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Formation of a Copper Specific Binding Site in Non-Native States of β -2-Microglobulin[†]

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ABSTRACT: A debilitating complication of long-term hemodialysis is the deposition of β -2-microglobulin $(\beta 2m)$ as amyloid plaques in the joint space. We have recently shown that Cu^{2+} can be a contributing, if not causal, factor at concentrations encountered during dialysis therapy. The basis for this effect is destabilization and incorporation of β 2m into amyloid fibers upon binding of Cu²⁺. In this work, we demonstrate that while β 2m binds Cu²⁺ specifically in the native state, it is binding of Cu²⁺ by nonnative states of β 2m which is responsible for destabilization. Mutagenesis of potential coordinating groups for Cu²⁺ shows that native state binding of Cu²⁺ is mediated by residues and structures that are different than those which bind in non-native states. An increased affinity for copper by non-native states compared to that of the native state gives rise to overall destabilization. Using mass spectrometry, NMR, and fluorescence techniques, we show that native state binding is localized to H31 and W60 and is highly specific for Cu^{2+} over Zn^{2+} and Ni^{2+} . Binding of Cu^{2+} in non-native states of $\beta 2m$ is mediated by residues H13, H51, and H84, but not H31. Although denatured β 2m has characteristics of a globally unfolded state, it nevertheless demonstrates the following strong specificity of binding: $Cu^{2+} > Zn^{2+} \gg Ni^{2+}$. This requires the existence of a well-defined structure in the unfolded state of this protein. As Cu²⁺ effects are reported in many other amyloidoses, e.g., PrP, α -synuclein, and A β , our results may be extended to the emerging field of divalent ion-associated amyloidosis.

The conversion of normally soluble proteins into amyloid fiber is a common feature of a number of clinical disorders, including Alzheimer's, type II diabetes, and spongiform encephalopathies (1). Furthermore, many proteins which do not form pathogenic fibers, e.g., myoglobin, have nevertheless been shown to convert to amyloid under laboratory conditions (2). Amyloid fibers formed from different proteins have many features in common. They are unbranched, resist proteolytic digestion, and display green birefringence upon staining with the histological dye Congo Red. The tertiary and quaternary structure of amyloid fibers has been determined from X-ray diffraction studies to be cross- β , with the β -strands arranged orthogonal to the fiber axis and hydrogen bonding parallel to the fiber axis (3). The kinetics of amyloid formation include a lag phase suggesting nucleation-dependent kinetics akin to crystallization (4). The similarity of fiber formation kinetics to crystallization includes the ability to bypass the lag phase by providing exogenous seed. Interestingly, while amyloids from different proteins share common histological and ultrastructural features, seeded reaction kinetics are dependent on using the seed of the same or

number of systems have been shown to display this property,

including mammalian prion (9), $A\beta$ from Alzheimer's (10),

α-synuclein from Parkinson's (11), immunoglobulin light

chains (12), and β 2m from dialysis-related amyloidosis (13).

closely related protein sequence. This has been used, for

example, to examine species barrier crossing in the yeast

prion system (5). Thus, despite apparent histological and ultrastructural similarities, amyloid fiber must have specific

atomic structures which mediate their assembly.

 β 2m is the 12 kDa polypeptide subunit that is necessary for the cell surface expression of the class I major histocompatibility complex (MHC)¹ (14). The turnover of MHC

MA 01701.

General determinants for amyloid formation have eluded identification; to date, only two structural features of polypeptides have been found to contribute. The first is a glutamine rich sequence such as that found in Huntingtin (6), and Sup35 from Saccharomyces cerevisiae prion (7). It is proposed that these sequences are stabilized by intersheet hydrogen bonding mediated by the glutamine side chains (8). The second is the presence of side chains which can interact with divalent metal, particularly Cu²⁺. An increasing

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¹ Abbreviations: β 2m, β -2-microglobulin; CD, circular dichroism spectroscopy; DRA, dialysis-related amyloidosis; ESI-MS, electrospray ionization mass spectrometry; MHC, major histocompatibility complex; NMR, nuclear magnetic resonance spectroscopy; NOESY, nuclear Overhauser effect spectroscopy; PAGE, polyacrylamide gel electrophoresis; SDS, sodium dodecyl sulfate; rWT, wild-type human β 2m from E. coli, possessing an additional N-terminal methionine; hWT, human-derived β 2m.

results in release of soluble β 2m followed by catabolism by the kidney. For patients suffering from kidney disease treated by dialysis, β 2m forms amyloid deposits principally in the joints, resulting in a variety of arthropathies. The circulating concentration of β 2m in dialysis-related amyloidosis (DRA) patients can be 5-50 times the normal level of 0.1 μ M. However, while elevated levels of protein are necessary for amyloid deposition, they are not sufficient. β 2m concentrations do not correlate with the pathology of DRA (15), nor does β 2m readily aggregate in vitro even at millimolar concentrations (pH 7) and salt concentrations isotonic with serum (16). We have recently shown that Cu^{2+} binds to $\beta 2m$ in a manner which is specific and at concentrations which can be encountered during dialysis therapy (13). Furthermore, the binding of Cu^{2+} by $\beta 2m$ results in destabilization and subsequent amyloid fiber formation. In this work, we have identified the residues responsible for binding and destabilization of the protein through the combined use of mutagenesis and analysis by native state mass spectrometry, NMR, and fluorescence.

MATERIALS AND METHODS

Chemicals. Buffers and salts obtained from either Sigma-Aldrich (Milwaukee, WI) or J. T. Baker (Phillipsburg, NJ). Cell culture media were purchased from Becton Dickinson (Sparks, MD).

Wild-Type and Mutant β2m Overexpression and Purification. β2m was expressed in Escherichia coli using plasmid pHN1, a gift from D. C. Wiley (Harvard University, Cambridge, MA). BL21(DE3) competent cells were transformed and grown in Luria broth in the presence of ampicillin (100 µg/mL). Cultures were grown at 37 °C to an OD at 600 nm of 0.6 and then induced with 500 μ M IPTG for 2 h. Cells were harvested by centrifugation and lysed by sonication. Inclusion bodies were extracted, washed six times with 1 M urea, 100 mM Tris, 100 mM EDTA, and 10 mM methionine (pH 8.0), and resuspended in 8 M urea, 100 mM Tris, 100 mM EDTA, and 10 mM methionine (pH 8.0) for 24 h at 4 °C. Insoluble material was removed by centrifugation, and the soluble protein was refolded by dialysis against 25 mM potassium phosphate, 150 mM KCl, 1 mM EDTA (pH 7.4), and 10 mM methionine at 4 °C. After dialysis, the protein was concentrated by centrifugation using Vivascience (5 kDa cutoff) and run over a gel filtration column with Sephedex-75 resin at 4 °C and pH 7.4. The final purity was assessed by SDS-PAGE and electrospray ionization mass spectrometry (ESI-MS). Site specific mutations of β 2m were made using Quickchange (Stratagene, La Jolla, CA) and sequenced (Keck facility, Yale University) to confirm the mutation. The mutant plasmids were purified as wild-type β 2m and confirmed by SDS-PAGE and ESI-MS. Humanderived β 2m was purified as previously described (13). Briefly, urine from Dent's disease patients (Yale School of Medicine Human Investigation Committee protocol 11522) was filtered (0.2 μ m) and applied directly to a Cu²⁺-charged NTA-Superflow column (Qiagen), washed (three times) with 5 mM KP (pH 7.4) and 150 mM KCl and five times with no salt and 1 mM imidazole. The protein was fractionated upon elution with 10 mM imidazole. β 2m-containing fractions were pooled, concentrated, and further purified by size exclusion as performed for E. coli-derived β 2m.

