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Directing Noble Metal Ion Chemistry within a Designed Ferritin Protein*†‡

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ABSTRACT: Human H ferritin (HuHF) assembles from 24 four-helix bundles to form an ∼500 kDa protein with an 8 nm internal cavity. HuHF provides a useful model for studying the transport of metal ions in solution to buried reaction sites in proteins. In this study, HuHF was redesigned to facilitate noble metal ion (Au3+, Ag+) binding, reduction, and nanoparticle formation within the cavity. Computationally determined amino acid substitutions were targeted at four external and four internal surface sites. A variant with a total of 96 cysteines and histidines removed from the exterior surface and 96 non-native cysteines added to the interior surface retained wild-type stability and structure, as confirmed by X-ray crystallography, and promoted the formation of silver or gold nanoparticles within the protein cavity. Crystallographic studies with HuHF variants provide insight into how ferritins control access of metal ions to interior residues that perform chemistry.

Polypeptides play important roles in the biological synthesis of many minerals (e.g., silicates (1), carbonates (2), and phosphates (3)) and biologically derived particles of copper (4), iron (5), silver (6, 7), and gold (8). The molecular details of such syntheses, however, are not always well understood. Polypeptide-mediated biomineralization typically occurs under ambient temperatures and pressures and leads to structures (e.g., magnetic iron oxide particles and corals) that are ordered on both nano- and macroscale dimensions. Despite long-standing interest in biomineralization processes, few systems involving such nanoparticles and their peptides and metal ions have been structurally characterized in molecular detail (1, 9). For this reason, we have embarked on a program to investigate and reengineer ferritin proteins. We demonstrate in this work that sequence and structural changes to human H ferritin (HuHF),1 which were guided by computation and verified by X-ray crystallography, can yield a protein with new abilities to form gold and silver nanoparticles within its interior.

Noble metal nanoparticles have useful optical and electronic properties, as well as promising applications for biomedical (10–12), catalysis (13, 14), sensing (15), and nanoelectronics (16). While water-soluble silver and gold colloids have been described elsewhere using organic passivating agents (17–19), protein-based assemblies are bio-compatible and provide well-defined structures that can be modified site-specifically. HuHF is composed of 24 four-helix bundles that self-assemble to form a hollow spherical protein assembly, Mw ∼500 kDa, with roughly 8 nm inner and 12 nm outer diameters. Thus, ferritin provides a vehicle for directing the synthesis of inorganic materials within a preorganized protein framework.

Previously, horse spleen apoferritin (HSAF) was employed in the synthesis of metallic silver (20, 21), gold (22), palladium (23), copper (24), cobalt (25), and nickel (26) nanoparticles. In reactions of HSAF with Ag+ (20, 21), Cu2+, and a reducing agent, small particles were formed, indicating incomplete loading of the ferritin cavity. Some ions, such as Fe3+, gain entry to the interior cavity of ferritin through narrow, protein-spanning channels, whereas other metals are believed to interact preferentially with residues on the protein exterior and are largely excluded from the cavity (27). Limited evidence exists for mono- or trivalent metals diffusing through the 3-fold channel (26, 28, 29). In one example, the presence of Au+ and sulfide yielded an Au2S core within HSAF (29). Other examples involved reduction of Au3+ or Ag+ in the presence of HSAF, which produced particles that exceeded 8 nm in diameter (20, 22). These particles were clearly identified on the outer protein surface by TEM, with no evidence of metal deposition inside the cavity.

Dominguez-Vera et al. reported similar results in the synthesis of Ag0 particles with HSAF using NaBH4 as the reducing agent (21). UV–vis measurements showed that the silver surface plasmon resonance (SPR) absorbance band was red shifted by more than 20 nm from typical silver nanoparticles in solution, which may result from particle aggregation. Similarly, TEM images of the unstained Ag-HSAF

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‡ Abbreviations: BSA, bovine serum albumin; CD, circular dichroism; DLS, dynamic light scattering; FPLC, fast protein liquid chromatography; GuHCl, guanidine hydrochloride; HSAF, horse spleen apoferritin; HuHF, human H-ferritin; ICP-OES, inductively coupled plasma optical emission spectroscopy; IPTG, isopropyl β-D-galactopyranoside; MPD, 2-methyl-2,4-pentanediol; MOPS, 3-(N-morpholino)propanesulfonic acid; PDB, Protein Data Bank; RFZ, rotation function Z score; SASA, solvent-accessible surface areas; SCADS, statistical computationally assisted design strategy; SPR, surface plasmon resonance; TEM, transmission electron microscopy; TFZ, translation function Z score; Tm, melting temperature; wtH, wild-type human H-ferritin.
samples indicated close association of the 4 nm silver particles. These results suggest that metal nanoparticle synthesis occurs readily on the more accessible ferritin exterior surface.

