and a partially disordered state (D). However, the relative population of the S-state of human ISCU is lower (28%) than the corresponding population of \textit{E. coli} IscU (80%) under comparable solution conditions. Human cysteine desulfurase (NFS1) alone and in the NFS1-ISD11 complex were tested in vitro for the S-state of human ISCU (Fig. 1A, red box), and the clones coded for a SUMO fusion containing an N-terminal His-tag. The genes were cloned into the pE-SUMO-Kan vector (Lifesensors; Malvern, PA) by using BsaI and XhoI restriction sites in the PCR gene fragments and vector. DNA primers used in these experiments were ordered through either the University of Wisconsin-Madison Biotechnology Center or Integrated DNA Technologies, Inc. (Corvallis, IA). Restriction enzymes were purchased from either Promega (Madison, WI) or New England Biolabs (Ipswich, MA). All PCR DNA primers used for cloning included nine additional base pairs at the 5’ ends upstream of the restriction sites to make digestion more efficient at the termini of PCR products. DNA ligation and construction of the expression plasmid were carried out in 10-\textmu l reactions with a PCR-based ligation thermo cycling program (30 s at 30 °C and 30 s at 10 °C repeated 800 times for 12 h). The ligation reaction was heat-inactivated at 65 °C for 25 min and then used to transform chemically competent 10G cells (Lucigen; Madison, WI), which were plated onto YT plates (containing 50 μg/ml kanamycin) and incubated at 37 °C overnight. An \textit{E. coli} recombinant colony archive was constructed by picking individual colonies and storing them in 30 μl of 20% sterile glycerol. 3-μl aliquots were removed for use in a PCR colony screen (20-μl reaction using Promega 2× PCR master mix) to identify positive clones. The original PCR primers used for isolating the target genes were employed in this PCR colony screen, and the reactions were analyzed on 1% agarose gels in TAE buffer (40 mM Tris acetate, 1 mM EDTA, pH 8.0). To prepare plasmids for DNA sequence analysis, the \textit{E. coli} colony glycerol stocks that yielded positive recombinants according to the PCR colony screens were grown overnight at 37 °C in 2–3 ml of CircleGrow® broth (MP Biomedicals; Santa Ana, CA) in the presence of 50 μg/ml kanamycin. All DNA sequencing reactions were carried out on a Bio-Rad Dye Pelletier Thermal Cycler at the University of Wisconsin-Madison Biotechnology Center, and SeqMan software (DNASTAR; Madison, WI) was used to analyze and identify targets with the correct DNA sequence.

We chose sites for production of single amino acid variants of ISCU to match those known to affect the D ⇄ S conformational equilibrium of \textit{E. coli} IscU (35). The positions of these residues on \textit{Mus musculus} Zn²⁺ bound ISCU (a model for the S-state) are shown in Fig. 1B (PDB code 1WFZ). \textit{M. musculus} ISCU shares 98% sequence identity with human ISCU (Fig. 1A). Genes for these variants were produced by using the Polymerase Incomplete Primer Extension (PIPE) site-directed mutagenesis method (36).

Production of Proteins—Single colonies containing validated genes for the target proteins (human ISCU variants, NFS1, ISD11, mtHSP70, and HSC20 and \textit{E. coli} IscS) were picked from the YT or MDAG plates (37) and grown in 1 ml of CircleGrow® broth or YT with 1% glucose (plus appropriate antibiotics) for 1–3 h at 37 °C at 250 rpm and then transferred to 50–100 ml of MDAG medium (with appropriate antibiotics) and grown overnight at 25 °C. For large scale protein production, 1 liter of unlabelled Terrific Broth (TB) auto-inducing medium or M9 isotopic medium was prepared, and 500-ml aliquots were transferred into sterile PET soda bottles (38, 39). Each 500-ml aliquot was inoculated with 10 ml of the overnight MDAG
starter culture, and the cell cultures were grown at 37 °C (250 – 320 rpm) for 2–5 h before dropping the growth temperature to 10–25 °C for 24–36 h. For induction with isopropyl-1-thio-β-D-galactoside (IPTG), the cell cultures were grown to A600 of 1.0–1.5; then the temperature was dropped to 10–25 °C, and IPTG (0.1–0.2 mM final concentration) was added when the temperature had stabilized (after about 15–30 min). For stable isotope labeling, we used an M9-based medium consisting of 100 ml/liter 10× M9 salts (70 g/liter Na2HPO4, 30 g/liter KH2PO4, and 5 g/liter NaCl), 1 ml of 1000× metal mix, 1 ml of vitamin mixture (40), 30 mg/liter thiamine, 0.5 ml of 0.2 M CaCl2 (0.1 mM final concentration), 2–5 drops of sterile antifoam, 2 ml of 1M MgSO4 (2 mM final concentration), [U-13C]glucose (2–4 g/liter), [15N]H4Cl (1 g/liter), plus the appropriate antibiotics (chloramphenicol to 35 g/ml and kanamycin to 50–100 g/ml). At the end of cell growth, the cultures were harvested by centrifugation for 30 min at 4000 × g in a centrifuge with a JS-4.0 rotor (Beckman Coulter; Brea, CA). The cell pastes were stored at −80 °C until needed for protein purification.

