Visualization of Transient Protein-Protein Interactions that Promote or Inhibit Amyloid Assembly

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SUMMARY

In the early stages of amyloid formation, heterogeneous populations of oligomeric species are generated, the affinity, specificity, and nature of which may promote, inhibit, or define the course of assembly. Despite the importance of the intermolecular interactions that initiate amyloid assembly, our understanding of these events remains poor. Here, using amyloidogenic and nonamyloidogenic variants of β2-microglobulin, we identify the interactions that inhibit or promote fibril formation in atomic detail. The results reveal that different outcomes of assembly result from biomolecular interactions involving similar surfaces. Specifically, inhibition occurs via rigid body docking of monomers in a head-to-head orientation to form kinetically trapped dimers. By contrast, the promotion of fibrillation involves relatively weak protein association in a similar orientation, which results in conformational changes in the initially nonfibrillogenic partner. The results highlight the complexity of interactions early in amyloid assembly and reveal atomic-level information about species barriers in amyloid formation.

INTRODUCTION

The assembly of proteins into amyloid fibrils is a complex process requiring specific and sequence-dependent polymerization of initially unfolded or partially folded monomers into fibrils with elaborate cross-β architectures (Greenwald and Riek, 2010). Despite recent insights into the structural characteristics of the amyloid fold (Eisenberg and Jucker, 2012; Fitzpatrick et al., 2013), the process of amyloid assembly is less well understood in structural terms. Assembly from natively folded precursors is commonly initiated by partial unfolding (Chiti and Dobson, 2009). These nonnative species then combine, generating an array of oligomeric intermediates that are transiently populated and usually heterogeneous in mass and conformation (Cremades et al., 2012; Smith et al., 2010). Although recent advances in structural methods have enabled the conformational properties of rarely populated, partially folded monomers of aggregation-prone proteins to be determined (Eichner et al., 2011; Jahn et al., 2006; Neudecker et al., 2012), the nature of the first protein-protein interactions that initiate amyloid formation remains unclear. Early in assembly an array of biomolecular interactions arise, some of which may be productive for amyloid formation, whereas others may be unproductive, with the potential to inhibit or retard amyloid assembly (Campioni et al., 2010; Cremades et al., 2012). The course of amyloid assembly thus may depend on the stability and lifetime of the productive interactions versus their unproductive counterparts. From this viewpoint, identifying and characterizing different biomolecular interactions early in amyloid assembly are crucial for a full understanding of the structural, kinetic, and thermodynamic determinants of amyloid formation and for interpreting phenomena such as species barriers in prion formation. Such information could also pave the way toward the design of molecules able to define or control the course of amyloid assembly.

Amyloid formation is highly specific, with only proteins of closely related sequence capable of copolymerization into amyloid fibrils (Sarell et al., 2013a). Copolymerization may occur by cross-seeding, in which monomers of a different sequence are capable of extending preformed fibrils from a related protein (Gasson et al., 2003; Guo et al., 2013). In other cases, copolymerization may occur prior to the critical nucleation step of fibrillation. In this case, monomers or small oligomers coassemble into assembly-competent species in the early stages of amyloid assembly (Middleton et al., 2012; Sarell et al., 2013b). One such example can be found in prions, proteins that possess at least one conformation that is infectious by being able to transmit their structural and pathological properties onto noninfectious prion monomers (Chien and Weissman, 2001; Tessier and Lindquist, 2009). Interestingly, when prion molecules are transferred to different species they can lose their high infectivity, establishing a species barrier (Chien et al., 2003, 2004), or can confer their toxic conformation onto previously innocuous proteins of a related species (Sindi and Serio, 2009). The molecular determinants of species barriers, however, remain unclear.

Here we have explored the nature of protein-protein interactions in the first steps of amyloid formation of β2-microglobulin (β2m), a 99-residue protein that forms amyloid deposits in dialysis-related amyloidosis (DRA) (Gejyo et al., 1985). Despite being the main constituent of fibrils in DRA, wild-type human β2m (hβ2m) is not capable of forming amyloid-like fibrils on an experimentally tractable timescale in vitro at neutral pH without the addition of...
external factors or cosolvents (Calabrese and Miranker, 2009; Eichner and Radford, 2011). By contrast, a truncated variant of β2m in which the N-terminal six amino acids are deleted (∆N6), a species that is found in amyloid fibrils in DRA (Esposito et al., 2000), is able to form amyloid fibrils spontaneously at neutral pH in vitro (Eichner et al., 2011; Esposito et al., 2000). NMR studies have shown this variant to be a close structural mimic of a folding intermediate of full-length hβ2m that contains a nonnative trans X-prolyl bond at Pro32 (I) (Figure 1A), the formation of which has been shown to initiate aggregation (Eichner et al., 2011; Jahn et al., 2006). Importantly, ∆N6 is able to convert hβ2m into an aggregation-competent state at neutral pH when added in a substoichiometric molar ratio (Eichner et al., 2011), despite the high structural and sequence similarity of the two proteins (sequence identity 70%, sequence homology 90%, root-mean-square deviation 0.91 Å) (Figures 1A and 1B). This phenomenon was further investigated here by measuring the kinetics of fibril formation of ∆N6 at pH 6.2 (the pH optimum for ∆N6 fibril formation in vitro; Eichner et al., 2011) to which mβ2m had been added in different molar ratios. To account for the effect of protein concentration on the kinetics of amyloid formation, the total protein concentration was maintained at 60 μM in all experiments. Figure 2A shows that ∆N6 assemblies into fibrils able to bind thioflavin T (ThT) with lag-dependent kinetics typical of β2m amyloid formation (Xue et al., 2008), whereas mβ2m does not form fibrils under the conditions employed. Measured over more than ten replicates, the mean lag time of ∆N6 assembly was 32.7 ± 3.8 hr, after which time long straight fibrils typical of amyloid formed (Figure 2Bi). When mβ2m was mixed with ∆N6 in substoichiometric molar ratios, 4:1 ∆N6:mβ2m or 2:1 ∆N6:mβ2m, the mean lag time

Figure 1. Comparison of β2m Variants
(A) Structures of hβ2m (Protein Data Bank [PDB] ID code 2XKS), ∆N6 (PDB ID code 2XKU) (Eichner et al., 2011), and mβ2m (PDB ID code 1LK2) (Rudolph et al., 2004) (left to right). (B) Secondary structure and sequence alignment of hβ2m and mβ2m. Regions identical in sequence are shown in gray. Regions in close spatial proximity to P32 (BC loop, DE loop, FG loop) are highlighted in the structures in (A) and with dashed boxes in (B). The disulfide bond is shown in yellow in (B). P32 is shown as yellow spheres in (A) and highlighted in red in (B). Positions of the spin labels are shown as spheres and sticks in the structure of ∆N6 in (A) and with an asterisk in (B).

that lead to the promotion (∆N6-hβ2m) or inhibition (∆N6-mβ2m) of fibril formation. The results reveal that the surfaces involved in the inhibition and promotion of fibrillation are similar. However, the spatial distribution and chemical properties of the generated ensembles differ in detail, sufficient to alter the affinities of these interactions and the effects of the biomolecular collision on the conformational properties of the monomeric precursors involved. Our findings highlight the complexity of the first steps in amyloid assembly, wherein protein association via similar binding surfaces results in different molecular outcomes. They also reveal information about the origins of species barriers in amyloid formation and identify the surfaces to target by molecular design to enable the course of amyloid assembly to be controlled and/or defined.