The sequence and structures of wild-type proteins can be manipulated so as to control nanoparticle formation. One effort to bias the formation of nanoparticles within the interior of ferritin focused on developing a protein chimera (30). An exogenous polypeptide capable of reducing silver was appended to human L ferritin and displayed in the interior cavity, which promoted production of small silver particles. Such exogenous peptides are often unstructured, however, and may occlude interior volume in an uncontrolled manner. An alternate and potentially less disruptive approach is to redesign the sequence of a known ferritin, especially because these proteins can be robust with respect to a wide variety of mutations (31). In order to identify nonnatural ferritin nanoreactors with metal binding properties that are tuned to achieve new functions, we have employed computational design for the judicious choice of protein mutations.

Computational redesign of existing proteins allows the simultaneous variation of many residues and has previously yielded structures with novel properties (32, 33). Applications include the creation of maquettes (34) that bind porphyrins (35, 36) or metals (37) and proteins with catalytic activity (38). Many sequence search methods use pruning and dead-end elimination (39), Monte Carlo (40), and simulated annealing (33, 41) methods. Complementary to these approaches are probabilistic approaches to protein design (42, 43), which yield the site-specific probabilities of the amino acids among sequences likely to fold to a given three-dimensional structure (41, 44). With their capabilities for considering large numbers of simultaneous interresidue interactions and multiple substitutions, such computational methods are useful for identifying appropriate positions and mutations when engineering proteins. Based on their stability and high symmetry, ferritins are particularly amenable to redesign, as exemplified by a Dps protein that we engineered to contain a cavity with 86% apolar surface area (31). Using a similar approach, we have now modified up to 192 residues on the interior and exterior surfaces of HuHF, in order to control noble metal binding, reduction, particle nucleation, and growth within the protein template.

We hypothesized that removal of reactive cysteine and histidine residues on the exterior surface and introduction of additional soft cysteine ligands on the interior surface would promote the formation of gold and silver particles within the cavity. Precedent for this approach appears in work involving other protein assemblies. Within cowpea chlorotic mottle virus, changing native interior basic residues to acidic glutamates promoted the formation of an iron oxide core (45). Cysteine residues introduced into the interior of cow pea mosaic virus yielded a protein that maintained its structure and stability (46). Similarly, the addition of cysteine residues on the exterior of viral proteins proved to be useful for synthesizing gold clusters (47) and binding pre-formed particles (9). While these systems highlight the potential benefits of incorporating one or two external cysteine residues on the binding of premade gold particles or gold clusters, they were not designed for incorporation and nucleation of size-constrained noble metal particles. In the present study, the manipulation of histidine and sulfur ligands on both the HuHF exterior and interior surfaces was shown to improve solubility and have minimal effects on structure and stability, while creating a versatile biotemplate for the synthesis of metallic silver or gold nanoparticles. HuHF provides a useful model for studying how proteins are able to control the access of solution metal ions to residues at buried sites that perform chemistry.

**MATERIALS AND METHODS**

**Reagents.** Bactotryptone and yeast extract were purchased from Difco. IPTG was purchased from Fisher Biotech. Ampicillin sodium salt was purchased from Fisher Bioreagents. Molecular biology grade GuHCl was purchased from Promega. MOPS was purchased from Sigma. Gold(III) chloride was purchased from Acros Organics. 2,2′-Dipyridyl was purchased from Aldrich. All other chemicals were purchased from Fisher Scientific and were of ACS grade or higher, unless otherwise specified.

**Computational Determination of Mutations.** On the exterior HuHF surface, all four solvent-exposed cysteines and histidines, C90, C102, H13, and H105, were selected for mutation, in order to generate mutant H4. A second mutant was generated, H8, which included the four exterior mutations of H4 and four non-native cysteines on the interior surface.

The X-ray crystal structure used in the computational modeling was human H ferritin (PDB code 2FHA) with Gln-86 substituted for the wild-type Lys (48). Sites on the interior surface helices (B and D) were selected based on their SASA, as calculated with the software NACCESS using a probe radius of 1.4 Å (49). Residues with more than 50% solvent-accessible area were considered as possible mutation sites. Residues having favorable intra- or intersubunit interactions were discarded from the preliminary set: K49 and K53 were close to the 4-fold axis; K71 and K157 interacted with neighboring residues via interchain salt bridges, whereas N139 made favorable hydrogen bonds, H57 and H60 on the B-helix were also maintained as wild type, in order to maximize metal coordination within the cavity. The remaining five interior surface sites (E64, T135, E140, K143, and K157) were considered for possible substitution with cysteine. SCADS was employed to estimate the amino acid probabilities at each site considered for mutation (37, 42). To facilitate the calculation, a symmetry approximation was implemented, given the 24 identical subunits of HuHF, by imposing that every chain have the same sequence with equivalent residues having identical rotamer states (31, 44). Interatomic nonbonding energies were determined using the AMBER potential with a distance-dependent dielectric constant and a modified hydrogen-bonding term with an interatomic cutoff of 8 Å (42). No constraints involving solvation preferences (42) or secondary structure propensities were imposed. For the exterior surface design, all amino acids were allowed at sites 90 and 102 and all amino acids but Cys were considered at sites 13 and 105. For each of the five interior sites selected for mutation, only Cys and wild-type residues were allowed in each position.