Determination of the Oxidation State. Oxidized and reduced wild-type and mutant β2m were analyzed by reverse-phase HPLC using a Vydac C18, 300 Å,4.6 mm × 250 mm column. Gradient separations were performed using an acetonitrile/water gradient (1%/min) in the presence of 0.1% trifluoroacetic acid, with a flow rate of 1 mL/min. Samples were denatured (6 M guanidinium chloride) in the presence or absence of freshly prepared dithiothreitol (\sim 1 mM) overnight prior to being applied to the column. The fraction of protein in the reduced state was determined from a comparison of the integrated peak area at elution times corresponding to fully oxidized or reduced β2m standards. The dynamic range of this approach permits us to confidently detect as little as 1% residual reduced protein in our samples.

Circular Dichroism (CD). CD experiments were performed on an Aviv model 215 spectrometer. Far-UV spectra (190–240 nm) were collected in a 1 mm path-length cell with 30 μ M β 2m in 25 mM potassium phosphate and 150 mM KCl (pH 7.4). Near-UV spectra (240–330 nm) of 100 μ M β 2m in the same buffer were recorded in a 3 mm path-length cuvette.

Fluorescence. Metal binding to the native state and the chemical stability of $\beta 2m$ were monitored by following changes in intrinsic fluorescence on a PTI Quantamaster C-61 with slit widths of 2 nm and excitation at 283 nm. In metal binding experiments, protein was diluted to 2.5 μ M with a buffer containing the stated metal salt, 25 mM MOPS, and 150 mM potassium acetate at (pH 7.4). Chemical stability measurements were taken under the same buffer conditions over a range of urea concentrations. For chemical denaturation, samples were allowed to equilibrate for at least 12 h prior to data acquisition. Results are expressed as the average emission wavelength, $\langle \lambda \rangle$, calculated from the emission between 300 and 450 nm (13, 17).

Mass Spectrometry. Spectra were acquired from 200 to $5000 \, m/z$ with signal averaging for 1 min using a Micromass LCT electrospray time-of-flight mass spectrometer. Atmospheric pressure ionization was performed using borosilicate glass capillaries drawn and sputter coated in-house. All mass determinations were made using a minimum of three charge states. External calibrations were performed with 20 mM CsI in $\rm H_2O$ ionized under matched instrument conditions. The temperature of the mass spectrometer ionization source was maintained at 20-25 °C. Protein ion counts of metal-bound and unbound forms of $\beta 2 \rm m$ were determined from the +7 charge state.

Calculations. Multiple-sequence alignment was performed using the mature human wild-type sequence as a query string to fasta33t (18) at www.ebi.ac.uk. Default parameters were used apart from limiting the upper expectation value to 10^{-10} which had the effect of restricting hits to alternative variants of β 2m. Consensus analysis was performed using MView version 1.41.8 (19). Molecular graphics and solvent exposure calculations were performed using MolMol (20, 34). The extent of solvent exposure was determined on residue side chains using a 1.4 Å probe radius. All curve fitting in this work was performed using the NonlinearRegress function in Mathematica 4.0 (Wolfram Research, Inc., Champaign, II.)

NMR. NMR spectra were recorded in house using a Varian Unity Plus instrument with proton resonance of 600 MHz. NOESY spectra were collected using a standard pulse

sequence with a mixing time of 200 ms. Two thousand points were collected in t_2 , and 256 increments were collected in t_1 . Data were processed using NMR PIPE (21) and analyzed using Sparky (22). The apo sample was 400 μ M β 2m, 0.4 mM d_{12} -EDTA, 10% D₂O, 150 mM KCl, and 25 mM potassium phosphate buffer at pH 7.0 and 37 °C. The holo sample was 500 μ M β 2m, 60 μ M CuCl₂, 10% D₂O, 150 mM KCl, and 25 mM phosphate buffer at pH 7.0 and 37 °C. TMSP was included in both samples for reference.

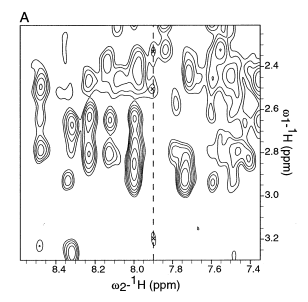
RESULTS

In this work, we seek to delineate the role of divalent metal in native state binding and destabilization of $\beta 2m$. The side chain ligand most likely to be involved in these interactions is histidine. Therefore, a series of four mutants have been prepared in which each histidine residue of $\beta 2m$ has been changed to a noncoordinating moiety. The affinity of a divalent ion for protein reflects both chemical and structural effects, the latter resulting from the three-dimensional orientation of the coordinating moieties of the protein. We have therefore made comparative measurements of $\beta 2m$ in the presence of Cu^{2+} , Zn^{2+} , and Ni^{2+} .

Native State Structure. The native state copper binding site was initially elucidated by a combination of ¹H NMR and intrinsic fluorescence. ${}^{1}H^{-1}H$ NOESY spectra of $\beta 2m$ acquired in the presence and absence of CuCl2 indicate that H31 is involved in binding. The basis for this observation by NMR is the paramagnetic broadening of resonances near the binding site. To avoid complications from both nonspecific paramagnetic broadening and protein aggregation at high concentrations of protein and copper (13), substoichiometric (1:8) amounts of Cu²⁺ were used. Paramagnetic broadening resulted in the diminishment of peaks previously assigned (16) to the side chain of H31. For example, crosspeaks involving the C_{ϵ} proton of H31 (Figure 1A) are absent in spectra in which Cu²⁺ is present (Figure 1B). This is similar to previously reported ¹H-¹H TOCSY observations (23) in which H31 and also H13 resonances are affected by the presence of Cu²⁺. Under our conditions, the cross-peaks of the C_{ϵ} proton are completely eliminated. As substoichiometric amounts of copper are being used, this indicates that chemical exchange of copper is rapid compared to the relaxation time.