RESULTS

mβ2m Kinetically Inhibits ∆N6 Assembly
In previous studies, we have shown that mβ2m is able to inhibit the assembly of ∆N6 into amyloid-like fibrils when added in a stoichiometric ratio (Eichner et al., 2011), despite the high structural and sequence similarity of the two proteins (sequence identity 70%, sequence homology 90%, root-mean-square deviation 0.91 Å) (Figures 1A and 1B). This phenomenon was further investigated here by measuring the kinetics of fibril formation of ∆N6 at pH 6.2 (the pH optimum for ∆N6 fibril formation in vitro; Eichner et al., 2011) to which mβ2m had been added in different molar ratios. To account for the effect of protein concentration on the kinetics of amyloid formation, the total protein concentration was maintained at 60 μM in all experiments. Figure 2A shows that ∆N6 assemblies into fibrils able to bind thioflavin T (ThT) with lag-dependent kinetics typical of β2m amyloid formation (Xue et al., 2008), whereas mβ2m does not form fibrils under the conditions employed. Measured over more than ten replicates, the mean lag time of ∆N6 assembly was 32.7 ± 3.8 hr, after which time long straight fibrils typical of amyloid formed (Figure 2Bi). When mβ2m was mixed with ∆N6 in substoichiometric molar ratios, 4:1 ∆N6:mβ2m or 2:1 ∆N6:mβ2m, the mean lag time...
increased to 63.2 ± 3.8 and 91.0 ± 6.2 hr, respectively (Figure 2A), although fibrils formed over the 1 week (120 hr) time course of the experiment using a 5:1 molar ratio of the two proteins (Figure 2B). When the two proteins were mixed in a 1:1 molar ratio, complete inhibition ensued (Figure 2A). The dependence of the lag time on the concentration of mβ2m added (Figure S1A available online) suggests that inhibition of fibrillation is a kinetically determined process. Accordingly, increasing the molar ratio of mβ2m to ΔN6 delays, but does not inhibit, the formation of amyloid. In support of this notion, the mixtures of ΔN6:mβ2m that did not show evidence of fibril formation after 120 hr were incubated for longer periods of time (≥350 hr) and the extent of fibril formation was again measured using ThT fluorescence and negative-stain electron microscopy (EM). These experiments showed that fibrils were able to form after extended incubation times, with the lag time depending on the excess of mβ2m added (Figures S1B–S1D). These findings confirm that the interaction between ΔN6 and mβ2m retards fibril assembly but, because fibrils are thermodynamically favored, the kinetic barrier to their formation is eventually overcome.

To identify whether mβ2m is incorporated into fibrils when mixed with ΔN6, the aggregates formed in samples containing...
different molar ratios of ΔN6:mβ2m after 350 hr were collected by centrifugation, depolymerized by incubation at 100°C in SDS-PAGE loading buffer or by incubation in 1,1,1,3,3,3-hexafluoro-2-isopropanol (HFIP), and subjected to analysis by SDS-PAGE or electrospray ionization mass spectrometry (ESI-MS) (Experimental Procedures; Supplemental Experimental Procedures). As a control, fibrils were assembled from ΔN6 alone, incubated subsequently with the same concentrations of monomeric mβ2m, and analyzed in a similar manner. The results of these experiments showed that mβ2m associates with the ΔN6 fibrils to a similar extent irrespective of whether the protein was added pre- or postassembly (Figures 2C and 2D). These results indicate that mβ2m is not incorporated into the ΔN6 fibrils but associates with the fibrillar surface subsequent to assembly. By contrast, hβ2m has been shown to be incorporated into the fibril core when incubated with ΔN6 in a 1:1 ratio at pH 6.2 (Sarell et al., 2013b).

**Different Binding Affinities for the Inhibition and Promotion of Fibril Assembly**

To investigate the interfaces involved in the inhibition (ΔN6-mβ2m) or promotion (ΔN6-hβ2m) of amyloid assembly, NMR studies were carried out by mixing 15N-labeled ΔN6 with 15N-labeled mβ2m or hβ2m (80 μM) and monitoring the chemical shift perturbations upon binding using 1H-15N HSQC spectra (Experimental Procedures). For both interaction types, the exchange was found to be in the intermediate-to-fast regime (data not shown), giving rise to small, but significant, chemical shift changes upon binding.

In the case of the inhibitory complex (ΔN6-mβ2m), changes in the 1H-15N HSQC spectrum, including chemical shift differences and exchange line broadening, were observed for a subset of resonances, even when the proteins were mixed in substoichiometric ΔN6:mβ2m ratios. Residues that show significantly altered chemical shifts upon binding are localized in the BC and DE loops in the apical region of mβ2m (Figure 3A). By contrast, an excess (≥80 μM) of 15N-labeled ΔN6 was required to observe significant chemical shift changes in the spectrum of 15N-labeled hβ2m (Figure 3B). In this case, the residues experiencing significant chemical shift differences include the N-terminal regions, the B strand, and the BC and DE loops (Figure 3B). Globally fitting the resulting data (Supplemental Experimental Procedures) yielded Kd values of 68 ± 20 μM for the mβ2m-ΔN6 interaction and 494 ± 180 μM for the interaction between ΔN6 and hβ2m (Figures 3C and 3D). Together, these data suggest a larger interface for the ΔN6-hβ2m interaction (more...
residues experience significant chemical shift perturbations) in comparison to its inhibitory ΔN6-mβ2m counterpart, despite an ~7-fold decrease in binding affinity.

**Inhibition and Promotion of Fibril Formation Involve Similar Binding Interfaces**

Although chemical shifts are excellent probes of protein-protein interactions, they can be affected by long-range effects upon binding (Zhuravleva and Gierasch, 2011). Thus, we next sought to investigate the nature of the protein-protein interactions that lead to inhibition (ΔN6-mβ2m) or promotion (ΔN6-hβ2m) of fibril formation in more detail using PRE studies. The PRE depends on the distance between a paramagnet and adjacent nuclei and can provide long-distance (~30 Å) information quantified by the HN-Γ₂ PRE rate (Supplemental Experimental Procedures) for each amide proton (Clore and Iwahara, 2009). The PRE approach is ideally suited to the analysis of weak intermolecular associations (Clore and Iwahara, 2009), providing distance information that can be used to visualize transient and lowly populated (<0.5%) protein states (Tang et al., 2006, 2008) such as those occurring in the early stages of amyloid formation. To enable these experiments, variants containing a solvent-exposed cysteine were created in ΔN6 by mutating either S20 (AB loop), S33 (BC loop), or S61 (DE loop) to cysteine (Figure 1A) while maintaining the disulfide bond involving C25 and C80 (Experimental Procedures). Chemical modification with (1-oxyl-2,2,5,5-tetramethyl-3-pyrroline-3-methyl) methanethiosulfonate (MTSL) yielded ΔN6 molecules 100% labeled at a single site (Experimental Procedures). These chemically modified molecules were then used in PRE studies to map the interactions between 14N-labeled and MTSL-labeled ΔN6 with 15N-labeled hβ2m or mβ2m, each pair in a stoichiometric ratio (60 M each) at pH 6.2 and 25°C (Experimental Procedures). Under the conditions employed, and in the absence of agitation, fibril formation does not occur for either pair of proteins over the time course of the experiment (<40 hr). Accordingly, the difference in the 1H R₂ relaxation rates of the 15N-labeled protein (hβ2m/ mβ2m) in the presence of oxidized or reduced MTSL-labeled 14N-ΔN6 was measured (HN-Γ₂ rate) (Experimental Procedures) and used to map the interaction surfaces of the different protein pairs.