**wtH, H4, H8, and H9x Gene Construction.** The wtH gene was generously provided by Dr. Paolo Arosio and cloned into a PET-17b vector (Novagen) via PCR by cloning in Ndel and HindIII restriction sites (at 5′ and 3′ ends, respectively). The PCR product was cleaned with the QIAquick PCR
purification kit (Qiagen), doubly digested with NdeI and HindIII restriction enzymes (New England Biolabs) at 37 °C for 1.5 h, and purified on a 1.2% agarose gel by the QIAquick gel extraction kit (Qiagen). The gene was ligated to NdeI/HindIII doubly digested pET-17b vector using the T4 DNA rapid ligation kit (Roche). The wtH cDNA served as the template for the H4 mutant. The following mutations were made in a stepwise fashion using the QuickChange site-directed mutagenesis kit (Stratagene): H13D, C90R, C102A, and H105Q. Primers used for mutagenesis of wtH to generate the H4 mutant are as follows: H13D (forward) 5′-GTGCGCCAGAATCAGCAGAGAGTCTCAGAG-3′, HindIII restriction enzymes (37 °C, 1.5 h). A 1.2% agarose gel was used to purify the gene, and a T4 DNA rapid ligation kit (Roche) was used to ligate the H8 gene with previously NdeI/HindIII doubly digested pET-17b vector.

Purified H8 cDNA served as the template for the H9x mutant, with two additional mutations, K86Q and C130S. These mutations were made in a stepwise fashion using the QuickChange site-directed mutagenesis kit (Stratagene). Primers used for mutagenesis of H8 to create the H9x mutant are as follows: K86Q (forward) 5′-CCTCTGAGCCACATGAGAAGTTTCTGC-3′, HindIII restriction enzymes (37 °C, 1.5 h). A 1.2% agarose gel was used to purify the gene, and a T4 DNA rapid ligation kit (Roche) was used to ligate the H8 gene with previously NdeI/HindIII doubly digested pET-17b vector.

**Protein Expression and Purification.** wtH, H4, H8, and H9x were transformed into *Escherichia coli* BL21-Codon Plus (DE3)-RP competent cells (Stratagene). Ten milliliter LB starter cultures (100 μg/mL ampicillin) were inoculated from single colonies and grown overnight at 37 °C, with shaking at 225 rpm. These cultures were then used to inoculate 1 L LB cultures (containing 200 μg/mL ampicillin) and grown until an OD600 of 0.6–0.8 was achieved. The cells were induced with 1 mM IPTG for 4 h. Cells were pelleted by centrifugation (5000 rpm, 20 min). The pellets were stored at −20 °C until use.

Purification was performed using a modified procedure by Levi et al. (50). Cell pellets were redissolved in 20 mM Tris and 0.15 M NaCl buffer at pH 7.4 (TBS), and a protease inhibitor cocktail tablet (Roche) was added. Cells were lysed on ice by sonication (3 × 2 min) and centrifuged (5000 rpm, 30 min). The lysate supernatant was heat shocked at 70 °C for 10 min and centrifuged (5000 rpm, 30 min). Ammonium sulfate (520 g/L) was added (57), stirred at 4 °C until all the ammonium sulfate dissolved, and centrifuged (5000 rpm, 30 min), and the pellet was redissolved in minimal TBS. The protein was run through a Superoxid 200 10/300 GL column multiple times to remove other protein and DNA. Protein purity of >95% was confirmed by FPLC, SDS−PAGE gel electrophoresis, and UV−vis spectroscopy. Iron was removed using 2,2′-dipyridyl following the procedure of Levi et al. (52). Overall yields of iron-free apoferritin were 2 mg/L of LB for wtH and H4 and 5 mg/L for H8 and H9x, as determined by Bradford assay. Only pure ferritin was used for the studies presented in this paper.

**Protein Characterization.** Ferritin samples were analyzed by UV−vis spectroscopy with an Agilent 8453 spectrophotometer with temperature controller and magnetic stirrer (Agilent 89090A), using a quartz cuvette with 1 cm path length. The molecular weights of the ferritin four-helix bundle subunits were determined at the Wistar Institute Proteomics Facility using an Applied Biosystems Voyager System 6030 MALDI-TOF mass spectrometer and by SDS−PAGE using a 15% denaturing gel with SDS. CD spectra were collected on an Aviv 62DS spectropolarimeter in the wavelength range 190−260 nm (far-UV) using a 1 mm quartz cell. Standard buffer is defined as 50 mM phosphate and 50 mM NaCl at pH 7.5.

