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Monitoring the Interaction between $\beta_2$-Microglobulin and the Molecular Chaperone $\alpha$B-crystallin by NMR and Mass Spectrometry: $\alpha$B-CRYSTALLIN DISASSOCIATES $\beta_2$-MICROGLOBULIN OLIGOMERS

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Monitoring the Interaction between \(\beta_2\)-Microglobulin and the Molecular Chaperone \(\alpha B\)-crystallin by NMR and Mass Spectrometry

\(\alpha B\)-CRYSSTALLIN DISSOCIATES \(\beta_2\)-MICROGLOBULIN Oligomers

Background: \(\beta_2\)-Microglobulin (\(\beta_2\m\)) is a paradigmatic amyloidogenic protein.

Results: \(\textit{In vitro}\), the molecular chaperone \(\alpha B\)-crystallin affects the oligomerization and the fibrillogenesis of \(\beta_2m\) and its R3A mutant.

Conclusion: \(\alpha B\)-crystallin prevents \(\beta_2m\) aggregation at various stages of its aggregation pathway.

Significance: Molecular chaperones may be relevant to amyloid formation \(\textit{in vivo}\).

The interaction at neutral pH between wild-type and a variant form (R3A) of the amyloid fibril-forming protein \(\beta_2\)-microglobulin (\(\beta_2m\)) and the molecular chaperone \(\alpha B\)-crystallin was investigated by thioflavin T fluorescence, NMR spectroscopy, and mass spectrometry. Fibril formation of R3A\(\beta_2m\) was potently prevented by \(\alpha B\)-crystallin. \(\alpha B\)-crystallin also prevented the unfolding and nonfibrillar aggregation of R3A\(\beta_2m\).

From analysis of the NMR spectra collected at various R3A\(\beta_2m\) to \(\alpha B\)-crystallin molar subunit ratios, it is concluded that the structured \(\beta\)-sheet core and the apical loops of R3A\(\beta_2m\) interact in a nonspecific manner with the \(\alpha B\)-crystallin. Complementary information was derived from NMR diffusion coefficient measurements of wild-type \(\beta_2m\) at a 100-fold concentration excess with respect to \(\alpha B\)-crystallin. Mass spectrometry acquired in the native state showed that the onset of wild-type \(\beta_2m\) oligomerization was effectively reduced by \(\alpha B\)-crystallin. Furthermore, and most importantly, \(\alpha B\)-crystallin reversibly dissociated \(\beta_2m\) oligomers formed spontaneously in aged samples. These results, coupled with our previous studies, highlight the potent effectiveness of \(\alpha B\)-crystallin in preventing \(\beta_2m\) aggregation at the various stages of its aggregation pathway. Our findings are highly relevant to the emerging view that molecular chaperone action is intimately involved in the prevention of \(\textit{in vivo}\) amyloid fibril formation.

It is well recognized that there is a close association between a wide range of diseases and protein aggregation. For example, Alzheimer, Parkinson, Creutzfeldt-Jakob, and Huntington diseases are characterized by the accumulation of protein deposits or plaques in which the aggregated protein is arranged in highly ordered amyloid fibrils (1). Understanding the relationship between the formation of amyloid fibrils and protein aggregation diseases is the focus of much research activity. It is clear that amyloid fibril formation arises from the repertoire of partially folded intermediate states of proteins, and there is significant evidence that the process of aggregation of these states causes cellular toxicity which, in turn, contributes to the particular disease (1). Intriguingly, various reports have suggested that the cellular toxicity of the putative causative agent in Alzheimer disease, the amyloid \(\beta\) (\(A\beta\))^5 peptide, and other amyloidogenic proteins, does not arise from the insoluble amyloid deposits but rather from the soluble prefibrillar oligomers (2–6).

Molecular chaperone proteins interact with the intermediate states of proteins. Most likely, they are important in preventing...
β2-Microglobulin (β2m) is the nonpolymorphic light chain component of class I major histocompatibility complex (MHC-I) (10). In vivo, β2m causes amyloidosis via accumulation in the tissues of patients with renal failure as a result of long term hemodialysis (11). The structure of β2m and the molecular aspects of its amyloid transition have been extensively investigated in recent years, not only because of the clinical relevance of dialysis-related amyloidosis but also as a model for other amyloidogenic proteins (12–19).