The PRE data collected for the inhibitory interaction between 14N-labeled ΔN6 (S61C-MTLS) and 15N-labeled mβ2m are shown in Figure 4Aii and Figure S2A. Backbone assignments for mβ2m at pH 6.2 were obtained using standard triple-resonance NMR experiments and uniformly 15N/13C-labeled protein (Experimental Procedures). The results showed high Γ₂ values (Γ₂ > 60 s⁻¹) for residues in the BC and DE loops of mβ2m and lower Γ₂ values (<60 s⁻¹) for residues in the N-terminal 10 residues and the FG loop. These regions cluster on one side of mβ2m surrounding P32 (Figure 1A; Figure 4A, inset), a residue that undergoes cis-trans isomerization known to be required for amyloid formation from hβ2m (Eichner et al., 2011; Sakata et al., 2008). A similar PRE pattern was obtained when the spin label was attached at position 33 (Figure 4Ai). The results suggest that the region of mβ2m surrounding P32 is involved in the interaction with ΔN6 to create a heterodimer (as supported by analytical ultracentrifugation; see below) that kinetically inhibits amyloid formation. Consistent with this supposition, when the spin label is moved to position 20 on 14N-labeled ΔN6, the Γ₂ rates of mβ2m in the BC and DE loops are substantially reduced (<25 s⁻¹) (Figure 4Aiii), suggesting that S20 is distant from the site of interaction (Supplemental Experimental Procedures). These data suggest, therefore, that a head-to-head configuration of the ΔN6-mβ2m heterodimer, involving the BC and DE loops from both monomers, creates the inhibitory complex.

Having identified the protein-protein interactions that lead to the inhibition of ΔN6 fibril formation, we next investigated the interactions that lead to ΔN6-induced promotion of hβ2m fibril assembly. Again, 14N-labeled ΔN6 was spin labeled with MTSL at residues 61, 33, or 20 and PREs to 11N-labeled hβ2m were measured (Figure 4B; Figure S2B). In marked contrast with the results obtained for the ΔN6-mβ2m interaction, the magnitude of the Γ₂ values is reduced significantly when the spin-labeled ΔN6 variants are mixed with hβ2m (compare Figures 4Aii and 4Bii with Figures 4Bi and 4Biii), consistent with the ~7-fold lower Kₐ of the hβ2m-ΔN6 complex (Figures 3C and 3D). Despite the differences in magnitude of the Γ₂ values for the two complexes, the pattern of HN-Γ₂ values obtained is similar to that for the ΔN6-mβ2m interaction, with the largest PREs observed for residues 55–65 in the DE loop and 26–34 in the BC loop when the spin label is attached at position 61 (Figure 4B). Residues in the N-terminal region (residues 2–10) showed increased PRE rates when the spin label is attached at position 33, which were not observed when MTSL was added at position 61 (Figures 4Bi and 4Bii). Again, only very small PREs were observed when MTSL was added at position 20 (Figure 4Biii). These results suggest that the promotion of hβ2m fibril formation also involves a head-to-head association of the two monomers.

**Distinct Conformational Ensembles with Structurally Similar Binding Surfaces**

To obtain more detailed insights into the protein complexes that give rise to the inhibition or promotion of amyloid formation, the PRE data were used in a rigid body/torsion angle simulated annealing approach to generate structural ensembles of the different complexes by minimizing the difference between the observed and calculated Γ₂ values. PRE data for each complex obtained using spin labels at positions 33 and 61 in ΔN6 were fitted simultaneously, along with data from chemical shift perturbations upon binding that were treated as ambiguous distance restraints (see below and Experimental Procedures). Data arising from spin-labeled ΔN6 at position 20 were not included (Supplemental Experimental Procedures). The population of the interconverting species was set to 18% in both cases based on the known Kₛ of each complex.

In a first series of simulated annealing calculations, the interconverting species were represented as a single conformer (N = 1) (Experimental Procedures). The results of this analysis revealed a head-to-head configuration for the association of ΔN6 with mβ2m in which the DE loops from each monomer make the majority of the intermolecular contacts (Figure 5A). Interestingly, the high Q factor (0.54; Figure S2C; Supplemental Experimental Procedures) suggests that multiple conformations are required to satisfy the experimental restraints. In exchanging
Figure 4. Interaction Interfaces in Different Protein Complexes

(A) Per-residue $\Gamma_2$ rates of m$\beta_2$m (60 $\mu$M) when MTSL is attached to S61 (i), S33 (ii), or S20 (iii) on $\Delta$N6 (60 $\mu$M) colored according to their amplitude (blue, not assigned; gray, insignificant; yellow, $>20$ s$^{-1}$; red, $>60$ s$^{-1}$) at pH 6.2, 25°C. The structure of m$\beta_2$m as a surface representation colored by the amplitude of the $\Gamma_2$ rates is shown (insets). Red crosses indicate residues for which the $\Gamma_2$ rate is either too large to appear on this scale or resonances broadened beyond detection when the spin label is oxidized and hence the $\Gamma_2$ rate cannot be measured. Blue dots represent proline or overlapping resonances, and blue crosses denote residues for which the assignments are missing. Error bars were calculated from the noise level in the experiment.

(B) As in (A) but for the interaction between 14N- and MTSL-labeled $\Delta$N6 (60 $\mu$M) and 15N-labeled h$\beta_2$m (60 $\mu$M). The structure of h$\beta_2$m is colored according to the amplitude of the $\Gamma_2$ rates after extrapolation to the same % bound as in (A) (blue, not assigned; gray, insignificant; yellow, $>9 \times 4$ s$^{-1}$; red, $>16 \times 4$ s$^{-1}$). Note that the scale is expanded in (B).

(C) The distribution of the m$\beta_2$m molecules in the $\Delta$N6-m$\beta_2$m complex, with the m$\beta_2$m ensemble shown as a pink surface around $\Delta$N6 (cartoon). The 50 top-scoring ensembles (N = 2, 2 × 50 structures) were included in the calculation.

(D) As in (C) but for the $\Delta$N6-h$\beta_2$m association. The pose of $\Delta$N6 is identical to (C) and the ensemble of h$\beta_2$m subunits is colored in blue. The BC, DE, and FG loops of $\Delta$N6 are highlighted in green, yellow, and blue, respectively, and the positions of the spin label (S20, S33, and S61) are shown as spheres.
systems the observed PRE rate is the weighted population average of the species in solution, as long as those are in the fast exchange regime (Iwahara et al., 2004). In this case, the PRE methodology allows the visualization of the ensemble of the interconverting species. Increasing the number of conformers to two \((N = 2)\) results in a significant decrease in the \(Q\) of the interconverting species. Increasing the number of conformers to three \((N = 3)\) results in an even greater decrease, with a significant decrease \((Q = 0.36)\) when \(N\) is increased to three \((N = 3)\) (Figures S2C and S2D). Similar analysis of the ensemble for the \(D_{N6}\) molecules cluster around the DE loop of \(m_{β_2m}\), whereas the second cluster of \(ΔN6\) molecules locates opposite the edge strands D and C (Movie S2). On the other hand, the \(ΔN6-h_{β_2m}\) interaction is more heterogeneous, extending to both sides of the apical region of \(ΔN6\) (around P32) (Figure 4D; Figure S3B; Movie S1). The volume of the \(ΔN6-m_{β_2m}\) density map is calculated to be \(7,157\) Å\(^3\), whereas that of the \(ΔN6-h_{β_2m}\) cluster is almost twice as large \((13,670\) Å\(^3\); a cutoff of 40% was used in both cases; Table S1). Interestingly, the distributions of \(m_{β_2m}\) and \(h_{β_2m}\) molecules around \(ΔN6\) do not completely overlay. Areas showing high intermolecular contacts unique to the \(ΔN6-h_{β_2m}\) complex involve the BC and FG loops of \(h_{β_2m}\) (Figure 4D; Figure S3B). A correlation between the hydrophobic surface area of \(m_{β_2m}\) (shaped mainly by the region surrounding the DE loop) and the distribution of \(ΔN6\) molecules is observed, indicating that this interaction interface is predominantly hydrophobic in nature, with residues F56, W60, and F62 participating in key intermolecular contacts (Figure 4D; Figure S3B; Movie S1). By contrast, the apical region of \(h_{β_2m}\) (DE, BC, and FG loops) displays less solvent-exposed hydrophobic surface area and a greater predominance of charged residues that reflect the differences in the sequence of the proteins in these regions (Figure 1; Figure S3D; Table S1; Movie S2). Together, the results indicate that inhibition of \(ΔN6\) fibril formation involves a “specific” head-to-head protein association driven by hydrophobic interactions with \(m_{β_2m}\). On the other hand, the \(ΔN6-h_{β_2m}\) interaction, although also adopting a head-to-head configuration, is weaker, more heterogeneous, and involves electrostatic interactions. Whether these data reflect the formation of a range of “encounter complexes”
between ΔN6 and hβ2m that is not observed for the ΔN6-mβ2m interaction, or whether they report on the transient formation of higher-order oligomers between ΔN6 and hβ2m, remains to be resolved.