Samples for thermal denaturation were prepared with 0.1 mg/mL protein in standard buffer. Ellipticity was measured at 222 nm by CD, and the temperature was increased from 4 to 96 °C in 2 °C steps. Samples were allowed to equilibrate for 1 min at each temperature. The melting process resulted in loss of signal as secondary structure was lost and the protein precipitated from solution. Melting temperatures were determined from the maximum of the first derivative plot of percent unfolded vs inverse temperature.

Samples for GuHCl denaturation were prepared with 0.1 mg/mL protein and concentrations of GuHCl varying between 0 and 7 M in standard buffer. The samples were mixed by vortex and allowed to stand at room temperature for 24 h, in order to achieve equilibrium at each guanidine concentration. The ratio of Trp fluorescence (λ316 nm/λ360 nm) was determined with a Varian Cary Eclipse fluorescence spectrophotometer operated with the Cary Eclipse Bio software package, upon excitation at 295 nm.

**DLS** was performed on a DynaPro instrument (Wyatt Technology Corp.), and the resulting data were analyzed with DYNAMICS Version 6.7. Samples were centrifuged at 15000 rpm for 10 min prior to measurement. A 5% intensity cutoff was applied to the data to compensate for fluctuations in signal. The polydispersity was less than 25% for all samples. Protein characterization data are summarized in Table 1.

**Table 1: Characterization of Wild-Type and Designed Ferritin Proteins**

<table>
<thead>
<tr>
<th>protein</th>
<th>Ms (calc)</th>
<th>Ms (exp)</th>
<th>DLS (radius, nm)</th>
<th>[GuHCl]1/2 (M)</th>
<th>Tm (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>wtH</td>
<td>21226</td>
<td>21336</td>
<td>6.7</td>
<td>4.0</td>
<td>83</td>
</tr>
<tr>
<td>H4a</td>
<td>21085</td>
<td>21066</td>
<td>6.3</td>
<td>6.2</td>
<td>90</td>
</tr>
<tr>
<td>H9b</td>
<td>20982</td>
<td>20982</td>
<td>6.4</td>
<td>4.7</td>
<td>80</td>
</tr>
</tbody>
</table>


**Metal Binding, Nanoparticle Synthesis, and Characterization.** Experiments were performed to determine the stoichiometry of Ag⁺ binding to 24-mer apoH4 and apoH8. Protein...
(0.2 mg/mL) in 50 mM MOPS buffer, pH 7.5, was titrated with aliquots of 10 equiv of Ag\(^+\) at 60 s intervals and monitored by UV–vis spectroscopy at 25 °C with stirring at 300 rpm. No protein loss was observed during this loading procedure. A difference spectrum (Ag\(^+\)–protein minus protein in solution) was obtained for each sample, in order to observe the thiolate-to-silver charge-transfer band. These crystals grew to typical dimensions of 0.2 mm × 0.2 mm × 0.1 mm within 2 days and exhibited diamond-like morphology that were suitable for data collection. These crystals grew to typical dimensions of 0.2 mm × 0.2 mm × 0.1 mm within 4 days. To prepare the Hg\(^{2+}\)–H8 complex, crystals of H8 were gradually transferred to 100 mM sodium acetate (pH 4.6), 20% (v/v) 2-propanol, and 20 mM thimerosal and soaked for 10 days. During this time crystals were washed twice with the same solution containing 20 mM thimerosal.

Crystals of H8 were cryoprotected by adding 2-methyl-2,4-pentanediol (MPD) to the mother liquor [final concentration = 45% (v/v)] and yielded diffraction data to 2.72 Å resolution at the Brookhaven National Laboratory (beamline X29A, λ = 1.0399 Å, 100 K). Data were processed with HKL-2000 (55) and scaled with SCALPACK (55) (Table 2). Diffraction intensities measured from these crystals exhibited symmetry consistent with Laue group P422 (unit cell parameters a = b = 171.3 Å, c = 190.5 Å). Analysis of reflection conditions suggested the space group P422 or P422_1. Crystals of Hg\(^{2+}\)–H8 were cryoprotected with a solution containing 45% (v/v) MPD and 0.1 M sodium acetate (pH 4.6) and yielded diffraction data to 3.06 Å resolution at the Brookhaven National Laboratory (beamline X29A, λ = 1.0399 Å, 100 K, unit cell parameters a = b = 171.2 Å, c = 190.4 Å). Due to the small size of the Hg\(^{2+}\)–H8 crystals, soaking experiments, and the high flux of the synchrotron beam, moderate crystal damage affected
data collection with an increased mosaicty of ~0.2°, which likely contributed to the relatively high $R_{merge}$ value of 0.144.