Among the amyloidogenic proteins, β2m has the unusual property of not requiring mutation or proteolysis to form fibrils. In vitro, however, the transition does not occur spontaneously at neutral pH but requires moderately acidic conditions and proper ionic strength or the presence of chaotropes (20–22). The formation of β2m fibrils at neutral pH occurs upon the addition of seeds and 2,2,2-trifluoroethanol, surfactants, or glycosaminoglycans (23, 24). Furthermore, temperature and pH conditions similar to those occurring upon inflammation in periarticular compartments enhance aggregation (25) and trigger fibril formation in the presence of collagen fibers (26) and heparin (27). The availability of several variants of the natural protein with different stabilities and amyloidogenic properties and the recent finding of a naturally occurring fibrillogenesis of the natural protein with different stabilities and amyloido-

disordered N- and C-terminal domains, the latter of which also contains a short, highly mobile, and unstructured C-terminal extension (34–36). Crystal and NMR structures have recently become available for the α-crystallin domain (37–39).

In addition to its potent ability to prevent amorphous protein aggregation of numerous target proteins, αB-crystallin also inhibits ordered amyloid fibril formation, e.g. by the Aβ peptide (22, 40, 41), apolipoprotein C-II (42), α-synuclein, the principal component of Lewy bodies in Parkinson disease (43), κ-casein (44–46), ataxin-3 (47), and β2m under acidic conditions (22, 48). Furthermore, αB-crystallin inhibits the aggregation of the serpin, α1-antichymotrypsin (49).

This study exploited the complementary insight obtainable from NMR spectroscopy and mass spectrometry (MS) into the early stages of β2m amyloid fibril formation and the nature of the interaction between αB-crystallin and β2m. For this purpose, we used wild-type β2m and R3Aβ2m under conditions of a convenient time scale for spectroscopic monitoring of processes that precede and lead to β2m aggregation, along with the effects of αB-crystallin on these processes. By use of 1H NMR spectroscopy, we determined that a nonspecific interaction occurs between the exposed surface of R3Aβ2m and αB-crys-

The αB-crystallin inhibits the aggregation of the serpin, α1-antichymotrypsin (49).

EXPERIMENTAL PROCEDURES

Materials—Recombinant β2m and R3Aβ2m were prepared as described previously (14, 19). The B subunit of α-crystallin was employed for all the experiments except for the DOSY measurements in which bovine eye lens α-crystallin (Sigma), composed of ~3:1 αA-αB-crystallin, was used. Recombinant human αB-crystallin was prepared as reported by Rekas et al. (43, 50). Protein concentrations were determined spectrophotometrically at 280 nm. For a 1-cm optical path length, A1% = 16.91 (β2m), 17.04 (R3Aβ2m), 6.93 (αB-crystallin), and 7.11 (α-crystallin). Protein integrity was monitored before and after experimentation by MS.

Thioflavin T Fluorescence Assay—Thioflavin T (ThT) fluorescence assays (440:490 nm excitation/emission) to monitor amyloid fibril formation of R3Aβ2m (500 μg/ml) were under-

NMR Spectroscopy—NMR spectra were obtained at 500.13 MHz with a Bruker Avance spectrometer on 0.1–0.9 mM protein samples dissolved in H2O/D2O (90:10 and 95:5 v/v) with 50–70 mM phosphate buffer, 50–100 mM NaCl, and pH * in the range 6.5–6.7 (where pH * is the pH meter reading without iso-

to effect correction). When necessary, samples were centrifuged and/or filtered using filters with a threshold of 0.22 μm (Millipore, Bedford, MA) or 0.02 μm (ANOTOP type, What-

man, Maidstone, UK). Studies were carried out at 310 K apart from a few experiments acquired at 313 K, which did not show
any NMR-detectable differences. The samples containing two proteins were prepared using αB-crystallin/R3Aβ2m at molar subunit ratios of 1:60, 1:15, 1:5, 1:2, and 1:1, and α-crystallin/β2m molar subunit ratios of 1.49 × 10^5, 1.95, 1.60, and 1.35. The β2m mutant and the stock α-crystallin solutions employed for measuring the signal attenuations were filtered using 0.02-μm microfilters with either lyophilized or un-lyophilized purified β2m proteins being used. The 1H two-dimensional total correlation spectroscopy (TOCSY) (52) NMR spectra were collected and processed as reported previously (19, 53). Based on the signal-to-noise ratio, amplitude errors typically ranged within ±5%, although much larger errors are to be expected for very weak resonances. Values of average accessible surface area of backbone amide and Cα hydrogen atoms were computed with the software available at the public domain bio-computation suite of EMBL, using a probe radius of 0.14 nm and the 20 deposited NMR structure coordinates of R3Aβ2m (19). Molecular structures were generated with InsightII (Accelrys Inc., San Diego), MOLMOL (54), and Swiss-Pdb Viewer software packages.