**Mutation of Aromatic Residues Prevents Inhibition of ΔN6 Assembly by mβ2m**

To confirm that the head-to-head association of ΔN6 with mβ2m is involved in inhibition of fibril formation, two amino acid substitutions (F56E and W60E) were introduced into mβ2m at sites that were found to participate in the majority of intermolecular contacts between the two molecules (Figure 5A; Figure S4A). The ability of this variant to bind to ΔN6 and to inhibit fibril assembly was then monitored using NMR and ThT fluorescence assays, respectively. When 14N-labeled F56E/W60E mβ2m (160 μM) was mixed with 15N-labeled ΔN6 (80 μM) at pH 6.2, only small changes in the chemical shifts of ΔN6 (~20% in comparison to wild-type mβ2m) were observed in the BC, DE, and FG loops (Figure 5B; Figures S5A and S5B), consistent with the proteins no longer interacting tightly. Consistent with these observations, F56E/W60E mβ2m is unable to inhibit ΔN6 fibril assembly when added in a 2-fold molar excess (Figure 5C; Figures S5C and S5D), conditions under which wild-type mβ2m delays the onset of amyloid for more than 120 hr (Figure 2A; Figure S1C). The interaction of wild-type mβ2m with ΔN6 prevents the formation of oligomeric species by the latter protein as observed by sedimentation velocity analytical ultracentrifugation (AUC) (Figure 5D), resulting in a monomer-dimer (~80:20) equilibrium, consistent with a specific interaction as suggested by the analysis of the PRE data. Notably, under identical conditions, the F56E/W60E variant abolishes the ability of the murine protein to dissociate preformed oligomers of ΔN6 (Figure 5D).

**Binding-Induced Unfolding versus Rigid Body Docking: A Rationale for the Outcome of Biomolecular Collision**

To investigate why biomolecular collision of hβ2m or mβ2m with ΔN6 results in different outcomes of assembly, the effect of ΔN6 binding on the conformational dynamics of each monomer was measured using hydrogen-deuterium (H/D) exchange. In each case, the rate of H/D exchange of monomeric (unbound) hβ2m/mβ2m was compared with its ΔN6-bound counterpart at pH 6.2, using samples in which the protein concentrations of hβ2m/mβ2m + ΔN6 were adjusted to generate complexes containing a similar percent (~20%) of ΔN6-bound hβ2m/mβ2m monomer. These experiments showed that the H/D exchange rates of mβ2m are unaffected (kex increases by less than ~1.3-fold)
upon interaction with $\Delta N6$ (Figure 6A; Figure S6A). By contrast, the addition of $\Delta N6$ to $h\beta2m$ causes a 2- to 3-fold increase in the H/D exchange rates of residues throughout the sequence of $h\beta2m$ (Figure 6B; Figure S6B), consistent with an increase in global dynamics of the protein upon interaction with $\Delta N6$. These results were confirmed using a variety of $\Delta N6$ concentrations for both complexes, ranging from 40 to 320 $\mu$M.

Close examination of the chemical shift changes that occur when $^{15}$N-labeled $\Delta N6$ is added to $^{15}$N-labeled $m\beta2m$ reveals that the residues that undergo significant chemical shift changes also experience increased PRE rates (BC and DE loops), confirming that these regions of the protein form the interaction interface (Figure 3A). On the other hand, residues in the N-terminal region including the AB loop of $h\beta2m$ (residues 12–13) show significant chemical shift changes upon binding to $\Delta N6$ (Figure 3B) but minor PREs (Figure 4B), consistent with these residues not being involved in the interface of the lowest-energy structures of the $\Delta N6$-$h\beta2m$ complex (Figure S3B). These observations suggest that the binding of $\Delta N6$ to $h\beta2m$ provides sufficient energy to alter the conformation of the N-terminal 12 residues of $h\beta2m$ (observed previously by H/D exchange and relaxation NMR methods; Eichner et al., 2011) such that a more amyloidogenic conformation is adopted. By contrast, the nonamyloidogenic (and thermodynamically less stable) $m\beta2m$ ($\Delta G_{un}^\text{mouse} = -10.7 \text{ kJ/mol}, \Delta G_{un}^\text{human} = -22.5 \text{ kJ/mol}$; C. Pasley and S.E.R., unpublished data) is not affected significantly by binding. Differences in cooperativity or local stability of the interacting monomers thus dictate the progress of amyloid assembly.

Finally, the consequences of binding on the conformational properties of $\Delta N6$ were investigated by measuring the changes in the chemical shifts of $^{15}$N-labeled $\Delta N6$ (80 $\mu$M) upon titration with $^{15}$N-labeled $m\beta2m$ (80 $\mu$M) or $^{15}$N-labeled $h\beta2m$ (480 $\mu$M) (~45% $\Delta N6$ bound in each case) (Figures 6C and 6D). Significant chemical shift differences were observed for residues in the BC and DE loops of $\Delta N6$ upon binding to $h\beta2m$ and $m\beta2m$, consistent with the head-to-head structure of both complexes. The larger number of $\Delta N6$ residues showing chemical shift differences observed upon binding and the greater $\Delta$ observed for the $\Delta N6$-$h\beta2m$ complex are consistent with the larger interface of this interaction, but could also suggest that $\Delta N6$ responds to binding $h\beta2m$ by undergoing conformational change. The picture that emerges, therefore, is that the promotion of $h\beta2m$ fibril formation by $\Delta N6$ involves weak binding that nonetheless leads to conformational changes in one or both of the interacting partners. By contrast, the $\Delta N6$-$m\beta2m$ complex, even though employing a similar head-to-head interaction, involves the formation of a relatively specific, tight binding, inhibitory complex with little or no effect on the conformational properties of the interacting partners.

**DISCUSSION**

**Protein Interaction Surfaces and the Molecular Mechanism of $\beta2m$ Aggregation**

Amyloid fibrils share similar structural features based upon a cross-β core, irrespective of the organism of origin, the protein involved, or the sequence of the protein precursor (Eisenberg and Jucker, 2012). Despite their similarity in structure, amyloid fibrils can be beneficial to the organism concerned, whereas for others amyloid formation is deleterious (Otzen, 2010). For each scenario, mechanisms have evolved that either facilitate assembly or protect against the accumulation of aggregation-competent proteins, depending on whether the fibrils are beneficial or not (Bucciantini et al., 2002; Otzen, 2010; Maji et al., 2009). One such example can be found in prions, proteins that possess at least one amyloid-competent conformation that is infectious by being able to transmit its structural and pathological properties onto innocuously folded prion monomers (Sindi and Serio, 2009). When prion molecules are transferred between species, they can lose their infectivity or allow propagation depending on the organism involved, establishing a so-called species barrier (Chien et al., 2003; Tessier and Lindquist, 2009; Baskakov, 2014). The precise molecular details of how and why species barriers occur between very similar proteins remain unclear. $\Delta N6$ has been shown to possess prion-like properties in its ability to convert $h\beta2m$ in an aggregation-prone conformation by biomolecular collision (although the protein is not infectious) (Eichner et al., 2011). Here we show that the prion-like characteristics of $\Delta N6$ are not only limited to its ability to convert $h\beta2m$ into an amyloid-competent conformation but also in its ability to experience species barriers (when the molecule interacts with $m\beta2m$, amyloid assembly is inhibited). The results show that aggregation propensity is not simply related to the kinetic and/or thermodynamic properties of the proteins involved (the least stable $\beta2m$ variant studied here [m$\beta2m$] inhibits assembly, whereas propagation involves interaction of $\Delta N6$ with the most stable variant [h$\beta2m$]). Instead, the fate of amyloid assembly involves a fine interplay between molecular recognition and protein plasticity, which is governed by the precise location and chemical properties of the interfaces involved in the first biomolecular interaction events.