The Matthews coefficient of 2.76 Å$^3$/Da (solvent content = 55.5%) indicated that the asymmetric unit of H8 crystals contained 12 H8 monomers. The crystal structure of H8 was solved by molecular replacement with the program Phaser (56) using a hexamer of human ferritin as a search probe (PDB accession code 2FHA, less ions and water molecules) (48). The correct space group of H8 crystals was determined to be P422 based on the fact that only with this space group were very strong peaks found for the translation function with correspondingly high values of log-likelihood gain (LLG) (57). Using data in the 23–3.2 Å resolution range, the optimal solution for the positioning of two hexamers in the asymmetric unit yielded a total LLG of 6304, $RFZ = 14.8$ and $TZF = 49.9$ for the position of the first hexamer and $RFZ = 17.3$ and $TZF = 74.6$ for the second hexamer. The Eulerian angles for the orientations of the two hexamers were $\alpha = 135.2°$, $\beta = 44.9°$, $\gamma = 89.9°$ and $\alpha = 287.2°$, $\beta = 90.0°$, $\gamma = 90.0°$, respectively, and the corresponding translational searches yielded $x = 0.53$, $y = 0.23$, $z = -0.30$, and $x = -86.3$, $y = 0.32$, $z = 89.8$. The packing of the two hexamers in the asymmetric unit yielded a 24-mer assembly with 432 point group symmetry by the application of the symmetry operators for space group P422.

Iterative cycles of model building with the graphics program O (58) and refinement against 50.0–2.72 Å resolution data using torsion angle dynamics as implemented in CNS (59) improved the protein structure, as monitored by the values of $R$ and $R_{merge}$. A restrained noncrystallographic symmetry was used during all refinement processes with an appropriate weighting scheme (CNS) as judged by $R_{merge}$ and the residual Fourier map as refinement progressed. Group $B$-factors were utilized during refinement, and the majority of water molecules were automatically fit into the electron density peaks using a cutoff of 3.0σ, which improved the $R$ and $R_{merge}$ values to those reported in Table 2. A total of 368 water molecules were included in the refinement. Strong peaks aligned on the 3-fold axes located 2.2–2.9 Å from E131/E134 were modeled as Ca$^{2+}$ ions from the crystallization medium. Additional strong peaks aligned on the 4-fold axes located 3.1 Å from H173 of the terminal E-helices were tentatively identified as Zn$^{2+}$ ions based on inspection of the Bijvoet difference Fourier map calculated with anomalous data collected at the wavelength corresponding to the Zn edge. Disordered segments at the N- and C-termini (M1–S4 and D177–S182) were absent in the experimental electron density and are omitted from the final model.

The crystal structure of Hg$^{2+}$–H8 was solved by the difference Fourier method and refined as described for the structure of H8 with exception that in later cycles the refinement was completed with PHENIX (60). Strong peaks were observed in the simulated annealing map and identified as mercury ions. Disordered segments at the N- and C-termini (M1–S4 and D177–S182) were absent in the experimental electron density and are omitted from the final model. During refinement process we noted in the residual Fourier map a broad ~5σ peak adjacent to C64. This peak was ambiguous but may correspond to disordered thimerosal or MPD from the crystallization medium.

Crystal Structure Determination for H9x. Crystals of H9x were prepared in a similar manner as described for H8, and the Au$^{3+}$–H9x complex was prepared by soaking crystals in a solution of 0.5 mM AuCl3 of mother liquor with the absence of Ca$^{2+}$ for 1 week. X-ray diffraction data were collected from crystals of the Au$^{3+}$–H9x complex at the Cornell High Energy Synchrotron Source (CHESS) on beamline A1a and diffracted to 2.80 Å resolution. These crystals were isomorphous (space group F432, a = 181.6 Å) with the cubic crystal structure of human H chain ferritin (Table 2).

The crystal structure was solved by molecular replacement using the monomer from the H8 structure as a search probe for rotation and translation functions. The structure was refined as described above with the exception that in the latter cycles the refinement was completed with PHENIX (60). Strong peaks were observed in the simulated annealing map and identified as Au$^{3+}$ ions.

RESULTS

Computational Design. Exterior Cys and His residues were targeted for mutation, so as to minimize protein aggregation and nanoparticle nucleation at the protein outer surface. Cysteines and histidines are excellent ligands for noble metal ions. Furthermore, C90 and C102 were shown previously to promote ferritin aggregation through sulfur oxidation chemistry (61). Thus, calculations were performed to help identify optimal replacements for C90, C102, H13, and H105.

Probabilistic computational design methods were used to determine the likelihoods of the amino acids at these positions. The most probable residue at site 90 was Arg. The resulting model structure suggested that the guanidinium group of R90 can hydrogen bond with the backbone oxygen of D92 and form a salt bridge with E94. The most probable amino acids at site 102 were Ser and Ala. Alanine was selected due to its inertness toward metal binding, high α-helix propensity, and proximity to nearby hydrophobic residues (Trp93 and Met37). At position 13 on the protein exterior, Asp was the most probable residue, stabilized by a potential interhelical salt bridge with K124. For site 105, Glu, Lys, and Gln were preferred and had comparable probabilities. The wild-type crystal structure suggested that an ionizable residue at site 105 may disrupt a network of hydrogen-bonding and ionic interactions involving K109, Q7, R9, E101, and R156. The neutral H105Q mutation was selected to minimize interference with these wild-type interactions.