The diffusion coefficients of a 0.44 mm solution of wild-type β2m at 310 K (93.7 v/v H₂O/D₂O, 100 mM NaCl, 70 mM phosphate buffer, pH 6.6), as a function of the α-crystallin content (at the ratios reported above), were measured using the convolution-compensated two-dimensional double-stimulated echo diffusion ordered spectroscopy (DSTE DOSY) (55) pulse sequence to record matrices of 2048 (t2) by 80 points (t1). The gradient strength was varied linearly from 2 to 95% of its maximum value (61.1 G/cm). The lengths of the diffusion interval (200 ms) and diffusion gradient (2.5 ms) were optimized to obtain a substantial decay of the signal (95% loss, at least). The data analysis along t1 was performed using the Bruker DOSY software by imposing a single exponential fitting. Water suppression was achieved with the addition of an excitation sculpting WATERGATE module (56, 57).

Mass Spectrometry—MS measurements were performed on lyophilized wild-type β2m solutions dissolved in 100 mM ammonium acetate buffer, pH 6.82, at a final concentration of 120 μM, in the absence or in the presence of αB-crystallin. All sets of measurements were carried out using β2m solutions divided into two equal portions that were or were not filtered through 0.02-μm filters. For the first set of experiments, β2m was dissolved in a final volume of 500 μl at the concentrations reported above. The sample was divided into 2 aliquots. The 1st aliquot was the β2m control solution, and the second one was mixed with αB-crystallin in 100 mM ammonium acetate buffer, pH 6.82, at a final ratio of 30:1 β2m/αB-crystallin on a molar subunit basis.

In the second set of experiments, concentrated solutions of αB-crystallin in 100 mM ammonium acetate were added to 50 μl of an aged (20 days old) 120 μM β2m solution at a ratio of 30:1 β2m/αB-crystallin on a molar subunit basis. Before electrospray ionization (ESI), the samples were diluted to a final concentration of 20 μM in 100 mM ammonium acetate buffer, pH 6.82. This concentration was chosen because the physiological concentration of circulating β2m is 0.3 μM in healthy individuals but is elevated 20–60-fold in long term hemodialysis patients (58, 59).

The native MS experiments were performed using an ESI time-of-flight (TOF) LCT mass spectrometer equipped with a Z-spray nanoflow electrospray source (Micromass UK Ltd., Manchester, UK). Aliquots of 2 μl of the 20 μM protein solutions were infused into the mass spectrometer using an in-house pulled and gold-coated borosilicate glass needle. ESI-TOF-MS parameters were set as follows: capillary voltage, 1.2–1.5 kV; sample cone voltage, 120–150 V; extraction cone voltage, 50 V; source pressure, 6.7 mbar; TOF analyzer, 1.2 × 10^6 mbar. Spectra were recorded in the positive ion mode within the m/z range 500–10,000. For mass calibration, an aqueous 20 mg/ml CsI solution was used. The data were processed with MassLinks Version 4.1 Software (Micromass).

**RESULTS**

Amyloid Fibril Formation of R3Aβ2m Is Prevented by αB-crystallin—At the global level, amyloid fibril formation of R3Aβ2m was monitored via ThT binding. ThT fluorescence is markedly enhanced upon binding to the cross-β-sheet structure associated with the amyloid fibril structure. The ThT binding curves for an agitated R3Aβ2m solution, in the absence and presence of αB-crystallin, are shown in Fig. 1A. Interestingly, the ThT curves reproducibly displayed a biphasic profile, with an initial increase in fluorescence occurring after a lag phase of some 100 h and a plateau phase extending up to about 250 h, followed by a second much larger increase in fluorescence. Transmission electron microscopy images of R3Aβ2m, taken 400 h after the commencement of incubation, confirmed the extensive presence of fibrillar species (data not shown). The presence of increasing amounts of αB-crystallin led to the partial, and at higher concentrations, complete suppression of ThT fluorescence, due to the inhibition of fibril formation. When only partial suppression of ThT fluorescence increase was observed, i.e. at αB-crystallin/R3Aβ2m molar subunit ratios of 1:1000, 1:100, and 1:10, little if any inhibition of the first ThT fluorescence increment occurred, whereas the effect on the second ThT fluorescence increment was significant, implying that αB-crystallin preferentially interacts with the second phase of R3Aβ2m aggregation. Fig. 1B summarizes the amount of ThT fluorescence at the end of the experiments (400 h).