The intrinsic fluorescence of $\beta 2m$ is principally derived from its two indole side chains, W95 and W60. The fluorescence emission of $\beta 2m$ in the presence of Cu²⁺ both decreases in intensity and shifts to shorter wavelengths (Figure 6 and ref 13). At saturating levels of Cu²⁺, the maximal decrease in intensity is $\sim 40\%$ and the blue shift is 6 nm. Since a buried indole is expected to emit at shorter wavelengths, these properties are consistent with a model in which a solvent accessible tryptophan is quenched by interaction with Cu²⁺. W60 is more solvent exposed than W95 (75 vs 9%, respectively) (24), implying W60 is proximal to the Cu²⁺ binding site. Indeed, the crystal structure of $\beta 2m$ reveals that W60 and H31 are only ~ 7 Å apart (Figure 2B), further reinforcing the observation that H31 is part of the native state binding site.

The specificity for Cu²⁺ binding by the native state (13) suggests a physiological role for Cu²⁺ binding. This is supported by its multiple-sequence alignment analysis show-



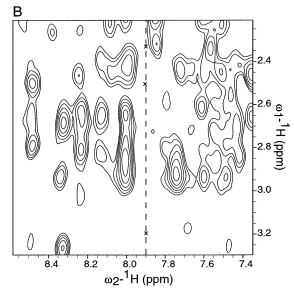
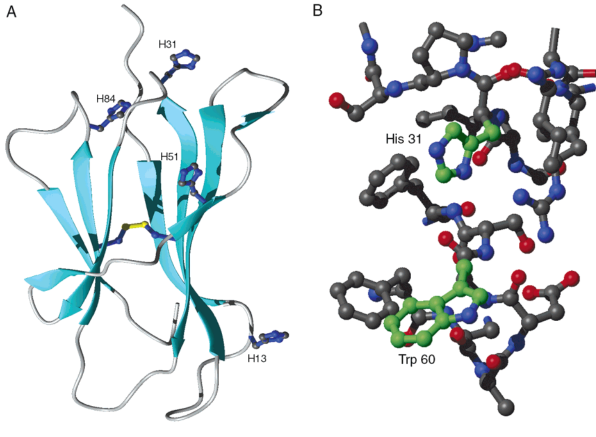


FIGURE 1: (A) $^{1}H^{-1}H$ NOESY spectrum of 400 μ M β 2m in the presence of 0.4 mM EDTA. (B) $^{1}H^{-1}H$ NOESY spectrum of 500 μ M β 2m in the presence of 60 μ M CuCl₂. The dashed line at $\omega_2 = 7.9$ ppm corresponds to H31 H ϵ (16). A total of nine peaks (of which three are shown) were present at this resonance in the EDTA spectrum; all of these were absent or significantly less intense in the presence of copper. Spectra shown are contoured at the same level renormalized to the average of six invariant crosspeaks in the aromatic region of the spectra.

ing conservation of H31 across 87 sequence variants (Figure 2C). Of the four histidine residues in human β 2m (Figure 2A), those at positions 31 and 84 are 100% conserved across species with as little as 37% total sequence identity. In contrast, those at positions 13 and 51 are more variable. This conservation correlates with the relative levels of solvent exposure of these residues; H13 and H51 are \sim 70% exposed (Table 1), while H31 and H84 are 38 and 0% exposed, respectively. Sequence conservation of H84 likely reflects a role in the packing of the protein core. In contrast, as residue H31 is not wholly buried, its conservation suggests a potential physiological role.

Mutant Protein Characterization. To identify the Cu^{2+} binding site and probe the role of Cu^{2+} binding and destabilization, each of the histidine residues in $\beta 2m$ (H13,



С									
Species	Identity	Sequence							
Human	100%	IQRTPKIQVYSRHPAENGKSNFLNCYVSGFHPSDIEVDLLKNGERIEKVEHSDLSFSKDWSFYLLYYTEFTPTEKDEYACRVNHVTLSQPKIVKWDRDM							
Orangutan	96%	IQRTPKIQVYTRHPAENGKSNFLNCYVSGFHPSDIEVDLLKNGVKIEKVEHSDLSFSKDWSFYLLYYTEFTPTEKDEYACRVNHVTLSQPKTVKWDRDM							
Mouse	70%	IQKTPQIQVYSRHPPENGKPNILNCYVTQFHPPHIEIQMLKNGKKIPKVEMSDMSFSKDWSFYILAHTEFTPTETDTYACRVKHDSMAEPKTVYWDRDM							
Chicken	47%	ADLTPKVQVYSRFPASAGTKNVLNCFAAGFHPPKISITLMKDGVPMEGAQYSDMSFNDDWTFQRLVHADFTPSSGSTYACKVEHETLKEPQVYKWDPEF							
Medaka Fish	37%	$\tt KTSPPTVQVYSRDPGQYGKENTLICHVSNFHPPDITIRLMEDDQELPEAKQTDLAFKQNWHFHLTKSAPFTPKQGARYTCKVTHSGLAKDYTWESNM$							
Consensus	90%	spnlovysp Pen Go Nhi, Cavot FHPenInlphi.Keg h ten oblumnewnFalbh st FTPent-nyeC+VnH h t.n.h W							

FIGURE 2: (A) Ribbon structure of β 2m (2CLR) with histidine and cysteine residues shown with a ball-and-stick rendering. (B) Ball-andstick rendering of atoms within 8.5 Å of the ϵ -carbon of H31. Carbon atoms of H31 and W60 are shown in green. (C) Summary of multiple-sequence analysis of 87 sequence variants of β 2m. A selected subset are shown along with the consensus analysis for all 87 sequences.

Table 1: Summary of Measured Free Energies of Unfolding and Dissociation Constants for Cu²⁺

	$\Delta G_{ m unfolding}^{ m apo}$ a	$m_{ m apo}{}^a$	$\Delta G_{ ext{unfolding}}^{ ext{holo}\ b}$	$m_{ m holo}{}^b$	$K_{ m d}$ c	$\Delta G_{ m u}^{{ m Cu}d}$	$\Delta\Delta G_{ m u}^{{ m Cu}\;e}$	exposure ^f
WT	33.0 ± 2.2	5.4 ± 0.4	12.0 ± 0.7	3.6 ± 0.2	2.8 ± 0.4	29.6	NA	NA
H13F	33.5 ± 3.4	5.0 ± 0.5	20.6 ± 0.7	3.7 ± 0.1	3.2 ± 0.4	21.2	8.4	70
H31F	43.9 ± 2.3	6.3 ± 0.3	17.0 ± 0.3	3.6 ± 0.2	41.0 ± 2.7	28.8	0.8	38
H51F	25.0 ± 0.3	4.0 ± 0.05	16.0 ± 0.9	3.4 ± 0.1	1.7 ± 0.3	18.9	10.7	67
H84A	22.0 ± 0.6	4.4 ± 0.1	14.0 ± 0.3	3.7 ± 0.06	1.5 ± 1.4	18.1	11.5	0