**Interactions that Result in Inhibition or Promotion of Amyloid Assembly**

Amyloid diseases are usually late-onset disorders, with symptoms appearing many decades into life, even for individuals carrying the most deleterious of mutations (Greenwald and Riek, 2010). Why this is the case remains unclear; possibilities include the time taken to nucleate fibril formation, and/or atrophy or overload of the proteostatic mechanisms that protect cells from protein misfolding and aggregation (Balch et al., 2008). Defining the nature of the complex network of protein-protein interactions that form in the earliest stages of amyloid assembly is of crucial importance; therefore, in our quest to understand the events that initiate protein aggregation at a molecular level. Such knowledge will also open the door to the design of inhibitors able to arrest amyloid formation by targeting specific surfaces that block the formation of fibrils and their toxic precursors, thereby halting the disease process at its outset.

Attempts to identify the intermolecular interactions that form early in amyloid assembly have remained a significant challenge as a consequence of the interactions’ heterogeneity and transient nature (Cremades et al., 2012). By exploiting the power of biomolecular NMR methods and applying them to $\beta2m$ sequences from different species, we have been able to define
Protein Interactions in Early Amyloid Assembly

the intermolecular surfaces that determine the course of amyloid assembly. Specifically, we show that the interaction of ΔN6 with mβ2m inhibits aggregation via trapping the amyloidogenic precursor (ΔN6) in kinetically stable dimers (K_d = 68 ± 20 μM). These involve the formation of a relatively well defined interface, stabilized by hydrophobic interactions involving the side chains of residues in the DE and BC loops of both molecules, including F56 and W60 (Figure 7, bottom). Interestingly, mβ2m is the least stable variant of the three β2m homologs studied here, as shown by its increased H/D exchange rates and decreased unfolding free energy relative to ΔN6 and hβ2m (T.K.K., C. Pashley, and S.E.R., unpublished data). Thermodynamically and kinetically unstable proteins, therefore, and not only their stable counterparts (e.g., antibodies or affibodies; Dumoulin et al., 2003; Hoyer et al., 2008), can act as efficient and specific inhibitors of aggregation. Surprisingly, the amyloid-promoting association of ΔN6 with hβ2m also involves a head-to-head interaction similar, but not identical, to that of the inhibitory complex. Consistent with this finding, the folding intermediate I_c of hβ2m that structurally resembles ΔN6 (Eichner and Radford, 2009; Eichner et al., 2011) was recently shown to form transient oligomers during folding that are also organized in a head-to-head configuration, although the structures formed and their implications for aggregation were not described (Rennella et al., 2013).

We show here that the amyloid-promoting interaction between ΔN6 and hβ2m is thermodynamically weaker than its inhibitory counterpart (K_d = 494 ± 180 μM) and involves multiple interaction sites that involve complementary electrostatic interactions between the interacting molecules that are not utilized in its inhibitory ΔN6-mβ2m counterpart. These differences in the interaction interfaces result in binding-induced conformational changes in hβ2m that are manifested by a 2- to 3-fold increase in its hydrogen exchange rates (Figure 7, top). This interaction also alters the conformation of the AB loop of hβ2m, as shown previously (Eichner et al., 2011). Accordingly, ΔN6 is able to act as protein saboteur, each molecule interacting with numerous copies of hβ2m, destabilizing the native fold of hβ2m and allowing P32 to relax from its native cis isomer to its trans form, which then traps the protein irreversibly in an aggregation-competent state. cis Pro32 in hβ2m, therefore, acts as a key switch in amyloid formation. Accordingly, any event that promotes relaxation of Pro32 to the trans conformer (mutation, formation of ΔN6 or its interaction with Cu^{2+} ions, chaperones, or proline isomerase) promotes formation of amyloid fibrils (reviewed in Eichner and Radford, 2011).

### Implications for the Origins of Transmissibility in Amyloid Diseases

The results presented reveal that subtle differences in the nature of protein-protein interactions can give rise to fundamentally different outcomes of amyloid assembly that depend on the affinity of the interaction, the stability of the interacting partners, and the chemical nature of the interacting surfaces. The results have significance that extends beyond the specific case of the β2m variants studied here. The catalytic templating model proposed to explain the conversion of the cellular human prion protein (PrP^C) to its infectious scrapie form (PrP^Sc) is one such case (Aguzzi et al., 2008). Accordingly, mutations that have little effect on the structural and thermodynamic properties of the monomeric PrP precursors (Ba et al., 2009) could alter the surface properties of the protein, influencing the network of intermolecular interactions formed, and hence lead to increased or decreased infectivity. Other amyloid proteins that are intrinsically disordered (such as Aβ40 and α-synuclein) are known to mutually enhance each other’s aggregation (Guo et al., 2013), possibly involving a similar mechanism of binding-induced conformational change. Indeed, heteropolymerization in amyloid assembly seems to be more common than initially anticipated (Sarell et al., 2013a). As shown here, protein association, response to binding, and the effect of transient intermolecular association on the course of assembly are all interlinked. Binding, even to similar surfaces, can cause a different response on the partners.
involved and thus lead to a different outcome of assembly. The HET-S/HET-s prion strains in filamentous fungi represent another example (Greenwald et al., 2010). HET-S, even though 97% identical in sequence to HET-s, does not aggregate, and can also inhibit the propagation of the prion form of HET-s by biomolecular interaction, resembling the effect of m\(\beta_2\)-m on \(\Delta N6\) assembly. A model for prion inhibition by HET-S has been proposed in which HET-S, although able to interact with HET-s and adopt the amyloid \(\beta\)-solennoid fold, is incompetent for further polymerization (Greenwald et al., 2010), further highlighting the observation that collision of similar proteins can result in different outcomes of assembly. Application of the approach taken here for \(\beta_2\)-m to other proteins involved in human disease, including the classic examples of species barriers in PrP propagation (Baskakov, 2014), prion compatibility in yeast and other fungi (Tessier and Lindquist, 2009), and other proteins purported to be infectious (Brundin et al., 2010), will reveal the similarities and distinctions between \(\Delta N6\)-induced conformational conversion and amyloid inhibition and the molecular events occurring in other systems.

As well as providing insights into the molecular origins of species barriers in amyloid formation, the results presented provide opportunities for the design of molecules to control amyloid disease by targeting intermolecular contacts in the specific surfaces involved. The design of small molecules able to disrupt protein–protein interactions and the generation of other reagents (antibodies, affibodies, or nucleic acid aptamers [Bunka et al., 2007] selected to bind to a specific surface) are exciting possibilities for future avenues of research. The complexity of amyloid assembly, especially in the cellular environment, may require multiple routes involving different strategies to delay, prevent, or revert disease to be deployed simultaneously (for example by combining interference of protein assembly with small molecules or aptamers in concert with regulation of the cellular mechanisms that recognize protein misfolding events). The ability to target the earliest biomolecular events in the aggregation cascade offers potential for a route toward amyloid therapy that will add to the arsenal of approaches currently being developed to combat these devastating disorders.