Buffers—The composition of the 1st immobilized metal affinity chromatography (IMAC) buffer was 20 mM Tris-HCl, pH 8, 300–500 mM NaCl, 0.1% Nonidet P-40, 1–2 mM β-mercaptoethanol or DTT, 1 mM phenylmethanesulfonyl fluoride (PMSF), 5–10% glycerol, and 5 mM imidazole. The composition of the 2nd IMAC buffer was the same as the 1st IMAC buffer except that it contained 250 mM imidazole. The SUMO-fusion cleavage buffer contained 20 mM Tris buffer at pH 8, 150 mM NaCl, 2 mM DTT (or β-mercaptoethanol), and 5–10% glycerol. The TND buffer consisted of 50 mM Tris-HCl, pH 8, containing 150 mM NaCl, 5 mM DTT, and 0.3% NaN3. The TKDM buffer consisted of 50 mM Tris-HCl, pH 7.5, containing 150 mM KCl, 5 mM DTT, and 10 mM MgCl2.

Protein Purification—All E. coli cell pastes were quickly thawed either on ice or at room temperature and then resuspended in 60–70 ml of lysis buffer: 1st IMAC buffer supplemented with Benzonase (Novagen, Millipore; Billerica, MA) or OmniCleavenuclease (Epicenter, Illumina; Madison, WI), tRNase (Novagen), RNase (Qiagen; Valencia, CA), and 0.1% Nonidet P-40 (Sigma). To break open the resuspended cells, we used sonication with a total time of 15–30 min at 4 °C, with a duty cycle of 2 s on and 4 s off. Cell lysates were clarified by high-speed centrifugation at 25,000 rpm for 30 min in a centrifuge with a JA 30.5Ti rotor (Beckman Coulter). The clarified protein solution was loaded onto a Qiagen Superflow FF or Ni-Sepharose column (GE Healthcare) IMAC resin at 1–5 ml/min. The IMAC column was washed first with ~10 column volumes of 1st IMAC buffer and second with 5–10 column volumes of wash buffer (1st IMAC buffer + 30 mM imidazole). The target protein was eluted with the 2nd IMAC buffer, and fractions were collected. SDS-PAGE was used to analyze and assess the purity of the eluted target protein in the collected fractions.

**FIGURE 2.** NMR evidence that ISCU exists in solution as two slow interchanging conformational states and that the S ⇄ D equilibrium is perturbed by single amino acid substitutions. A, 1H,15N HSQC NMR spectrum of ISCU. Because ISCU contains only one tryptophan (Trp-76), the presence of two cross-peaks in the boxed region and inset indicates the existence of two different conformational states. Assignments of individual peaks to the S- and D-states are indicated. B, 1H,15N HSQC NMR spectra of ISCU variants with shifted S ⇄ D equilibria. Whereas the substitutions D39V and N90A stabilize the S-state, substitutions D39A and H105A stabilize the D-state. C, expansions of the spectra in B show the Trp-76 side chain signals used to quantify the relative populations of the S- and D-states. All NMR spectra were collected at 600 MHz (1H) at 25 °C with solutions at pH 8.0.
The His-tagged, N-terminal SUMO fusion protein was digested with 0.5 mg of SUMO protease. The reaction was carried out in SUMO-fusion cleavage buffer either by desalting the fusion protein by size exclusion chromatography and adding SUMO protease or, more usually, by dialyzing the fusion protein in the presence of SUMO protease overnight at 4 °C against the cleavage buffer. The cleaved sample was loaded onto a freshly equilibrated subtractive IMAC column, which bound the cleaved His-tagged SUMO domain and allowed the cleaved target protein to pass through to a fraction collector. The purities of the target protein fractions were assessed by SDS-PAGE.