^a Free energy and m values of unfolding in the absence of added metal (Figure 4). ^b Free energy and m values of unfolding in the presence of 90 μ M CuCl₂ (Figure 4). ^c Dissociation constant for Cu²⁺ from β 2m (Figure 6). ^d The calculated free energy of copper dissociating from the unfolded state at 90 μ M Cu²⁺ ($\Delta G_{\rm f}^{\rm Cu}$) based on the model discussed in the text ($\Delta G_{\rm u}^{\rm Cu} = \Delta G_{\rm unfolding}^{\rm apo} + \Delta G_{\rm f}^{\rm Cu} - \Delta G_{\rm unfolding}^{\rm holo}$) is determined using $K_{\rm d}$ in column 3. ^e Relative contribution of each histidine to the calculated free energy of copper binding to the unfolded state ($\Delta \Delta G_{\rm u}^{\rm Cu} = WT\Delta G_{\rm u}^{\rm Cu} - \Delta G_{\rm unfolding}^{\rm Cu}$) mutant ΔG_u^{Cu}). Percent solvent exposure of the histidine side chain in 2CLR. Units of ΔG are kilojoules per mole. Units of m values are kilojoules per mole per molar. Units of K_d are micromolar.

H31, H51, and H84) was individually mutated to phenylalanine using standard methods. In the case of H84, mutation to Phe resulted in an unstructured protein (data not shown). This residue was therefore mutated to alanine. All constructs were purified by the same method used for the recombinant wild-type protein (rWT), producing H13F, H31F, H51F, and H84A β 2m mutants. The purity was assessed by SDS-PAGE and mass spectrometry. Only bands attributed to β 2m were visualized by SDS-PAGE, and masses determined by ESI-MS are within 0.1% of the calculated mass. The

oxidation state of the C25-C80 disulfide in β 2m was evaluated using a chromatographic approach (see Materials and Methods). Human-derived β 2m, (hWT), H13F, H51F, and H84A did not exhibit any detectable reduced protein (data not shown). Recombinant WT (rWT) and H31F each have \sim 9% reduced protein. For the stability and ligand binding measurements performed here, rWT and hWT are indistinguishable, suggesting that these small amounts of reduced material are not significant in measurements of global properties.

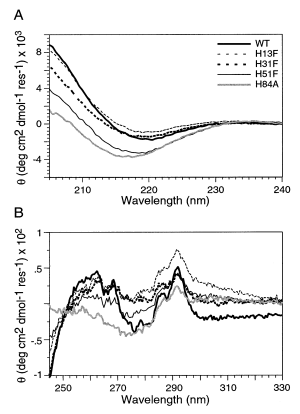


FIGURE 3: Far-UV (A) and near-UV (B) CD spectra of wild-type and mutant constructs of $\beta 2$ m. Near-identical profiles demonstrate that the global structures of wild-type and mutant $\beta 2$ m are identical. Small changes, particularly in the near-UV CD, are consistent with mutations occurring in aromatic regions of the polypeptide. WT spectra of the recombinant protein are shown.

All constructs used in this work are folded with secondary and tertiary structure consistent with the wild-type protein. Assessment of structure in rWT, H13F, H31F, H51F, and H84A β 2m was performed using far- and near-UV CD. All demonstrate weak negative ellipticity (Figure 3A) at 220 nm, as is characteristic of a β -sheet fold (Figure 2A). H51F and H84A, however, have somewhat more negative ellipticity at 220 nm than rWT. This may be attributed to the breaking of one or more hydrogen bonds upon mutation of the histidine, resulting in small changes to the secondary structure. In the near-UV CD spectrum, positive bands were observed at 270 and 290 nm (pH 7.4) for the wild type, and all mutations. These bands arise from the packing of aromatic groups into a well-defined and fixed conformation (Figure 3B). Individual mutations in β 2m vary only slightly compared to WT in near-UV CD, indicating that the mutants are both folded and have nearly identical atomic arrangements.

Wild-type and mutant constructs all show cooperative two-state transitions upon titration with urea (Figure 4), providing further evidence that the mutant proteins are folded. The stability (ΔG) and linear dependence of stability on urea concentration (m value) (25) of H13F are nearly identical to those of the wild type (Table 1). This is consistent with the fact that H13 is 70% solvent exposed, and does not participate in any intramolecular hydrogen bonding. Mutation of H84 and H51 results in destabilization of 11.0 and 8.0 kJ/mol, respectively, compared to WT. This is likely the result of lost hydrogen bonding and changes in solvent exposure. For example, the difference in buried surface area

upon mutating H84 to Ala is \sim 80 Å³, and two hydrogen bonds are lost (T86 and the backbone of S33) (24). In contrast, H31F displays a net stabilization of -10.9 kJ/mol. This suggests that the local structure around H31 is normally under strain.

Cu²⁺ Effects on Conformational Stability. The contribution of each histidine residue to Cu²⁺ induced destabilization was assessed by urea titration in the presence and absence of added Cu2+ (Figure 4). The most notable feature of this analysis is that destabilization by Cu²⁺ is enhanced rather than diminished in the H31F construct. Under near-physiological conditions (150 mM potassium acetate at pH 7.4 and 25 °C), WT β 2m is destabilized 21.0 kJ/mol by the addition of 90 μ M Cu²⁺ [Table 1 (33.0 – 12.0 kJ/mol) and Figure 4A,D]. In H31F, the protein is destabilized by 26.9kJ/ mol upon addition of Cu²⁺ (Figure 4C,D). All other constructs have a reduced sensitivity to Cu²⁺. H13F, H51F, and H84A are destabilized by 12.9, 9.0, and 8.0 kJ/mol, respectively. This reduced destabilization suggests that these three histidine residues are dominant in Cu²⁺ binding by nonnative states of β 2m. In contrast, the increased Cu²⁺ sensitivity of H31F compared to that of WT suggests that H31 interactions with Cu²⁺ are dominant in folded states of β 2m.

The specificity of Cu²⁺ induced destabilization was assessed by urea titration of wild-type β 2m in the presence of Ni²⁺ or Zn²⁺. At 90 μ M Ni²⁺ or Zn²⁺, β 2m is not destabilized (Figure 5A). Specificity was therefore determined using relatively high concentrations of Ni²⁺ and Zn²⁺ compared to the concentration of Cu^{2+} . At 20 μ M Cu^{2+} , WT β 2m is destabilized by 9.8 kJ/mol (Figure 5B). Addition of $400 \,\mu\text{M Ni}^{2+}$ to $\beta2\text{m}$ resulted in no detectable change to the urea titration profile (data not shown). At 2000 μ M Ni²⁺, the urea titration profile has shifted, giving a midpoint of 6.6 M urea compared to a midpoint of 6.1 M urea for no divalent ion. The free energy of unfolding, however, is within error of metal free β 2m (-30.6 kJ/mol). The analogous study with Zn²⁺ did yield statistically significant destabilization. Addition of 4000 μ M Zn²⁺ was required to destabilize β 2m by 14 kJ/mol. Clearly, non-native states of β 2m show a strong preference for the binding of Cu²⁺ over Zn²⁺ and Ni²⁺, suggesting well-defined positioning of the coordinating histidine side chains in these non-native states.