**EXPERIMENTAL PROCEDURES**

**Protein Preparation**

\(h\beta_2\)-m, \(m\beta_2\)-m, and \(\Delta N6\) (\(^{14}\)N- and \(^{15}\)N-labeled) and their variants were expressed and purified as described (Platt et al., 2005).

**Assembly of Amyloid-like Fibrils**

Samples containing 0.6–60 \(\mu\)M protein, 10 mM sodium phosphate buffer (pH 6.2), 83.3 mM NaCl (total ionic strength 100 mM), 0.02% (w/v) sodium azide, and 10 \(\mu\)M ThT were incubated at 37°C in sealed 96-well plates with agitation at 600 rpm (Supplemental Experimental Procedures).

**PRE Experiments**

The \(\Delta N6\) variants (\(^{15}\)N-labeled) C20S, C33S, and S61C modified with MTSL (Supplemental Experimental Procedures) were mixed with \(^{15}\)N-labeled \(h\beta_2\)-m or \(m\beta_2\)-m (60 \(\mu\)M, unless otherwise stated) in 10 mM sodium phosphate buffer (pH 6.2) and \(H_2\)-PRE data were measured as described in Supplemental Experimental Procedures.

**Simulated Annealing Calculations**

All structure calculations were performed using a torsion angle-simulated annealing protocol in XPLOR-NIH as described (Iwahara et al., 2004) (Supplemental Experimental Procedures).

**Kd Measurements**

Binding affinities for the complexes of \(m\beta_2\)-m and \(h\beta_2\)-m with \(\Delta N6\) were determined at pH 6.2 and 25°C by titrating 80 \(\mu\)M \(^{15}\)N-labeled \(m\beta_2\)-m with 0–320 \(\mu\)M \(^{15}\)N-labeled \(\Delta N6\) or 60 \(\mu\)M \(^{15}\)N-labeled \(h\beta_2\)-m with 0–480 \(\mu\)M \(^{15}\)N-labeled \(\Delta N6\) and measurement of the resulting chemical shift changes using \(^1\)H-\(^{15}\)N HSQC spectra (Supplemental Experimental Procedures).

**Hydrogen Exchange Experiments**

The rate of H/D exchange of samples of \(^{15}\)N-labeled \(h\beta_2\)-m or \(m\beta_2\)-m (80 \(\mu\)M) alone or mixed with \(^{15}\)N-labeled \(\Delta N6\) (160 or 40 \(\mu\)M, respectively) to produce \(\sim 22\%\) bound complexes in each case was measured at pH 6.2. Hydrogen exchange was measured using SOFAST-HMQC NMR methods as previously described (Schanda et al., 2003) (Supplemental Experimental Procedures).

**Additional Procedures and Further Information**

Detailed description of all other methods and protocols can be found in Supplemental Experimental Procedures.

**ACCESSION NUMBERS**

Assignments for the backbone atoms of \(m\beta_2\)-m have been deposited in the BioMagResBank under accession number 19772.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, six figures, one table, and two movies and can be found with this article online at http://dx.doi.org/10.1016/j.molcel.2014.05.026.

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Supplemental Information

Visualization of Transient Protein-Protein Interactions that Promote or Inhibit Amyloid Assembly
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Figure S1 - related to Figure 2: Kinetic inhibition of ΔN6 amyloid formation by mβ2m. (A) Plot of the average lag time of fibril formation for different molar ratios of ΔN6:mβ2m. The grey-shaded area represents experiments where the lag time could not be estimated because the protein mixtures did not show an increase in ThT fluorescence by the end of the experiment shown in Figure 2A (120h-red line). Error bars represent the standard error of the mean. (B) Kinetics of amyloid assembly of mixtures of ΔN6:mβ2m: 30µM:30µM, (C) 20µM:40µM and (D) 10µM:200µM over timescales up to 400h monitored using ThT fluorescence. Negative stain EM images of the fibrils formed at the end of each reaction are inset (scale bar = 500nm).
**Figure S2** – related to Figure 4. Agreement between the experimental and back-calculated intermolecular PRE data for the ΔN6-mβ2m and ΔN6-hβ2m interactions. Intermolecular PRE profiles for the interaction between $^{14}$N-ΔN6 spin labeled at position 61 and $^{15}$N-labeled mβ2m (A) or $^{15}$N-labeled hβ2m (B). (C) Plots of the calculated Q factor versus the number of ensemble members for the interaction of ΔN6 with mβ2m. Dashed red lines represent the best Q factor possible for each dataset, if the quality of the fit is comparable to the error of the experimental data and is calculated as described in (Tang et al., 2008). (D) Experimental and calculated PRE data for the ΔN6-mβ2m interaction when the spin label is attached at position 61 (left), 33 (middle) or 20 (right). Predicted PRE rates for an ensemble size of 2 (N=2) are shown in black lines while red dots denote the experimentally measured PRE rates. Data arising from the spin label on position 20 were not used in the fitting. The calculated $\Gamma_2$ rates shown represent the average values (per residue), back-calculated from 50 independent calculations (50x2 structures). All ensemble members were equally weighted during the calculations. (E) As in (C) and (F) as in (D) but for the association of ΔN6 with hβ2m.
Figure S3 – related to Figure 4. Analysis of the ΔN6-mβ2m and the ΔN6-hβ2m interaction ensembles. (A) Contact map for the 50 best scoring ensembles for the ΔN6-mβ2m or (B) the ΔN6-hβ2m interaction. Every non-hydrogen atom with an intermolecular distance less than 4Å to any other (non-hydrogen) atom is identified as a contact. The number of contacts for atoms of each residue is color-coded as shown in the color-bar. (C) Structures mβ2m colored according to the number of intermolecular contacts, using the same color scale as in (A) (left). A surface representation of the protein, colored according to its electrostatic potential (red
negative, blue positive, ±2k_B T) is shown in the right. (D) As in (C) but for hβ_2m. The pose of the proteins is the same in both cases, with the BC, DE and FG loops at the top of the molecule.
**Figure S4 – related to Figure 5.** The roles of F56 and W60 in the interface of different complexes. (A) Normalized frequency histograms of the number of contacts that F56 (left) or W60 (middle) of mβ2m make with residues of ΔN6 in the ΔN6-mβ2m complex. The 50 best-scoring ensembles (N=2, 50x2 structures) were analysed. These histograms essentially represent horizontal slices of the heat maps shown in Figure S3. The rightmost panel shows the position of F56 and W60 of mβ2m (sticks) in the top 10 ensembles. ΔN6 is shown as red cartoon and the positions of the spin labels (S20, S33 and S61) are highlighted in spheres. (B) As in (A) but for the ΔN6-hβ2m interaction. The distribution of F56 and W60 in the ΔN6-hβ2m complex is much more diverse in comparison its ΔN6-mβ2m counterpart. As a consequence, whilst the F56E/W60E mβ2m mutant prevents the association of ΔN6 with mβ2m, a more detailed mutational analysis in the interface of the ΔN6-hβ2m complex is required to unpick the roles of specific residues in the course of assembly (Figure S4B).
Figure S5 –related to Figure 5. Residual interactions between F56E/W60E mβ2m and ΔN6 are not sufficient to inhibit fibrillation. (A) Additional examples of resonances of $^{15}$N-labeled...
ΔN6 (80µM, red) that show chemical shift changes upon the addition of 14N-labeled mβ2m (green), but not its F56E/W60E variant (160µM, blue) at pH 6.2, 25°C. (B) Changes in the chemical shifts of 80µM 15N-labelled ΔN6 upon addition of 160µM 14N-labelled mβ2m (green) or F56E/W60E mβ2m (blue) (pH 6.2, 25°C). Residues that are broadened beyond detection because of exchange line broadening in the spectrum of ΔN6:mβ2m have an arbitrary value of 1. The small chemical shift differences observed in the F56E/W60E mβ2m:ΔN6 sample suggest a residual interaction with increased $K_d$. (C) Negative stain electron micrograph of 20µM ΔN6 mixed with two molar equivalents of F56E/W60E mβ2m (top) or wild-type mβ2m (bottom). Bar represents 500nm. (D) Aggregation kinetics of ΔN6, F56E/W60E mβ2m or wild-type mβ2m (60µM each alone) followed by ThT fluorescence. Three example traces of each protein variant are shown.
**Figure S6 – related to Figure 6.** H/D exchange rates of mβ2m and hβ2m upon interaction with ΔN6. (A) Plots of peak intensity versus time after the initiation of H/D exchange for example residues of 15N-labeled mβ2m alone (80µM, left column) or in the presence of 40µM 14N-labeled ΔN6 (right column). Solid grey dots represent the raw data and solid lines show fits to single exponentials. Error bars were calculated from the noise level of the experiment. Open symbols show the residuals of the fits. (B) as in (A) but for 15N-labeled hβ2m alone (80µM) or in the presence of 160µM 14N-labeled ΔN6.
Supplemental Movies