**NMR Spectroscopy**—For NMR samples the TND and TKDM buffers were modified to contain 10% D2O for the frequency lock. All NMR spectra were collected on 600 MHz (1H) Bruker BioSpin (Billerica, MA) NMR spectrometers equipped with a z-gradient cryogenic probe. All sample temperatures were regulated at 25 °C. NMRPipe software (41) was used to process the raw NMR data, and SPARKY software (42, 43) was utilized to visualize and analyze the processed NMR data.

\[ %S = \frac{[S]}{[S] + [D]} \times 100 \quad \text{(Eq. 1)} \]

We carried out three or more independent FMLR analyses of each spectrum to determine reproducibility and estimate errors (shown as error bars).

**Circular Dichroism Spectroscopy**—The sample buffer used in circular dichroism (CD) experiments contained 20 mM NaH2PO4 and 50 mM NaCl at pH 8. The solutions were placed in 1-mm path length quartz cuvettes. The concentration of ISCU variants was 20 μM. Far-UV CD spectra of ISCU variants were collected with an Aviv 202SF CD spectrophotometer (Aviv Biomedical; Lakewood, NJ) at 25 °C. Secondary structure content was estimated from the CD spectra by using K2D2 software (45).

**Size Exclusion Chromatography (Gel Filtration)**—Analytical gel-filtration studies were conducted with Hi-Load 75 Column (GE Healthcare) at room temperature. To investigate the interaction between HSC20 and ISCU (WT or N90A), a 2:1 (molar ratio) mixture of HSC20:ISCU in TKDM buffer was injected. The protein sample was eluted at 1 ml/min flow rate with TKDM buffer as the elution buffer, and 2 ml fractions were collected using an automatic fraction collector (GE Healthcare). Eluted fractions were analyzed by SDS-PAGE.

**ATPase Assays**—ATPase assays were carried out in TKDM buffer containing 0.1 mM ATP. The ATPase activity of mTHSP70 was determined at 25 °C by using an EnzCheck Phosphate Assay kit (Invitrogen) to measure the rate of phosphate release rate as described previously (46).

In Vitro Fe-S Cluster Assembly—A published protocol (47) was used to assemble Fe-S clusters in vitro. All samples were prepared in an anaerobic chamber (Coy Laboratory; Farmingdale, NY) filled with 90% N2 gas and 10% H2 gas. The reconstitution mix in TND buffer at pH 7.5 consisted of 50 μM ISCU or ISCU(N90A), 250 μM Fe(NH4)2(SO4)2, and 1 μM cysteine desulfurase (NFS1, NFS1-ISD11, or E. coli IscS). The reaction

### Table 1

| Structural property | WT | D39V | N90A | D39A | H105A |
|---------------------|----|------|------|------|------|
| %S (determined by NMR) | 27.5 | 100 | 95.4 | 29.1 | 11.2 |
| Predicted α-helix (%) | 0.742 | 0.926 | 1.01 | 25.5 | 25.0 |
| Predicted β-strand (%) | 16.2 | 25.5 | 25.0 | 16.2 | 25.5 |
| %S = ([S]/([S] + [D])) \times 100 (Eq. 1) |

We were able to assign NMR signals of ISCU(N90A), a variant that fully populates the S-state. We collected three-dimensional CBCACONH and HNCACB spectra from a sample of ISCU(N90A) labeled uniformly with 13C and 15N and used the data to carry out sequential backbone assignments.
was initiated by adding 250 μM l-cysteine into the reconstitution mix to make the final volume equal 1 ml. The reaction was carried out at 25 °C in a 10-mm path length quartz cuvette sealed with a rubber septum. Spectra were collected on a UV-1700 UV-visible spectrophotometer (Shimadzu; Kyoto, Japan) equipped with a temperature control utility. UVProbe 2.21 software (Shimadzu) was used to collect and analyze the data.