Native State Binding of Divalent Metal Ion. The binding constant of each mutant construct to Cu^{2+} was determined by changes in intrinsic fluorescence (13). H13F, H51F, and H84A have dissociation constants between 1 and 3 μ M (Figure 6 and Table 1) which are comparable to WT. H31F, however, binds $Cu^{2+} > 10$ -fold weaker than the wild type with a dissociation constant of $41.0 \pm 2.7 \,\mu$ M (Figure 6A,B). This clearly implicates H31 as the primary ligand in Cu^{2+} binding by the native conformation of β 2m. As binding of Cu^{2+} to the H31F construct is not eliminated, it is likely that other residues in the vicinity of H31 also participate in forming the binding pocket (Figure 2B).

The binding constant of WT β 2m to Ni²⁺ and Zn²⁺ was also determined by changes in intrinsic fluorescence. Ni²⁺ and Zn²⁺ bind much more weakly to β 2m than Cu²⁺ (Figure 6C,D). As Ni²⁺ is also a fluorescence quencher, its level of binding could be determined in the same manner as that of Cu²⁺. This yields a K_d of 400 μ M (Figure 6C), more than 100-fold weaker than the interaction of Cu²⁺ with β 2m. In



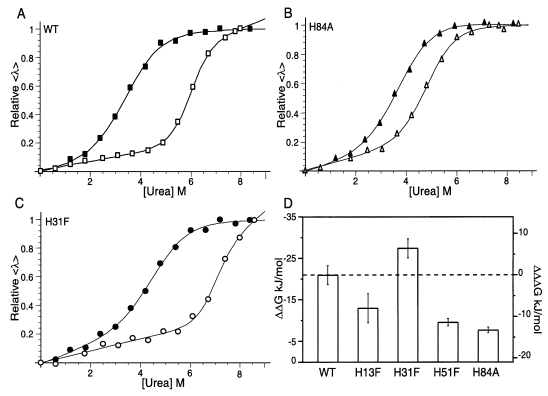


FIGURE 4: Chemical stability of β 2m and mutant variants measured by changes in intrinsic fluorescence. Urea denaturation of 2.5 μ M WT (A), H84A (B), and H31F (C) in the presence (filled symbols) and absence (empty symbols) of 90 μ M Cu²⁺. All denaturation data were fit to a two-state model (25). (D) Summary of these results for all mutants expressed on the left axis as ΔG of folding in the presence of $Cu^{2+} - \Delta G$ of folding in the absence of Cu^{2+} . To illustrate the relative increase in sensitivity of H31F to copper, and the relative decrease in sensitivity of all other mutants to copper, a labeled axis on the same scale is shown on the right but expressed relative to WT. A dotted line is drawn at the zero of the right-hand axis. Error bars are ± 1 standard error of the mean (SEM).

addition, the loss of fluorescence intensity upon Ni²⁺ binding is less than that observed for Cu²⁺ (only 20% compared to 40%). This disparity could be due to the different intrinsic quenching efficiencies of the two metals. Another factor, however, could be the geometry of coordinating moieties in the binding site. These could place Ni²⁺ in a different position with respect to W60, leading to a decreased level of quenching. Since Zn²⁺ is not a quencher, its binding constant to β 2m was determined by competition for the Cu²⁺ binding site by Zn²⁺ over a series of Cu²⁺ binding assays at seven different $\mathrm{Zn^{2+}}$ concentrations ranging from 0 to 6000 $\mu\mathrm{M}$ Zn^{2+} (Figure 6D). The binding constant of Zn^{2+} to $\beta 2m$ was determined by global analysis to be 1.5 mM, 500-fold above the binding constant for Cu²⁺. Clearly, the metal binding site of native β 2m is highly specific for Cu²⁺.

Metal Binding by Native State Mass Spectrometry. Direct assessment of metal binding to β 2m was made using native state electrospray ionization mass spectrometry (26). An electrospray mass spectrum of β 2m in the presence of Cu²⁺ contains a series of peaks corresponding to $\beta 2m \cdot (Cu^{2+})_k$ (for k = 0, 1, 2, ...). Measurement of dissociation constants by mass spectrometry is complicated by the absence of a 1:1 correspondence between the measured intensity and solution concentration. Furthermore, the necessary presence of weak volatile chelates (e.g., NH₃) complicates determination of the free divalent ion concentration. We have therefore combined hWT protein with expressed proteins to allow for internal control of these factors (Figure 7A). As the recombinant constructs all contain an additional N-terminal methionine, they have an increased mass (e.g., 131 Da for rWT-hWT) which permits their resolution from hWT in

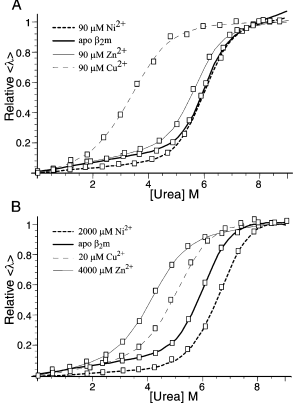


FIGURE 5: (A) Chemical stability of 2.5 μ M WT β 2m measured by changes in intrinsic fluorescence as a function of added urea in the absence or presence of 90 μ M Cu²⁺, Ni²⁺, or Zn²⁺. (B) Chemical stability in the absence or presence of 20 μ M Cu²⁺, 2000 μ M Ni²⁺, or 4000 μ M Zn²⁺.

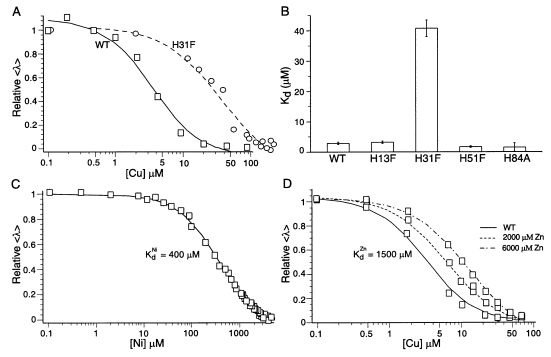


FIGURE 6: Measurement of the divalent ion affinity of mutant β 2m for the folded state by changes in intrinsic fluorescence. (A) Representative plots of copper titration of WT (\square) and H31F (\bigcirc) are shown with nonlinear fits of equilibrium dissociation with a 1:1 stoichiometry. (B) Summary of Cu²⁺ dissociation constants measured for each construct shown with error bars at ±1 SEM. (C) Assessment of Ni²⁺ binding by titration. A nonlinear fit is shown giving a K_d of 400 μ M. (D) Multiple copper titrations of β 2m at seven different concentrations of Zn²⁺ were performed. Three representative data sets with fits are shown. Global fit to all the data yield a K_d for Zn²⁺ of 1500 μ M.

heterogeneous mixtures. Comparisons of other closely related protein pairs suggest that it is reasonable to assume that instrument response will affect $hWT \cdot (Cu^{2+})_k$ and $rWT \cdot (Cu^{2+})_k$ equally for a given k within a given charge state in the same spectrum (27). Relative dissociation constants can therefore be accurately determined as

$$K_{d,r\beta2m}/K_{d,h\beta2m} = [r\beta2m \cdot (Cu^{2+})_0 \times h\beta2m \cdot (Cu^{2+})_1]/$$
$$[h\beta2m \cdot (Cu^{2+})_0 \times r\beta2m \cdot (Cu^{2+})_1] (1)$$

where each $r\beta 2m \cdot (Cu^{2+})_k$ is the peak intensity from a given recombinant $\beta 2m$, with $k Cu^{2+}$ ions bound.