Assessment of the Stability of R3Aβ2m Solutions in the Presence and Absence of αB-crystallin as Monitored by 1H NMR Spectroscopy—The 1H NMR spectrum of R3Aβ2m has been assigned (19) thereby enabling investigation of alterations in chemical shifts and intensities of its resonances upon interaction with αB-crystallin. Because of its relatively rapid rate of aggregation and precipitation, the mutant exhibits a more convenient aggregation time scale for monitoring by NMR than the wild-type protein. Depending on the procedures undertaken after expression, purification, and sample preparation, the time scale for R3Aβ2m aggregation was variable. We found that the timing of the onset of precipitation and its extent depended on several factors. Besides the expected effects of protein concentration, pH, ionic strength, and temperature, the onset of precipitation was also affected by filtering the freshly prepared solutions and, specifically, by the exclusion size threshold of the filter (0.22 or 0.02 μm). Different precipitation patterns were also observed.
when the freshly prepared solutions were centrifuged rather than filtered. Other sources of variability were whether the protein had been subjected to lyophilization after final purification by dialysis, as well as omitting the lyophilization step so as to utilize only frozen aliquots of purified protein solutions. Solutions of lyophilized protein that were used unfiltered or subjected to filtration through a 0.22-μm filter prior to use are referred to as ordinary samples. Solutions of protein not subjected to lyophilization that had been filtered through the 0.02-μm filter are referred to as nonordinary samples.

As already reported (19), the 1H NMR spectrum of an ordinary solution of R3Aβ2m, acquired immediately after dissolution, shows the same pattern as observed with wild-type protein (12), which is consistent with the strong conformational similarities between the two species (19). Data were collected using clear solutions prepared without filtering and, sometimes, with mild centrifugation. After a lag phase that could last one or several days depending on the concentration and previous history of the sample, the NMR spectrum underwent changes caused by the concomitant processes of unfolding and aggregation followed by precipitation, as reported previously (19). The progress of unfolding is highlighted by gradual loss of the iso- and cross-peaks in the one- and two-dimensional TOCSY spectra (e.g. TOCSY) (Fig. 2A, insets). The same pattern was also observed in NMR spectra at half of this ratio, i.e. 0.5:1.0 αB-crystallin/ R3Aβ2m (data not shown). The spectra shown in Fig. 2A are interpreted as evidence for a weak interaction between R3Aβ2m and αB-crystallin that introduces a limited broadening of the R3Aβ2m resonances due to transient mutual association.

Quantification of the effects of αB-crystallin on R3Aβ2m spectra was undertaken by integration of selected resonances and cross-peaks in the one- and two-dimensional TOCSY spectra. Fig. 2B documents the time-dependent intensity change of an upfield-shifted methyl peak of R3Aβ2m (L23 δ1-CH3). Similar results are shown also for amide cross-peaks from the two-dimensional TOCSY fingerprint (Fig. 2, C and D). In the absence of αB-crystallin, after a lag phase of ~50–60 h, the resonance intensity decreased in a linear time-dependent manner and achieved about 80% loss over the time course of the experiment (~195 h). In the presence of αB-crystallin, for the same resonance, the loss of intensity was slowed down 3.4-fold, on average (Fig. 2B). No detectable lag phase also occurred in the presence of αB-crystallin. Overall, in the presence of αB-crystallin, a 6-fold reduction in the increase in intensity of the aliphatic resonances that are characteristic of unstructured peptides was measured, compared with the corresponding measurements in the absence of αB-crystallin under the same conditions (data not shown). Moreover, the samples with αB-crystallin present exhibited hardly detectable precipitation over the same time period compared with samples without αB-crystallin, where significant precipitation occurred.