RESULTS

ISCU Populates Two Conformational States—Evidence for the structural heterogeneity of ISCU came from $^1$H,$^{15}$N HSQC NMR spectra that exhibited two sets of peaks for certain residues. The most prominent of these was the doubled $^1$H,$^{15}$N cross-peak from the side chain of Trp-76, the only tryptophan residue in the protein (boxed signals on Fig. 2A). The spectral analysis was clarified by comparison of $^1$H,$^{15}$N HSQC spectra of ISCU (Fig. 2A) with those of the four single-site mutants (Fig. 2B). Variants ISCU(D39V) and ISCU(N90A) yielded $^1$H,$^{15}$N HSQC spectra with sharp, well dispersed peaks, as expected for a well structured protein, whereas variants ISCU(D39A) and
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**TABLE 2**
Effect of added mtHSP70 on the %S of ISCU variants

| ISCU variant                   | Mole fraction of added mtHSP70 | %S (%)   |
|-------------------------------|--------------------------------|----------|
| ISCU                          | 0                              | 27.5     |
| ISCU(N90A)                    | 1                              | 95.4     |
| ISCU(H105A)                   | 1                              | 12.4     |

ISCU(H105A) yielded $^1$H,$^15$N HSQC spectra with broader, poorly dispersed peaks (particularly in the $^1$H dimension), as expected for a partially disordered protein. The spectrum of wild-type ISCU exhibited both sets of peaks. We thus assigned the sharper set of peaks to the structured state (S) and the broader set of peaks to the partially disordered state (D). Comparison of the Trp-76 peaks from ISCU (Fig. 2A) with those from the variants (Fig. 2C) allowed us to assign signals to the individual states. The %S values for the ISCU variants studied here are collected in Table 1.

We used CD spectroscopy to investigate the secondary structure of the ISCU variants. Far-UV (200–260 nm) CD spectra of ISCU variants that stabilize the S-state as shown by NMR exhibited secondary structure, whereas variants that stabilize the D-state as shown by NMR yielded CD spectra that could not be interpreted in terms of secondary structure (Fig. 3; Table 1). The CD spectrum of wild-type ISCU was consistent with the expected conformation of ISCU variants. Far-UV (200–260 nm) CD spectra of the structured variants of ISCU were found to be perturbed in the presence of 0.2 eq of unlabeled mtHSP70 (Fig. 5).

We hypothesized that the interaction between mtHSP70 and ISCU(N90A), which populates mainly the S-state, would be weaker than its interaction with ISCU. As anticipated, FMLR analysis of $^1$H,$^15$N TROSY-HSQC NMR spectra (Fig. 5, D and E) showed a much smaller effect: %S decreased from 95% in the absence of mtHSP70 to 83% at equimolar mtHSP70:U.$^1$N-ISCU(N90A) (Fig. 5F). The magnitudes of the chemical shift perturbations upon adding mtHSP70 were smaller for U.$^1$N-ISCU(N90A) than for U.$^1$N-ISCU (Fig. 6B).

We also investigated the interaction between mtHSP70 and ISCU(H105A), a variant that favors the D-state. The %S of U.$^1$N-ISCU(H105A) decreased from 12 to 0.54% upon the addition of equimolar mtHSP70 (Fig. 5, G–I; Table 2). The results again confirm that mtHSP70 preferentially binds to the D-state of ISCU.

**HSC20 Preferentially Binds to the S-state of ISCU**—Analytical gel-filtration chromatography was employed to investigate the interaction between ISCU and HSC20. Upon elution of a 2:1 (molar ratio) HSC20:ISCU mixture, a peak emerged at ~70 ml. This elution volume corresponds to the expected molecular mass of the HSC20-ISCU complex (~37 kDa) (Fig. 7A). SDS-PAGE of the elution fractions confirmed that the peak eluting at 70 ml contained both ISCU and HSC20 (Fig. 7B). We further followed two-dimensional $^15$N TROSY-HSQC NMR spectra of U.$^1$N-ISCU upon titration with unlabeled HSC20 (Fig. 8, A and B). FMLR analysis showed that %S increased from ~22 to ~31% upon the addition of equimolar HSC20 (Fig. 8C). The results indicate that HSC20 binds preferentially to the S-state of ISCU, in analogy to the finding that *E. coli* HscB binds preferentially to the S-state of IscU (15). By transferring the backbone assignments for ISCU(N90A) to the $^1$H,$^15$N HSQC spectrum from the S-state of ISCU, we were able to follow chemical shift perturbations $\Delta \delta_{HN}$ (as given by Equation 2) of U.$^1$N-ISCU upon titration with unlabeled HSC20 (Fig. 8, G and I),