The Cu²⁺ binding behavior of hWT and that of rWT are indistinguishable (Figure 7A,B). At 40 µM protein and 120 μM Cu²⁺, the dominant peak is $\beta 2m \cdot (Cu^{2+})_1$. A small percentage (13%) of protein can be seen binding a second Cu²⁺. In a control experiment using hen lysozyme under matched solution conditions, a single Cu²⁺ is seen, accounting for 11% of the protein (data not shown). Since hen lysozyme is not a Cu²⁺ binding protein, this suggests that the small population of $\beta 2m$ with two Cu^{2+} ions is a reflection of nonspecific adduct formation resulting from the ionization process. The peak corresponding to $\beta 2m \cdot (Cu^{2+})_1$ includes contributions from both specific binding and nonspecific adduct formation. The contributions of the latter may be estimated by looking to the peak corresponding to $\beta 2m \cdot (Cu^{2+})_2$. For rWT, this peak is 20% of the height of the peak at $\beta 2m^{\bullet}(Cu^{2+})_1$. The same relative proportion applied to $\beta 2m^{\bullet}(Cu^{2+})_0$ indicates that only 5% of the peak at $\beta 2m^{\bullet}(Cu^{2+})_1$ is due to nonspecific adduct formation. The relative K_d (eq 1) for rWT and hWT is 1.2, indicating that recombinant and human-derived protein are quantitatively as well as qualitatively identical.

Copper binding measurements of β 2m constructs were conducted by mixing hWT with H13F, H31F, H51F (MW = 11870 Da), or H84A (MW = 11794 Da). Only H31F reveals behavior clearly distinct from that of the wildtype protein (Figure 7D). A total of 90% of hWT is in Cu²⁺bound states (k = 1 or 2). In contrast, 30% of H31F shows addition of only one Cu2+ ion. This is only slightly larger than expected for nonspecific adduct formation. Supporting this is a Cu²⁺ titration of H31F in which we observe that binding of one Cu²⁺ by H31F parallels binding of a second Cu²⁺ by hWT (data not shown). Using eq 1, we can conclude that the relative K_d of H31F for copper is minimally 13-fold greater than that of hWT. A more accurate measurement cannot be made as this is at the limit of interference from nonspecific adduct formation. The H13F and H51F mutants each bind copper with qualitative behavior which is indistinguishable from that of the hWT protein (Figure 7C). The relative dissociation constants measured using eq 1 are 1.1 and 0.5, respectively. These compare well with the values of 1.1 and 0.6 calculated from the ratio of dissociation constants obtained by fluorescence (Figure 6B). The relative $K_{\rm d}$ of H84A could not readily be determined by this approach since hWT·(Cu²⁺)₂ and H84A·(Cu²⁺)₀ are partially overlapping. Overall, we conclude that direct assessment of Cu²⁺ binding agrees with our fluorescence studies, indicating that H31 is the primary location for native state Cu²⁺ binding (Figure 6).

Assessment of Ni²⁺ binding by fluorescence (Figure 6C) does not necessarily reflect binding to the Cu²⁺ binding site. Competition of Ni²⁺ for the Cu²⁺ binding site can, however, be directly observed by ESI-MS. The high concentrations of Ni²⁺ required for binding result in ESI-MS spectra which are obscured by nonspecific adduct formation intrinsic to

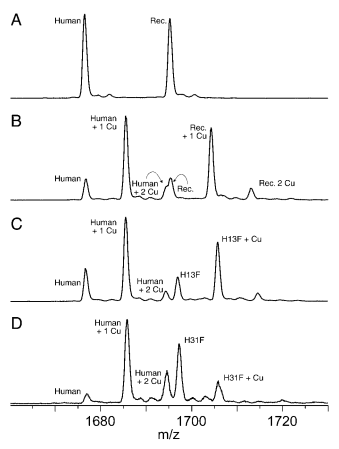


FIGURE 7: Mass spectra of folded $\beta 2m$ in the presence or absence of added divalent copper. To facilitate the comparison of mutant and wild-type forms of $\beta 2m$, equimolar mixtures of human-derived and recombinant $\beta 2m$ were prepared. The total protein concentration is 40 μ M. (A) Spectra of human and recombinant $\beta 2m$ in the absence of added metal. Only the +7 charge state is shown yielding two peaks separated by m/z 18.7. This corresponds to the additional methionine residue of recombinant $\beta 2m$. (B) Human and recombinant $\beta 2m$ in the presence of 120 μ M CuCl₂. (C) Human wild-type $\beta 2m$ and H13F in the presence of 120 μ M CuCl₂. (D) Human wild-type $\beta 2m$ and H31F in the presence of 120 μ M CuCl₂.

the electrospray process. Therefore, we have performed experiments using Ni²⁺ in the presence and absence of 60 μM Cu²⁺. Copper binding to $\beta 2m$ was competed at 60 μM Cu^{2+} with 60 μ M, 600 μ M, and 6 mM Ni²⁺. At a 1:1 Cu^{2+} : Ni²⁺ ratio, only Cu²⁺ binding is detected (Figure 8B). The $\beta 2$ m·(metal²⁺)₁ – $\beta 2$ m·(metal²⁺)₀ mass difference in Figure 8B is 61.3 Da which allows 94.6% of the β 2m·(metal²⁺)₁ peak to be attributed to Cu²⁺. At Ni²⁺ concentrations of 600 μM, the spectra become more complex due to nonspecific adduct formation (Figure 8C). Addition of 60 μ M Cu²⁺ is nevertheless able to cause a significant mass shift (Figure 8D). Mass measurement of $\beta 2m^{\bullet}(\text{metal}^{2+})_1 - \beta 2m^{\bullet}(\text{metal}^{2+})_0$ peaks in Figure 8D shows that 78% of the mass shift can be attributed to Cu²⁺. Increasing the concentration of Ni²⁺ to 6 mM gives spectra which are indistinguishable in the presence and absence of $60 \,\mu\mathrm{M}$ Cu²⁺ (Figure 8E,F). Thus, we observe competition of Ni²⁺ for the Cu²⁺ binding site at a 100-fold excess of Ni2+, a molar ratio comparable to the difference in the observed K_d , measured by changes in intrinsic fluorescence (Figure 6).