In the absence of αB-crystallin, the timing for the lag phase and subsequent loss of NMR resonance intensity (Fig. 2, B and
FIGURE 2. A, 500 MHz 1H NMR spectra of an ordinary sample of R3Aβ2m (−0.9 mM, 310 K, pH 6.6) in the absence (lower trace) and presence (upper trace) of an equimolar amount of αβ-crystallin subunit. The only apparent consequence of the sHsp addition is a general broadening, with conservation of the spectral pattern of R3Aβ2m. The insets show a portion of the H−H connectivity (fingerprint) region of the corresponding two-dimensional TOCSY spectra. The boxed cross-peaks arise from the unstructured and flexible C-terminal extension of αβ-crystallin. Only this region of 12 amino acids is present in the spectrum by virtue of its fast local tumbling as opposed to the extremely slow overall motion of αβ-crystallin oligomer (−650 kDa) that causes the extreme broadening of other resonances (34–36, 60). Assignments are indicated for R3Aβ2m NH to α-CH cross-peaks (12, 19). Arrows mark visibly attenuated R3Aβ2m connectivities for Q69 and T86. B, time evolution of the resolved upfield-shifted methyl resonance (L23 δ(CH3)), which is diagnostic for the correctly folded species, of an ordinary sample of R3Aβ2m obtained from 500 MHz 1H one-dimensional NMR spectra, in the absence (circles) and in the presence of αβ-crystallin at equimolar (squares) or halved concentration (triangles) with respect to R3Aβ2m. The average rate of loss of signal intensity was (5.6 ± 0.5) × 10−3 h−1 in the absence of αβ-crystallin and (1.5 ± 0.5) × 10−3 or (1.8 ± 0.5) × 10−3 h−1 for 1:2 or 1:1 αβ-crystallin/R3Aβ2m mixtures, respectively. The errors on the rates were estimated by considering the deviations obtained when assessments were made using absolute resonance amplitudes rather than self-normalized values (53) to account for the uncertainties introduced by the scaling procedure. Generally speaking, such uncertainties arise when resonances with smaller linewidths overlap pre-existing signals, as well as from the general linewidth increase upon addition of αβ-crystallin. C, two-dimensional TOCSY fingerprint details obtained from an 82-h-old solution of R3Aβ2m (0.8 mM, 310 K, pH 6.6). The boxed cross-peaks represent the NH-αCH connectivities whose time evolution is displayed in D, respectively, from Phe-30, in the folded protein, and from an unknown residue, in the unfolded soluble species. D, time dependence of the two-dimensional TOCSY cross-peak amplitudes of C (Phe-30 = squares, unassigned = diamonds). In the absence of αβ-crystallin, the evolution of the resolved fingerprint cross-peaks showed typically a similar trend as that obtained by considering the evolution of the resolved methyl resonances in one-dimensional 1H spectra (6). Indeed, the slope of the decreasing Phe-30 cross-peak amplitude is −5.9 × 10−3 h−1, i.e. in absolute value the same, within the experimental error, as the slope of the unknown cross-peak intensity.

D) matches the lag phase and first rise of ThT fluorescence (Fig. 1). Following a latency period of 60 ± 20 h, the NMR signal loss is due to concomitant precipitation and formation of large soluble aggregates that are poorly observable or totally unobservable by NMR due to their very large linewidth. The total loss of the NMR signal occurs around 200–250 h, i.e. at the time as the first plateau in the ThT fluorescence data (Fig. 1). So, the initial increase of ThT fluorescence could coincide with the loss of the NMR intensity due to formation of large prefibrillar aggregates. We hypothesize that these prefibrillar aggregates, possibly containing some extent of intermolecular cross-β-structure, occur along nonamyloid and/or slow amyloid-forming pathways that are present at high β2m concentration and are enhanced by solution agitation (60).

Mapping R3Aβ2m-αβ-crystallin Interaction by NMR Spectroscopy—By contrast to the situation in the previous section with ordinary samples of R3Aβ2m, neither unfolding nor precipitation of R3Aβ2m was detected over an extended time...
period in nonordinary solutions prepared from un-lyophilized protein that had been filtered through 0.02-

\[ H_9262 \]

m filters. Thus, analysis of NMR spectra of these samples in the presence of \[ B \]-crystallin was not affected (over typical measurement time intervals) by systematic uncertainties due to the loss of soluble protein or the onset of resonances with unknown linewidth due to \[ R_3A \]

\[ H_9252 \]

2m aggregation. Therefore, measurements of the attenuation of resonances induced by \[ B \]-crystallin were undertaken using \[ B \]-crystallin/\[ R_3A \]

\[ H_9252 \]

2m at molar ratios suitable for NMR quantification. Attenuation of two-dimensional TOCSY fingerprint (\( H^{N}-H^{H} \)) cross-peaks was measured at 1:15 and 1:5 molar subunit ratios (data not shown). Qualitatively, identical patterns were observed at both ratios, and therefore, only the more precise results at the higher ratio (1:5) are reported (Fig. 3A).