Calculations were also performed to identify interior sites where cysteines could be stably incorporated so as to promote metal ion binding and nanoparticle formation within the protein cavity. Site 135 exhibited a strong preference for the wild-type residue, particularly at low energies, and as a result Thr was retained at this position. At each of the four remaining sites (E64, E140, K143, E147), Cys had a probability comparable to the wild-type residues: E64, $P_{Cys}/P_{wt} = 2.0$; E140, $P_{Cys}/P_{wt} = 1.3$; K143, $P_{Cys}/P_{wt} = 0.5$; E147, $P_{Cys}/P_{wt} = 0.4$. Note that cysteines at sites 143 and 147 are less probable than wild type, suggesting that these mutations could destabilize the protein.

Biophysical Characterization. Upon expression and purification, the three proteins were studied for their assembly and stability. Dynamic light scattering (DLS) measurements indicated that H4 and H8 had virtually identical hydrody-
nomic radii to wtH. Each protein had a similar retention time by size exclusion chromatography (Figure S1, Supporting Information). Minimal 24-mer aggregation was observed for both H4 and H8 as shown by the disappearance of peaks corresponding to trimer (3 × 24-mer, elution at ~8 mL) and dimer (2 × 24-mer, elution at ~9 mL) with wtH. Multimerization of the 24-mer has been observed in other ferritins such as HSAF (62). H4 and H8 were more stable to guanidine denaturation than wtH (Table 1, Figure S2, Supporting Information), as monitored by tryptophan fluorescence that red shifted upon ferritin disassembly. Thermal denaturation experiments, in which loss of α-helical content at 222 nm was monitored by CD spectroscopy, determined the \( T_m \) (Figure S3, Supporting Information). The free energy of folding was calculated for wtH (~47 kJ·mol⁻¹), H4 (~53 kJ·mol⁻¹), and H8 (~52 kJ·mol⁻¹) as previously described (Figure 1) (31).

H8 X-ray Crystallography. The X-ray crystal structure of H8 (Figure 2, PDB code 2Z6M) confirmed proper assembly of the 24-mer holoprotein. Interestingly, a tetragonal crystal form resulted, which was the first time HuHF has been crystallized without the K86Q mutation. This crystal form was previously observed for horse spleen ferritin (but not for any human ferritin) with different unit cell parameters \((a = b = 147.2 \text{ Å}, c = 152.6 \text{ Å}; \text{PDB accession code IIES})\) (63). The H8 structure was refined to final \( R \) and \( R_{free} \) values of 0.215 and 0.252, respectively (Table 2). The root-mean-square deviation of 172 Cα atoms between H8 and wild type (PDB code 2FHA) (48) was 0.38 Å, indicating close structural homology. The cysteines in the cavity remained reduced under the conditions employed for crystallization. Electron density maps showing some of the engineered cysteine residues and the \( \text{Zn}^{2+} \) binding site are shown in Figure 3.

Silver and Gold Nanoparticle Synthesis. \( \text{Ag}^+ \) titrations were performed with 0.2 mg/mL H8 and monitored by UV–vis difference spectroscopy to determine the accessibility of cysteines within the protein interior. Absorbance at 291 nm may be assigned to a thiolate → Ag(I) charge-transfer band (Figure 4) (53). These data were consistent with H4 containing 1.0 ± 0.8 Cys per subunit (C130, located in the 3-fold symmetric pores) and H8 containing 4.3 ± 2.9 cysteines per subunit (four designed Cys mutations and C130). H4 serves as a useful control in these experiments, as the only difference between H4 and H8 is the incorporation of 96 cysteine residues within the H8 interior. These experiments indicate that cysteines introduced within the cavity are largely accessible to \( \text{Ag}^+ \) ions in solution.

Silver nanoparticles were synthesized by adding 1000 equiv of AgNO₃ to each apoferritin sample (0.2 mg/mL in 50 mM MOPS buffer, pH 7.5) and stirring at room temperature for 18 h while illuminating (60 W bulb, equivalent to 10 mW/cm² over the visible range, measured at the sample). All reactions with wtH, H4, and H8 turned yellow-brown within several hours. However, only the silver–H8 sample remained intensely yellow after centrifugation and elution through a gel filtration column and also exhibited a typical SPR band for silver nanoparticles, \( \lambda_{max} = 412 \text{ nm} \) (Figure 5). In MOPS buffer alone and in protein solutions that were not exposed to light, no color change was observed after 18 h, which confirmed that both the protein and light were required to form silver particles. The centrifugation and gel filtration purification steps ensured the removal of bulk silver.