$$\Delta \delta_{HN} = [(\Delta \delta_H)^2 + (\Delta \delta_N/6)^2]^{1/2}$$

where $\Delta \delta_H$ and $\Delta \delta_N$ are the chemical shift changes in the $^1$H and $^15$N dimensions, respectively.
Examples of the several peaks from U-^{15}N-ISCU that exhibited chemical shift perturbation upon binding HSC20 are shown in Fig. 9A. The residues of ISCU showing the largest chemical shift perturbations (Leu-31 and Val-32) (Fig. 8G) correspond to hydrophobic residues on the first $\beta$-strand of the three-dimensional structure of $M$. musculus Zn$^{2+}$-ISCU (PDB code 1WFZ) (Fig. 8I). We speculate that the first $\beta$-strand of ISCU provides the binding interface for the ISCU-HSC20 interaction.

We also investigated the interaction between HSC20 and the S-state favoring ISCU variant ISCU(N90A). Similar to ISCU, analytical gel-filtration results showed the elution of a peak corresponding to the expected mass of the HSC20-ISCU(N90A) complex (~37 kDa) (Fig. 7C). SDS-PAGE of the eluted fractions confirmed that this peak contained both ISCU and HSC20 (Fig. 7D). Two-dimensional $^{15}$N TROSY-HSQC NMR spectra of U-^{15}N-ISCU(N90A) were acquired as a function of added unlabeled HSC20 (Fig. 8, D and E). Several peaks from U-^{15}N-ISCU(N90A) exhibited chemical shift perturbations upon binding HSC20, and examples of these are shown in Fig. 9B. %S increased from ~94 to ~99% after the addition of 1 eq of HSC20, confirming that HSC20 binds preferentially to the S-state of ISCU(N90A) (Fig. 8F). Although more peaks of ISCU(N90A) were broadened beyond detection upon binding HSC20 as indicative of a tighter complex, the pattern of chemical shift perturbations (Fig. 8H) was similar to that for ISCU, suggesting that the first $\beta$-strand of ISCU(N90A) interacts with HSC20 (Fig. 8F).
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FIGURE 8. Interaction between HSC20 and ISCU. A, two-dimensional $^1H$/$^15N$ TROSY-HSQC spectra of U-$^15N$-ISCU (left panel), U-$^15N$-ISCU in the presence of equimolar unlabeled HSC20 and diluted by a factor of 2.25 (middle panel), overlay of the NMR spectra from the left and middle panels (right panel). B, expansions of the Trp-76 $^1H$/$^15N$ peaks from the spectra in A. C, %S calculated from FMLR analysis of the intensities of the Trp-76 cross-peaks assigned S and D under the conditions indicated. D, $^1H$/$^15N$ TROSY-HSQC spectra of U-$^15N$-ISCU(N90A) (left panel), U-$^15N$-ISCU(N90A) in the presence of 1 eq of unlabeled HSC20 and diluted by a factor of 2.25 (middle panel), overlay of the NMR spectra from the left panel and middle panels (right panel). E, expansions of the Trp-76 $^1H$/$^15N$ peaks from the spectra in D. F, %S calculated from FMLR analysis of the intensities of the Trp-76 cross-peaks assigned S and D under the conditions indicated. G, chemical shift perturbation of ISCU signals ($\Delta$\textsubscript{\textdelta}HN) upon the addition of 1.0 eq of HSC20 plotted as a function of ISCU residue number. Red triangles indicate residues whose NMR peaks were broadened beyond detection upon addition of HSC20. H, chemical shift perturbations ($\Delta$\textsubscript{\textdelta}HN) for residues of ISCU(N90A) upon the addition of 1 eq of HSC20. I, chemical shift perturbations of ISCU signals resulting from HSC20 binding mapped onto the structure of Zn$^{2+}$ bound M. musculus ISCU (PDB code 1WFZ) (47). Residues with $\Delta$\textsubscript{\textdelta}HN > 0.04 ppm are colored blue; residues whose NMR peaks were broadened beyond detection are colored red. J, chemical shift perturbations of ISCU(N90A) resulting from HSC20 binding mapped on the structure of Zn$^{2+}$ bound M. musculus ISCU with color coding as in I.
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In Vitro Fe-S Cluster Assembly Assay—To investigate the physiological importance of these findings, we carried out in vitro Fe-S cluster assembly assays. The UV spectra of the assembly mixture collected as a function of time showed the growth of peaks at 456 and 400 nm, which are characteristic for [2Fe-2S] and [4Fe-4S] clusters, respectively (47). This result indicates that ISCU serves as the scaffold protein for both types of Fe-S cluster (Fig. 10A). We found that NFS1 alone could catalyze Fe-S cluster assembly on ISCU and that the addition of ISD11 increased the Fe-S cluster assembly rate (Fig. 10B). We also investigated in vitro Fe-S cluster assembly on ISCU catalyzed by E. coli cysteine desulfurase (IscS) and found that the assembly rate was faster than that catalyzed by an equivalent concentration of NFS1-ISD11 (Fig. 10C). To investigate the effect of the conformational states of ISCU on cluster assembly, we repeated the reaction replacing ISCU by ISCU(N90A), a variant that is primarily in the S-state. Compared with wild-type ISCU, cluster assembly on ISCU(N90A) occurred at a much slower rate (Fig. 10D). The addition of Zn²⁺, which stabilizes the S-state (data not shown), inhibited the rate of cluster assembly on ISCU (Fig. 10E).