**Movie S1, related to Figure 4.** The mβ2m-ΔN6 and hβ2m-ΔN6 complexes involve different subunit orientations of a common head-to-head dimer. Movie animation of the structural ensembles shown in Figures 4C and 4D. ΔN6 is shown as cartoon representation with its BC loop highlighted in green, the DE loop in yellow and the FG loop in blue. The ensemble of mβ2m molecules around ΔN6 is shown as a pink surface on the left hand side, while the hβ2m ensemble is shown as a blue surface in the right hand side.

**Movie S2, related to Figure 4.** The mβ2m-ΔN6 and hβ2m-ΔN6 complexes show different chemical properties in the interface. Mβ2m (left hand side) and hβ2m (right hand side) are shown as a surface representation coloured according to their electrostatic potential (±2k_B T), with the BC, DE and FG loops on the top. The ensemble of ΔN6 molecules around mβ2m and/or hβ2m is shown as green and yellow mesh respectively. This representation is essentially the complementary picture of the ensembles shown in Figures 4C and 4D (where mβ2m and/or hβ2m were shown as weighted atomic probability density maps). Note the high correlation between the distribution of ΔN6 molecules around mβ2m with the hydrophobic surface of the latter. By contrast, part of the ΔN6 density map locates opposite the negatively charged part of the BC loop of hβ2m. The electrostatic surface potential was calculated using APBS (Baker et al., 2001) and movies were rendered in Pymol (Schrodinger, LLC, 2010).
Supplemental Tables

Table S1 – related to Figure 4. Analysis of the interfaces of different β2m complexes. The buried surface area is calculated as the sum for the two subunits for each complex and is measured using XPLOR-NIH (Schwieters et al., 2003). Interface residues were identified as NACCESS (Hubbard and Thornton, 1993). Errors represent the standard deviation.

|                  | ΔN6 - mβ2m complex | ΔN6 - hβ2m complex |
|------------------|--------------------|--------------------|
| Density volume (Å³) | 7,157              | 13,670             |
| Buried surface area (Å²) | 1481±290         | 1359±357           |
| % Hydrophobic residues in the interface | 48.6±14.5        | 43.81±12.6         |
| % Charged residues in the interface | 20.4±12.0        | 31.0±14.8          |

1 Within each ensemble member
Supplemental Experimental Procedures

Protein preparation

Mβ2m and F56A/W60A mβ2m were purified using the protocol described in (Eichner et al., 2011), but the refolding buffer was adjusted to pH 8.5 and gel filtration was carried out in 10mM sodium phosphate, pH 8.2. S20C, S33C and S61C mutants of ΔN6 were refolded in 10mM TrisHCl buffer containing 0.64M L-arginine and/or 2mM reduced glutathione and/or 0.2mM oxidized glutathione before purification as in (Ladner et al., 2010).

Assembly of amyloid-like fibrils

Each experiment was repeated over at least 10 replicate samples and the median and/or mean lag time and standard error of the mean (SEM) were determined. Analysis of the soluble and insoluble material in each reaction was carried out at the end of fibril growth (120h or 350h) by collecting insoluble material by centrifugation (15,000g, 20min) and analysis by SDS-PAGE or ESI-MS (the latter subsequent to depolymerization by incubation for 10h in 100% (v/v) 1,1,1,3,3,3-hexafluoro-2-isopropanol (HFIP) (Sarell et al., 2013). Fibril morphology was analyzed using negative stain EM.

Negative-stain EM

Carbon coated copper grids were prepared by the application of a thin layer of formvar with an overlay of thin carbon. Samples were centrifuged (14,000g, 10min) and the pellets were resuspended in fresh 10mM sodium phosphate buffer, pH 6.2, diluted to a final protein concentration of 12µM with deionized water and then applied to the grid in a drop-wise fashion. The grid was then carefully dried with filter paper before it was negatively stained by
the addition of 18 µl of 2% (w/v) uranyl acetate. Micrographs were recorded on a Philips CM10 or a JEOL JEM-1400 electron microscope.

**Analytical ultracentrifugation**

For sedimentation velocity experiments, a total volume of 450µl sample in 10mM sodium phosphate buffer, pH 6.2, 83.3mM NaCl was inserted in standard double-sector Epon centerpieces equipped with sapphire windows, inserted in an An60 Ti four-cell rotor. Sample concentrations included 60µM or 120µM (ΔN6 alone), 60µM ΔN6 mixed with 60µM mβ2m or 60µM F56E/W60E mβ2m at pH 6.2. Absorbance data at 280nm were acquired at a rotor speed of 50,000rpm at 25°C. Data were analyzed using the c(s) continuous distribution of the Lamm equations with the software SEDFIT (Brown and Schuck, 2006).

\[
D(s) = \frac{\sqrt{2}}{18\pi} k Ts^{-\frac{1}{2}}(\eta(f/f_0)_{w})^{-3/2}(1 - \frac{\bar{\eta}}{\eta})^{1/2}
\]

where D(s) is the diffusion coefficient, k Boltzmann’s constant, T the temperature in K, s the sedimentation coefficient, f is the frictional coefficient, f_0 the frictional coefficient of a compact smooth sphere, η the solvent viscosity, ρ represents the solvent density and \(\bar{\eta}\) the partial specific volume.

**Backbone assignments of mβ2m**

Backbone assignments for mβ2m were obtained using triple resonance NMR techniques (HNCA, HNCO, HNCACB, HN(CO)CACB) and samples containing 500-750µM uniformly labeled \(^{13}\)C/\(^{15}\)N mβ2m in 10mM sodium phosphate buffer, pH 6.2, 83.3mM NaCl. Assignments are deposited in BMRB (19772).
**PRE experiments**

The ΔN6 variants (14N-labeled) S20C, S33C and S61C (1-2mg/mL) were incubated with 5mM DTT for 20min, and then labeled immediately with MTSL by incubation with a 40-fold molar excess of the spin label for 4h in 25mM sodium phosphate buffer, pH 7.0, 1mM EDTA at room temperature. Excess spin label was removed by gel filtration (PD10 column, GE Healthcare). Spin-labeled ΔN6 was used directly or stored at -80°C. In all cases 100% labeling resulted at a single site as revealed by ESI-MS (not shown). For each PRE experiment MTSL-labeled 14N-ΔN6 (10-60µM) was mixed with 15N-labeled hβ2m or mβ2m (60-150µM) and the difference of the proton R$_2$ rates between oxidized and reduced (by addition of 1mM ascorbic acid) MTSL-labeled 14N-ΔN6 was measured. Data were recorded at 25°C using a 1H-15N correlation based pulse sequence with 5-6 time-points (0.0016-0.016s) and at least 32 scans per incremental delay, utilizing a Varian-Inova 750MHz spectrometer equipped with a cryogenic probe. R$_2$ rates were extracted by fitting the relaxation data to single exponentials using in-house scripts. The HN-Γ$_2$ rate was then calculated as the difference between the R$_2$ rate in the paramagnetic versus the diamagnetic sample:

$$\Gamma_2 = R_{2,\text{para}} - R_{2,\text{dia}}.$$

Errors were calculated based on the noise of the experiment. The small PRE signal observed when ΔN6 is modified with MTSL at position 20 can be attributed to non-specific binding of the spin label itself, since addition of free MTSL results in a similar PRE profile. Thus, data arising from spin-labeled ΔN6 at position 20 were not included in quantitative analysis of the PRE experiments.