In reactions of H8 with \( \text{Au}^{3+} \), reduction by MOPS buffer gave a characteristic gold SPR band at 534 nm (Figure 6). In the absence of MOPS, with only protein in solution to reduce the \( \text{Au}^{3+} \) ions, gold nanoparticles were observed to form on the time scale of weeks instead of minutes. The SPR max of the \( \text{Au–H8} \) sample was slightly red-shifted compared to \( \text{Au–wtH} \) and \( \text{Au–H4} \). While wtH and H8 appeared to stabilize \( \text{Au}^0 \) in solution, \( \text{Au–H4} \) appeared to...

**Figure 1:** Free energy of folding dependence of wtH, H4, and H8 as a function of GuHCl concentration calculated on a per subunit basis, \( n = 24 \). Linear extrapolation to 0 M GuHCl yielded \( \Delta G_{10,0} \) per mole of subunit in the absence of denaturant. Least-squares fitting was used to obtain the solid lines.

**Figure 2:** X-ray crystal structure (2.72 Å resolution) of a designed human H ferritin (H8) expanded to show a single H8 four-helix bundle subunit with mutations made to the exterior (red) and interior (yellow) surfaces.
aggregate based on the elevated baseline compared to the other two samples. Such aggregation with H4 was not unexpected and is likely the result of fewer Au0—surface interactions in H4 compared to wtH.

Loading of H8 with silver or gold was examined under various reaction conditions. The described procedures were performed dozens of times over many months for each protein—metal combination and gave very reproducible results. Based on Bradford assays using both BSA and ferritin standards, ~50% of H8 was retained in solution during metal particle synthesis. By comparison, 100% of the H8 protein was retained during iron loading. According to ICP-OES, Ag–H8 contained an average of 270 Ag atoms per holo-protein, which corresponded to uptake of 27% of available silver in solution under these reaction conditions. ICP-OES showed that H8 contained an average of 440 Au atoms per holo-protein, indicating that H8 was able to take up 22% of available gold under these reaction conditions. Based on TEM images for the Au–H8 samples, a small number of gold particles larger than the ferritin cavity were retained postpurification. Thus, the loading number obtained via ICP-OES for the gold samples should be considered an upper limit for these experiments. The 2–4 nm diameter cores of Ag0 and Au0 observed in many ferritin shells by TEM were consistent with spherical clusters of 300–2000 atoms.

Fe-Loaded H8. Iron-loading experiments with H8, monitored by UV−vis spectroscopy, showed ferric oxide mineral formation upon addition of 2000 equiv of Fe2+ (data not shown). ICP-OES performed on these samples indicated 1800 equiv of iron (>90% incorporation) per H8 24-mer, as expected for an iron-mineralized ferritin protein. TEM images of iron mineral–H8 (Fe–H8) showed a white outer protein shell and darker inner metallic core (Figure 7), which appeared very similar to the Ag–H8 and Au–H8 TEM images. Similarities in TEM data between Fe–H8, where the mineral occupies the cavity, and Ag–H8 and Au–H8...
provided strong evidence that mutations made to the HuHF exterior and interior surfaces promoted silver and gold nanoparticle formation inside the protein cavity.

$\text{Hg}^{2+} - \text{H8 X-ray Crystallography.}$ To examine why higher levels of noble metal loading were not observed in H8, Au$^{3+}$ soaks with protein crystals were attempted. Unfortunately, the presence of the gold ions caused crystal diffraction to appear mosaic, and Ag$^{+}$ was characteristically insoluble under protein crystallization conditions. Hg$^{2+}$ was then used instead to probe metal–thiol interactions. The Bijvoet difference Fourier map calculated with anomalous data collected at a wavelength corresponding to the Hg edge and the phases from the refined model less the atoms of Hg$^{2+}$, contoured at 3$\sigma$ (red). Mercury occupancies are estimated to be 50%.

$\text{Au}^{3+} - \text{H9x X-ray Crystallography.}$ Preliminary studies were performed to explore noble metal loading in the absence of C130, by making two mutations to H8 to create a new HuHF protein, H9x. The first mutation, K86Q, was done for previous HuHF crystal structures (PDB codes 1FHA and 2FHA) (48, 64), in order to improve ferritin intermolecular contacts, crystal growth, and diffraction. This exterior mutation should have little effect on noble metal binding. The second mutation was replacement of C130 with serine, as this residue should be less reactive with Au$^{3+}$ but is
Cys and/or His residues. Similar phenomena were seen in solution despite the added interior cysteine residues. In the crystal structure of H9x soaked with AuCl3, no gold ions were observed within the 3-fold symmetric channels, whereas gold ions were indeed observed at engineered cysteines in the H9x interior (Figure 9). This provides additional evidence that mutations made to surface-accessible residues (i.e., C130) can help to direct metal ions to the HuHF interior. Notably, however, in studies of gold and silver nanoparticle formation, UV−vis spectroscopy, TEM, and ICP-OES gave very similar data for H8 and H9x. Thus, additional factors controlling noble metal ion loading, reduction, nanoparticle formation and stabilization remain to be identified in this system.