ISCU and HSC20 Stimulate the ATPase Activity of mtHSP70—We found that mtHSP70 exhibited a basal ATPase activity of ~0.10 ± 0.023 min⁻¹ (Fig. 11A, black) which is lower than that reported for E. coli HscA (~0.46 min⁻¹) (49). The addition of either 6 μM HSC20 or 15 μM ISCU to 1 μM mtHSP70 increased the basal ATPase activity of mtHSP70 by factors of ~1.7 and ~4.5, respectively (Fig. 11A, red and blue). The addition of both 15 μM ISCU and 4 μM HSC20 increased the basal ATPase activity of mtHSP70 by a factor of ~15 (Fig. 11A, green). Furthermore, we measured the effects of increasing concentrations of HSC20 alone (Fig. 11B), ISCU alone, and HSC20 + ISCU (Fig. 11C) on the ATPase activity of 1 μM mtHSP70. Based on the double-reciprocal plot, ISCU alone elicited a maximal stimulation of ~5-fold, and half-maximal stimulation occurred at ~1.5 μM ISCU. In the presence of 5 μM HSC20, a maximal stimulation of ~17-fold was observed upon adding ISCU, and the concentration of ISCU required for half-maximal stimulation was ~2 μM (Fig. 11D). In the presence of 5 μM HSC20, the addition of 24 μM ISCU(N90A) increased the basal ATPase activity of mtHSP70 only by a factor of ~7 (Fig. 11E). Based on the double-reciprocal plot, the maximal stimulation was ~8-fold, and the concentration of ISCU(N90A) required for half-maximal stimulation was ~3.5 μM (Fig. 11F).

DISCUSSION

The best studied ISC system for Fe-S cluster biosynthesis is from bacteria. An NMR structure has been determined for the S-state of E. coli IscU (50), and x-ray and NMR structures have been determined for Zn²⁺ complexes of Haemophilus influenzae IscU, Streptococcus pyogenes IscU (51), and Bacillus subtilis IscU (PDB code 1XJS, 10.2210/pdb1xjs/pdb). X-ray structures have been determined for the Aquifex aeolicus IscU-[2Fe-2S] complex (17), for E. coli IscS (12), for the E. coli IscS-IscU complex (53), for E. coli HscB (54), for the substrate binding domain of E. coli HscA complexed with the IscU recognition peptide ELPPVKIH (55), and for E. coli IscA (56). In addition, NMR studies of the E. coli ISC system have elucidated the roles of the S and D conformational states of IscU in the cycle of Fe-S cluster assembly and delivery (15, 16, 18, 50, 57). Despite sequence similarities of the homologous human mitochondrial proteins, the only three-dimensional structure determined to date of a protein from the human ISC system is that of HSC20 (human HscB) (58).