DISCUSSION

Elucidation of the residue specific effects of Cu^{2+} on $\beta 2m$ is central to understanding the mechanism of divalent induced

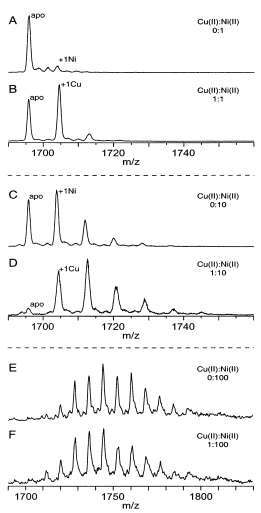


FIGURE 8: Mass spectra of $r\beta 2m$ in the presence of mixtures of Cu^{2+} and Ni^{2+} . The total protein concentration is 40 μ M. The Cu^{2+} concentration is 60 $\mu \dot{M}$ in panels B, D, and F. The Ni²⁺ concentration is 60 μ M in panels A and B, 600 μ M in panels C and D, and 6000 µM in panels E and F. Peaks corresponding to β 2m with one associated metal are annotated with the dominant metal contributing to the mass shift. Nickel-containing spectra are dominated by ladders of peaks which correspond to nonspecific association of Ni²⁺ which occurs during the ionization process. This is most notable in panels E and F. Solution binding of Ni²⁺ to the Cu²⁺ binding site is therefore inferred from the elimination of a mass shift associated with copper binding. The copper mass shift in the presence of Ni²⁺ is most clearly evident at a 1:1 Cu²⁺:Ni²⁺ stoichiometry (A and B). No copper mass shift is evident at a 1:100 Cu²⁺:Ni²⁺ stoichiometry (E and F), indicating that Ni²⁺ has displaced Cu²⁺ from the binding site.

fibrillogenesis in DRA. Here we have performed a range of experiments aimed at delineating between binding and destabilization. There are four critical observations which impact our understanding of this system. (i) The native state binding site for Cu^{2+} is restricted to the vicinity of H31 and W60. (ii) Native state binding of divalent ion is specific for Cu^{2+} over Zn^{2+} and Ni^{2+} . (iii) Cu^{2+} induced destabilization of $\beta 2m$ is mediated by residues which are not involved in native state binding. (iv) Destabilization of $\beta 2m$ by divalent ions is specific for Cu^{2+} over Zn^{2+} and Ni^{2+} .

Assertions of Cu²⁺ specificity when describing metal binding by proteins are complicated by the relatively high intrinsic affinity of the imidazole group for Cu²⁺ over other metals. Ni²⁺ and Zn²⁺ are commonly used for comparison with Cu²⁺ as a result of their identical charge, and nearly

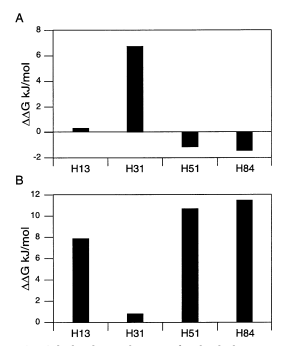


FIGURE 9: Calculated contributions of individual amino acids to the free energy of binding of $\mathrm{Cu^{2+}}$ to folded ($\Delta\Delta G_\mathrm{f}^\mathrm{Cu}=\mathrm{WT}\Delta G_\mathrm{f}^\mathrm{Cu}-\mathrm{mutant}\ \Delta G_\mathrm{f}^\mathrm{Cu}$) (A) and unfolded ($\Delta\Delta G_\mathrm{u}^\mathrm{Cu}$ from Table 1) (B) conformations of $\beta 2\mathrm{m}$. $\Delta G_\mathrm{f}^\mathrm{Cu}$ values are the free energies of copper binding at 90 $\mu\mathrm{M}$ $\mathrm{Cu^{2+}}$ using measured dissociation constants for each construct (Table 1, column 3).

identical ionic radii. The paradigm is that discrimination between these metals requires a well-defined three-dimensional macromolecular structure to hold coordinating ligands at fixed distances and geometry. The imidazole side chain of histidine and its related derivative 4-methylimidazole have intrinsic affinities for Cu^{2+} , Ni^{2+} , and Zn^{2+} of 63–66 μM , 0.9-1.2 mM, and 2.8-3.3 mM, respectively, in aqueous solution (28). This obeys the Irving-Williams series, which relates the decrease in ionic radii with stronger metal-ligand interaction. In this work, we have made comparisons of Cu²⁺ with Ni²⁺ or Zn²⁺ at 1:1 ratios, but also at ratios of approximately 1:10 and 1:100 for Cu²⁺:Ni²⁺ and Cu²⁺:Zn²⁺, respectively. This permits us not only to evaluate absolute specificity of copper effects but also to characterize the relative contribution of intrinsic chemistry and macromolecular structure.

Native state β 2m binds Cu²⁺ specifically with an affinity of 2.8 µM and maps via NMR (Figure 1) and mutagenesis to H31 (Figures 6 and 9A). As this binding constant is \sim 20fold tighter than the intrinsic affinity of imidazole for copper, it suggests that other substituents provide energetically favorable coordination to the metal. These ligands may include the protein backbone, water, and/or W60. This would be similar to coordination of Cu²⁺ by His, Gly backbone, water, and Trp recently reported for the octapeptide repeat of mammalian prion (29). Competition of Cu²⁺ with Ni²⁺ in mass spectra clearly shows Ni2+ and Cu2+ bind to the same site in β 2m (Figure 8). Competition of Cu²⁺ by Zn²⁺ in fluorescence spectra also shows Cu²⁺ and Zn²⁺ bind to the same site in β 2m (Figure 6). The affinities of β 2m for Ni^{2+} and Zn^{2+} are 400 and 1500 μM , respectively. These are $\sim^{1}/_{100}$ and $\sim^{1}/_{500}$ of the binding constant for Cu²⁺, and 10- and 5-fold, respectively, less than expected for binding by an unstructured imidazole. Thus, the additional substituents which enhance copper binding not only are increasing affinity but also are structured to enhance specificity for copper. The specificity of Cu^{2+} binding to $\beta 2m$ is therefore the result of a defined metal binding site.