**DISCUSSION**

**Biophysical Results.** Relative to wtH, the designed exterior mutations stabilized the protein (Figure 1, Table 1). The increase in H4 stability relative to wtH is most likely due to removal of the surface cysteines, which have been shown to cause ferritin aggregation (61), and introduction of stabilizing interactions involving surface exposed residues (C90R, H13D). With its 192 mutations guided by SCADS, it is remarkable that H8 exhibits nearly identical stability to H4 in solution despite the added interior cysteine residues.

**Silver and Gold Nanoparticle Synthesis.** The red-shifted Ag0 SPR band seen for wtH (λmax = 435 nm, with elevated baseline, Figure 5) is attributed to particle aggregation and partial solubilization of Ag0 via interactions with external Cys and/or His residues. Similar phenomena were seen previously in our laboratory with silver nanoparticle−HSAF assemblies (data not shown) and are similar to results observed with HSAF by Dominguez-Vera using NaBH4 as the reducing agent (21).

H4, which had the external Cys and His residues removed, serves as a useful control in these experiments, to allow direct comparison of the wild-type and cysteine-engineered protein interiors. H4 shows almost no SPR band, which indicates that H4 is unable to stabilize the silver particles that form in solution to the purification steps of centrifugation and gel filtration. In contrast, the purified Ag−H8 sample remains intensely yellow and exhibits a typical SPR band for silver nanoparticles, λmax = 412 nm. These data indicate that the cysteine-rich cavity is critical in forming stable silver nanoparticles at the protein interior. The removal of Cys and His residues from the protein exterior appears to favor reaction at these internal sites.

For the Au loading studies, clear differences are also observed between wtH, H4, and H8. H4 shows scattering (elevated baseline, Figure 6), which is indicative of poor particle solubilization, and results from the removal of exterior cysteine and histidine residues. wtH gives a characteristic SPR for gold, which indicates that the exterior wild-type surface (with particular emphasis on cysteine and histidine) stabilizes gold nanoparticles in solution as we saw previously for HSAF (22). However, the intensity of the Au SPR band is much greater for H8, providing evidence that a higher concentration of gold nanoparticles are stabilized in solution when grown within the protein interior.

**X-ray Crystallography.** As shown in Figure 8, the primary binding sites for the Hg2+ ions were the Cys130 residues at the interior opening of the 3-fold pore. While it is believed that C130 helps to facilitate metal ion transport into the protein, soft metal ions binding to these residues at the 3-fold pore, and subsequent reduction, could slow uptake and limit metal loading. Although there is no direct evidence for C130 nucleating gold and silver nanoparticle formation, reaction at this site has the potential to block the 3-fold channels and limit the size of the particles formed. Support for this hypothesis is provided by a crystal structure of H9x (Figure 9), in which C130 is replaced with serine, and Au ions soaked with the protein are observed at engineered cysteines in the H9x interior.

**Computational and Crystallographic Comparison.** In Figure 10, we compare the H8 structures determined via crystallographic refinement (Figure 2) with the calculated structures of the probable side chain conformational states. Each of the mutated sites is largely surface exposed, and a variety of populated side chain conformations may be expected. Nonetheless, the calculation yields local structure comparable to that observed in the experimentally determined structure, particularly with regard to C140 and C143 and the placement of the guanidinium group of R90. This comparison demonstrates the power of applying computation toward the design of ferritin proteins with new chemical reactivity.

In summary, variants of HuHF (H4 and H8) were designed with Cys and His residues removed from the outer surface. H8 also incorporated 96 non-native cysteines within the cavity, which was confirmed by X-ray crystallography. Both HuHF mutants formed very stable tetracosamers, but as designed, only H8 templated the synthesis of water-soluble, nonaggregating Ag0 and Au0 nanoparticles within the protein
interior, as indicated by UV-vis and TEM data. As expected, \( \textbf{H4} \) was relatively ineffective at stabilizing silver and gold nanoparticles in solution, based on the lack of available ligands on its exterior and interior surfaces.

This work illustrates how accessible protein surfaces may be engineered to control metal ion binding and subsequent chemistries. Ferritins support the wide-scale mutagenesis of residues important in metal surface chemistry, and it has become evident that ferritin, with eight 3-fold symmetric channels, can provide a useful model for studying the transport of metal ions within protein frameworks. Lessons learned from controlling the chemistry of noble metals in ferritin may be applied to other biomolecularly templated inorganic structures.

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SUPPORTING INFORMATION AVAILABLE

Figures S1–S3 providing FPLC traces, guanidine denaturation, and thermal denaturation curves for \( \text{wtH, H4, and H8} \). This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES

Noble Metal Ion Chemistry within a Designed Ferritin

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