**Simulated annealing calculations**

Simulated annealing calculations were carried out in XPLOR-NIH (Schwieters et al., 2003). To account for the flexibility of the MTSL side chain, the paramagnetic group was
represented as a 5 membered ensemble. The agreement between the experimental and the back-calculated data is described by the PRE Q factor (Tang et al., 2006) defined as:

$$Q = \left[ \sum_i i \left( \Gamma_{2,i}^{\text{obs}} - \Gamma_{2,i}^{\text{calc}} \right)^2 / \sum_i i (\Gamma_{2,i}^{\text{obs}})^2 \right]^{1/2},$$

where $\Gamma_{2,i}^{\text{obs}}$ is the observed $\Gamma_2$ value for residue i and $\Gamma_{2,i}^{\text{calc}}$ is the calculated $\Gamma_2$ value. All calculations were started from randomized starting positions.

The computational strategy employed included two PRE potential terms (arising from S61C-$\Delta$N6 and S33C-$\Delta$N6) and classic geometry restraints to restrict deviation from bond lengths, angles and dihedrals. To generate a dataset suitable for this analysis, PREs arising from position 61 and 33 for the $\Delta$N6-h$\beta_2$m interaction were (each) measured in two independent experiments and the average PRE value for each residue was used for fitting. For the $\Delta$N6-m$\beta_2$m interaction, resonances in the BC and DE loop are not visible in the spectrum of the oxidized sample when the spin label is attached at positions 33 or 61 when the proteins are mixed in a 1:1 molar ratio. To obtain an estimate for the $\Gamma_2$ rate for these residues, the PRE experiments (using spin-labeled $\Delta$N6 at positions 33 or 61) were repeated at different protein concentrations to: 1) improve the signal-to-noise ratio and 2) reduce the concentration of the bound complex so that a more accurate $\Gamma_2$ rate can be measured. PREs were then extrapolated to their values for a 1:1 complex using the measured $K_d$ and the resulting dataset was used for quantitative analysis of the structural properties of the complex. Resonances for which an estimation of the $R_2$ rate in the presence of oxidized spin label was not possible, were incorporated in the protocol as nOe-type of restraints with an upper bound of 11.5Å and a lower bound of 9Å. Additionally, chemical shift perturbations observed upon binding were incorporated as sparse, highly ambiguous intermolecular distance restraints as described in (Clore and Schwieters, 2003). As chemical shifts can be influenced by numerous factors upon protein-protein interaction, the treatment of the derived data undertaken here results in a loose
potential term that is unlikely to bias the structure calculation. Finally, the protocol also included a weak radius of gyration restraint (R_{\text{gyr}}) calculated as $2.2N^{0.38}$, where $N$ is the number of atoms in the complex. $R_{\text{gyr}}$ is required in order to prevent bias towards more extended structures and tends to underestimate the true value of the radius of gyration (Kuszewski et al., 1999).

The aforementioned potential terms were used in a rigid-body energy minimization/ simulated annealing in torsion angle space protocol to minimize the difference between the observed and calculated $\Gamma_2$ rates, starting from random orientations. The first step in the structure calculation consisted of 5000 steps of energy minimization against only the sparse chemical shift restraints, followed by simulated annealing dynamics with all the potential terms active, where the temperature is slowly decreased (3000-25K) over 4fs. During the hot phase (T=3000K) the PRE and nOe terms were underweighted to allow the proteins to sample a large conformational space and they were geometrically increased during the cooling phase. Proteins were treated as rigid bodies until the initiation of the cooling phase, where side chains were allowed to float (semi-rigid body calculation). The final step included torsion angle minimization using all potential terms. Ensemble calculations where the interacting species are represented as multiple states ($N>1$) were carried out as before but in this case the $\Gamma_2^{\text{cal}}$ is calculated as the average value between the conformers. The population of each conformer was set by specifying its weight in the calculation. For ensemble calculations the Q factor is calculated by averaging the predicted PRE value over all ensemble members corrected by their weight (the Q factor of the ensemble of ensembles).

In the case of the $\Delta N6$-m$\beta_2$m interaction when $N=1$, 7 out of 10 lowest energy structures share a backbone RMSD for the m$\beta_2$m subunit of 4.4Å, suggesting that the complex shown in
Figure 5A represents the main associating species in solution. On the other hand, a single conformer representation (N=1) for the ΔN6-hβ2m association yields a hβ2m subunit RMSD of 20.5Å for the 10 lowest energy structures, an observation that also supports the larger conformational ensemble between ΔN6 and hβ2m shown in Figure 4D. Buried surface area calculations were carried out in XPLOR-NIH (Schwieters et al., 2003).

**Fitting $K_d$ values**

The total chemical shift differences as each $^{15}$N-labeled protein was titrated with $^{14}$N-labeled ΔN6 (pH 6.2, 25ºC) was calculated using the function:

$$\Delta \delta_{tot} = \sqrt{(5 \delta^H)^2 + (\delta^{15}N)^2}.$$  

Residues for which the difference in chemical shift upon binding was ≥2 standard deviations from the mean were considered significant. These were used globally to extract the $K_d$ using the function:

$$\Delta \delta = \Delta \delta_{max} \left[ L_T \right] + \left[ U_T \right] + K_d \sqrt{\left[ L_T \right] \left[ U_T \right] + \left[ L_T \right] + \left[ U_T \right] + K_d} - \frac{4\left[ L_T \right] \left[ U_T \right]}{2\left[ L_T \right]},$$

where $\left[ L_T \right]$, $\left[ U_T \right]$ are the total concentrations of the labeled and unlabeled protein added respectively, and $\Delta \delta_{max}$ represents the highest value of the chemical shift difference upon titration. A total of 10 and 8 residues were fitted for the ΔN6:mβ2m and the ΔN6:hβ2m interactions respectively. Four representative examples are shown in Figures 3C and 3D for clarity. Errors on the measured peak positions were calculated as the standard deviation of the mean for residues that show insignificant chemical shift changes. $K_d$s were extracted by Monte Carlo analysis with 1000 steps performed using in-house scripts.

**Hydrogen exchange measurements**

Samples for H/D exchange NMR were made in 10mM sodium phosphate pH 6.2 buffer and
freezed-dried. On the day of the experiment, samples were dissolved in 100% (v/v) D$_2$O containing 83.3mM NaCl and hydrogen exchange was measured using SOFAST-HMQC NMR methods (Schanda et al., 2005) utilizing a 750MHz Varian Inova spectrometer (Agilent) equipped with a cryogenic probe. The dead time of the experiment was 5-10min and each spectrum was acquired for 10-15min. H/D exchange rates for mβ2m/hβ2m were measured at 25ºC and 37ºC respectively. Rates were extracted by fitting single exponentials (Figure S6).
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