Results presented here show strong parallels between the conformational properties of the human (ISCU) and E. coli (IscU) scaffold proteins as well as their functional properties. Both human ISCU and E. coli IscU can adopt two very different folded conformations; one more structured (S-state) and one partially disordered (D-state) (Fig. 2A). Thus, the scaffold protein can be categorized as a metamorphic protein (59). In both cases, the S ⇔ D equilibrium is affected by single site amino acid substitutions at positions 39, 90, and 105 (16) (Fig. 2, B and C). Human ISCU is much less structured (~28% S) than E. coli IscU (~80% S) under similar buffer, pH, and temperature conditions.
Another interesting difference is that the D39A substitution, which stabilizes the S-state of E. coli IscU (16, 50) and has been found to stabilize Fe-S clusters in Azotobacter vinelandii IscU and Schizosaccharomyces pombe Isu1 (60), was found to favor the D-state of human ISCU. Because we had earlier seen a parallel between the S-state fraction and cluster stability in E. coli IscU (16), this result appears at odds with an earlier study, which reported that human ISCU(D39A) supported cluster formation, whereas ISCU did not (61).

Prior studies of the human cysteine desulfurase have focused on the NFS1-ISD11 complex, because of difficulties in isolating the two subunits (26, 62, 63). In this study we successfully expressed and purified the two proteins separately. We show here that isolated ISD11 does not interact directly with ISCU.

**FIGURE 10.** *In vitro* Fe-S cluster assembly assays. A, UV-visible absorption spectra of ISCU during Fe-S cluster assembly catalyzed by NFS1-ISD11. Spectra were collected at 60-min intervals. Absorption (ABS) at 400 nm and 456 nm are characteristic of [4Fe-4S] and [2Fe-2S] clusters, respectively. B, time course of Fe-S cluster assembly followed at 456 nm. Black line, reaction containing ISCU and catalyzed by NFS1 alone. Red line, reaction containing ISCU and catalyzed by NFS1-ISD11. C, time course of Fe-S cluster assembly followed at 456 nm. Black line, reaction containing ISCU and a catalytic amount of E. coli IscS. Red line, reaction containing ISCU and a catalytic amount of NFS1-ISD11. D, time course of Fe-S cluster assembly followed at 456 nm. Red line, reaction containing ISCU and a catalytic amount of NFS1-ISD11. Black line, reaction containing ISCU(N90A) and a catalytic amount of NFS1-ISD11. E, time course of Fe-S cluster assembly followed at 456 nm in the presence of a catalytic amount of NFS1-ISD11. Red line, reaction containing ISCU. Black line, Reaction containing ISCU and 1 eq of Zn**2**.
On the other hand, isolated NFS1 binds to the D-state of ISCU (Fig. 4A, middle panel) as does the NFS1-ISD11 complex (Fig. 4A, bottom panel). Human ISCU contains the conserved LPPVK motif (Fig. 1A) found in E. coli IscU, which is recognized by the substrate binding domain of E. coli HscA (64). We found that human mtHSP70 interacts with the D-state of ISCU (Fig. 5, A and B). This result is parallel to the interaction between E. coli HscA and the D-state of IscU (18). The addition of 1 molar eq of mtHSP70 shifted the $S \leftrightarrow D$ equilibrium of ISCU completely to the D-state (Fig. 5, A and B). By contrast, variant ISCU(N90A), which has a stabilized S-state, remained primarily in the S-state upon the addition of 1 molar eq of mtHSP70 (Fig. 5, C and D).

Human HSC20, the putative human homolog of the specialized DnaJ type co-chaperones, has been reported to be involved in human mitochondrial Fe-S cluster biogenesis and mitochondrial iron homeostasis (34). Although human HSC20 and E. coli HscB share high structural similarity, the former contains an extra N-terminal rubredoxin-like domain not present in E. coli HscB or S. cerevisiae Jac1 (58). We found that human HSC20 preferentially binds and stabilizes the S-state of ISCU (Fig. 8, A–F). This result is parallel to the preferential interaction between E. coli HscB and the S-state of IscU (15). Residues on the first $\beta$-strand of ISCU, namely Gly-30, Leu-31, and Val-32, exhibited large chemical shift perturbations upon the addition of HSC20 (Fig. 8, G–J). We speculate that these residues are