Binding of copper by β 2m is favorable and therefore must stabilize the native state relative to unfolded state(s). To account for the net destabilization of β 2m by metal, the unfolded state must also bind Cu²⁺. In principle, any protein containing a buried histidine should yield this effect. Consider a hypothetical protein with a buried histidine which is wholly exposed upon unfolding. Given a dissociation constant of 4-methylimidazole for Cu²⁺ of 66 μM and a copper concentration of 90 μ M, for example, in Figure 4, this gives a net destabilization of ~0.7 kJ/mol. This is small compared to the value of 20-60 kJ/mol that is typical of protein stability and is consistent with the fact that histidine residues are frequently observed within the interior of protein structures. Of the four histidine residues of β 2m, only one is completely buried (H84; see Table 1). If it is assumed that all were buried and all were exposed upon unfolding to a random coil state, the net destabilization would be \sim 2.8 kJ/mol. This is on the order of $^{1}/_{10}$ of the observed $\Delta\Delta G$ upon metal binding (Figure 4D) which indicates that additional structural factors must contribute to the observed destabilization.

Analysis of structural factors relevant to destabilization of β 2m is conducted here by mutation at each histidine residue. This has demonstrated that native state binding and protein destabilization are structurally distinct events. Mutation of H31 to phenylalanine increases the sensitivity of β 2m to destabilization by Cu2+ by 5.9 kJ/mol over that of the wild type ($\Delta\Delta\Delta G$, Figure 4D). We further note that the level of native state binding to Cu^{2+} diminishes from 3 to 40 μ M upon mutation of H31 (Figure 6B). This corresponds to a relative change in binding induced stability ($\Delta\Delta G$) of 6.4 kJ/mol at 90 μ M Cu²⁺. Clearly, the change in binding energy is within error of the change in copper induced destabilization. This suggests that the entirety of the increased sensitivity of H31F to unfolding by copper is due to the loss of stabilization of the native state provided by ligand binding. We initially suggested that binding of Cu^{2+} by $\beta 2m$ was destabilizing in a manner akin to protonation (13); i.e., binding and destabilization could be mediated by the same residues. Others have also offered reasonable explanations for the creation of non-native states upon copper binding by the folded protein (23). However, the changes in energetics observed here for H31F are at odds with these explanations. If it is assumed that amyloidosis in β 2m requires destabilization of the folded state to populate amyloidogenic intermediates, then the structures formed upon native state binding of Cu²⁺ (Figure 2B) are unlikely to represent such an intermediate.

Histidine residues 13, 51, and 84 all contribute to the destabilization of β 2m. The simplest model in which to interpret these results is

$$\mathrm{UCu}_n \overset{K_\mathrm{d}^\mathrm{U}}{\longleftrightarrow} \mathrm{U} + n\mathrm{Cu} \overset{K_\mathrm{u}}{\longleftrightarrow} \mathrm{F} + \mathrm{Cu} \overset{K_\mathrm{d}^\mathrm{F}}{\longleftrightarrow} \mathrm{FCu}$$

Separate dissociation constants describe binding to either the unfolded state (K_d^U) or the folded state (K_d^F). The folding stability is characterized by K_u and is invariant to changes

in copper concentration. n is the stoichiometry of this binding and has two limiting conditions: n=1, which requires cooperative formation of a binding site in which all three histidine residues participate in the coordination of metal, and n=3, in which each histidine binds independently to a separate Cu^{2+} .

In the context of this model, the changes in stability due to copper (Figure 4) are solely the result of metal binding to the unfolded state. The $\Delta\Delta G_{\rm u}^{\rm Cu}$ values for the contribution of each histidine to the destabilization of $\beta 2{\rm m}$ (Figure 9B and Table 1) confirm that H31 does not participate in protein destabilization. Whereas H13, H51, and H84 each contribute on the order of 10 kJ/mol to the destabilization of the protein, H31 gives less than 1 kJ/mol. The sum of the free energies of H13, H51, and H84 mutants is 30.7 kJ/mol which is within 1.1 kJ/mol of the destabilization of the wild-type protein (29.6 kJ/mol, Table 1). This suggests that all of the destabilization is additive and mediated by these three histidine residues.

Destabilization is most likely mediated by the coordination of a single (n = 1) copper by H13, H51, and H84. The magnitude of the free energy contribution of each histidine (H13, H51, and H84) to destabilization each exceeds by greater than a factor of 10 the magnitude we calculate from changes in solvent exposure of a buried histidine (0.7 kJ/ mol; see above). In addition, since H13 and H51 are mostly exposed (\sim 70%) even in the folded state, there is an even narrower scope for single coordination of copper by individual histidine residues to contribute to the destabilization of this protein. Recent work has indicated a possible role for a number of residues, including H84, as being central to the formation of an acid molten globule state of β 2m (30). H84A is the least sensitive of our constructs to destabilization by Cu²⁺. It is possible, therefore, that acid molten globular structures are similar to copper induced conformational states.

The most compelling reason to assert the existence of a single copper center in non-native β 2m is the observation of specificity for Cu²⁺ over Ni²⁺ and Zn²⁺ for destabilization. When the stabilization of the native protein by metal ion binding is taken into account (see the above model), Cu²⁺ binding to the unfolded state contributes 17.1 kJ/mol at 20 μ M Cu²⁺ while Zn²⁺ contributes 18.4 kJ/mol at 4000 μ M. At 2000 μ M Ni²⁺, the contribution of binding to the unfolded state is 8.2 kJ/mol. Binding to the native state accounts for all observed effects due to Ni²⁺. The requirement for nearly 1000-fold greater Zn²⁺ than Cu²⁺ for an equivalent energy of binding to the unfolded state is consistent with changes in the affinity of three imidizole groups for these metals (28); i.e., the selectivity of Cu²⁺ over Zn²⁺ by the unfolded state is largely derived from differences in the intrinsic affinity of imidazole for these two metals. However, intrinsic affinities of three coordinating imidazoles are nearly identical for Zn²⁺ and Ni²⁺. The absence of any apparent binding by Ni²⁺ to the unfolded state of β 2m requires that we invoke protein structure to account for the selectivity. We conjecture that the nature of this structure is tetrahedral as Cu²⁺ and Zn²⁺ prefer this binding geometry. Ni²⁺, in contrast, prefers a square planer arrangement of its ligands.

The presence of a well-defined substructure in non-native, unfolded states of β 2m is exciting given our recent evidence

suggesting Cu²⁺ is central to amyloid formation and pathogenesis of DRA (13). Destabilization of protein native structure is generally a prerequisite for fibrillogenesis (31). However, wholly unfolded states of amyloid proteins, e.g., transthyretin, are not thought to contribute greatly to assembly. While specific formation by unfolded states seems to be implausible, it has recently been observed that local structure exists in unfolded states of several model systems, e.g., reduced lysozyme (32) and staphylococcal nuclease (33). Some amyloid proteins may therefore have sequences which, in the presence of divalent cation, give rise to well-defined substructures within an otherwise unfolded polypeptide chain. It is these structures that may then contribute to amyloid assembly. As Cu²⁺ has been recently implicated in Alzheimer's (10), Parkinson's (11), prion protein (9), and immunoglobulin light chain amyloidosis (12), the generality of this phenomenon may prove to be widespread.

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