![Figure 11](image-url)
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FIGURE 12. Working model for human mitochondrial Fe-S cluster biogenesis. 1, apoISCU in D=5S equilibrium. 2, complex formed between the cytochrome desulfurase complex (NFS1-ISD11) and the D-state of ISCU. 3, sulfur delivered to Cys residues of ISCU. 4, addition of iron to form a [2Fe-2S] cluster stabilizes the S-state of ISCU. 5, the co-chaperone (HSC20) binds to holo-SCU displacing the cytochrome desulfurase complex. 6, the J-domain of HSC20 binds to the ATPase domain of the chaperone (mtHSP70), bringing holo-ISCU close to the chaperone. 7, an acceptor protein containing free Cys-SH groups approaches. 8, attack of cysteine residues from the acceptor protein liberates residues of ISCU that bind to the chaperone leading to activation of its ATPase activity. 9, conversion of ATP to ADP leads to a conformational change in the substrate binding domain of the chaperone, which then binds preferentially to the D-state of ISCU releasing the holo-acceptor protein and HSC20. 10, exchange of mtHSP70-bound ADP with ATP (which involves an exchange factor, not shown) leads to the release of ISCU, which resumes its equilibrium between the S- and D-states.

involved in a hydrophobic interaction between ISCU and HSC20.

The cluster assembly assay indicated that both [2Fe-2S] and [4Fe-4S] clusters could be assembled on ISCU (Fig. 10A) as catalyzed by NFS1 alone or by the NFS1-ISD11 complex (Fig. 10B). The rate with NFS1-ISD11 was ~27% faster than with NFS1 alone. We found that E. coli IscS also assembled clusters on human ISCU at an even faster rate than with NFS-ISD11 (Fig. 10C). This result is in agreement with a recent finding that the human NFS1-ISD11 complex, in the absence of frataxin or its bacterial homologue CyaY, exhibited lower cysteine desulfurase activity than E. coli IscS (62). As catalyzed by NFS-ISD11, ISCU(N90A), the variant with a stabilized S-state, assembled clusters 2.5 times more slowly than ISCU (Fig. 10D). Similar results have been reported for E. coli IscU variants with stabilized S-states (16). As with the E. coli system (16), Zn\(^{2+}\) can also stabilize the S-state of human ISCU (data not shown), and the addition of Zn\(^{2+}\) was found to inhibit cluster formation (Fig. 10E).

The ATPase assay showed that mtHSP70 has a basal ATPase activity of ~0.10 min\(^{-1}\) at 25 °C, which is close to that of S. cerevisiae Ssc1 and E. coli DnaK (~0.12 min\(^{-1}\)) (52) but much lower than that of E. coli HscA (0.46 min\(^{-1}\)) (49). The lower ATPase activity of mtHSP70 compared with E. coli HscA can be attributed to the fact that mtHSP70 requires a nucleotide exchange factor (GrpEL1), which catalyzes the exchange of ADP for ATP, to reach maximal ATPase activity (48). The E. coli HscA/HscB chaperone system does not utilize a nucleotide exchange factor (14). ISCU and HSC20 individually enhanced the ATPase activity of mtHSP70 severalfold, and HSC20 plus ISCU together increased the ATPase activity still more (Fig. 11A). The synergic effect of HSC20 and ISCU in stimulating mtHSP70 ATPase activity is similar to that reported for stimulation of the ATPase activity of E. coli HscA by HscB and IscU (46). Unlike the E. coli system, in which the presence of HscB decreased the concentration of IscU required to stimulate the ATPase activity of HscA (46), the concentration of ISCU needed for half-maximal stimulation of mtHSP70 ATPase activity was the same or higher in the presence of HSC20 (Fig. 11, C and D). We found that ISCU(N90A) (95 %S) in the presence of HSC20 was half as effective as wild-type ISCU (28 %S) in stimulating the ATPase activity of mtHSP70 (Fig. 11, E and F), in agreement with our model in which the D-state of ISCU binds preferentially to mtHSP70. Together, these results are consistent with the proposed function of human mtHSP70 and HSC20 as the chaperone and co-chaperone, respectively, for human mitochondrial Fe-S cluster biosynthesis.

The findings above support a working model for human mitochondrial Fe-S cluster biogenesis (Fig. 12) that is analogous to one proposed for the E. coli system (35). In this model, conversion of ISCU to the D-state when bound to the cytochrome desulfurase ensures that its Cys residues are free of metal (e.g. Zn\(^{2+}\)) and capable of accepting sulfur. Cluster formation then stabilizes the S-state of ISCU, weakens its interaction with the cytochrome desulfurase (NFS1-ISD11), and strengthens its interaction with the co-chaperone (HSC20), which binds preferen-
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