To avoid a complex range of local mobilities, the quantitative attenuation analysis was restricted to connectivities from backbone hydrogens, i.e. a class of nuclei with relatively homogeneous motional properties (61). The fingerprint resolution of the TOCSY spectrum was not always sufficient to estimate individual cross-peak amplitudes, so only 67 out of 94 expected \( H^{N}-H^{H} \) connectivities could be integrated, which introduced discontinuity in the data of Fig. 3A. The interaction of \[ R_3A \]

\[ H_9252 \]

β2m with \[ B \]-crystallin is very dynamic due to a weak, nonspecific association between the two proteins. No chemical shift changes occurred, so the observed broadening of cross-peaks is ascribed

FIGURE 3. A, auto-scaled attenuation percentages (53) of \[ R_3A \]

\[ H_9252 \]

β2m fingerprint \( H^{N}-H^{H} \) TOCSY cross-peaks due to the presence of \[ B \]-crystallin. Data were obtained from integration of two-dimensional TOCSY spectra of 0.1 mM \[ R_3A \]

\[ H_9252 \]

β2m nonordinary samples, in the absence and presence of \[ B \]-crystallin at a molar ratio of 1:5 \[ B \]-crystallin/\[ R_3A \]

\[ H_9252 \]

2m. Because of resolution limits, only 67 out of 94 expected \( H^{N}-H^{H} \) connectivities could be integrated. The missing attenuation data were from Ile-1, Gln-2, Arg-12, Asn-17, Ser-20, Phe-22, Gly-29, Ser-33, Val-37, Leu-40, Arg-45, Lys-48, Glu-50, Ser-52, Leu-54, Ser-55, Ser-57, Lys-58, Trp-60, Ser-61, Leu-65, Thr-73, Glu-74, Tyr-78, His-84, Val-85, and Ser-88 cross-peaks, with no amide connectivity occurring for the prolyl residues at positions 5, 14, 32, 72, and 90 and the N-terminal methionine (M0). The attenuations exhibited a qualitatively identical pattern at a 1:15 molar ratio. However, greater attenuation (whose maximum was around 50%), which was more convenient for quantification, was observed at the higher \( B \)-crystallin ratio. The location of the \( \beta \)-strand segments according to the NMR solution structure (19) is given at the bottom of the figure. The attenuation trend along the \[ R_3A \]

\[ H_9252 \]

β2m sequence exhibited a clear analogy with the results of the experiments performed at an equimolar concentration on an ordinary sample (Fig. 2A).

B, schematic representation of \[ β2m \] secondary structure showing data, i.e. the attenuation levels of \[ R_3A \]

\[ H_9252 \]

β2m HN-H \( \alpha \) TOCSY connectivities due to \( B \)-crystallin interaction observed in nonordinary samples. Class I MHC heavy chain (backbone schematic and space filling) is shown in transparency. The crystal structure coordinates of the complex were used (Protein Data Bank code 3HLA (10)). A three-color code was employed to report \[ R_3A \]

\[ H_9252 \]

β2m experimental attenuation, as indicated explicitly. The residue positions for which no attenuation data could be measured is colored gray. The small schematic aside shows the \( \beta \)-strand naming scheme of \( β2m \). The drawing was prepared with PyMOL (DeLano Scientific LLC).
to the high effective mass that transiently is experienced by R3Aβ2m upon interaction with the chaperone.

At first glance, the profile of the fingerprint cross-peak attenuations appears to be spread over the whole R3Aβ2m sequence, except for the segment of residues 50–70 where the data set is sparse (Fig. 3B). The attenuations did not show any specific dependence on amino acid type, or any correlation with the secondary and tertiary structure, or a high surface exposure of the NH-C=H moieties. In detail, the largest attenuation values of HN-Hα correlations were mainly located in the inter-strand loop regions, namely the A-B and E-F loops and the C′ strand (five out of the nine most attenuated HN-Hα connectivities with an attenuation range of 40–50%). Therefore, the upper apical region of R3Aβ2m (Fig. 3B) may be preferentially involved in the interaction with αB-crystallin. Other backbone nuclei that experience comparable signal amplitude reductions are spread over the molecule. Fig. 3B provides a color-coded structural overview of the involvement of R3Aβ2m regions in interaction with αB-crystallin, as inferred from the NMR data, and shows that the contact patches are disseminated over the whole surface of R3Aβ2m. Among them, the regions of β2m in contact with the heavy chain of MHC-I (in transparent format) are readily apparent (10). Therefore, the weak, nonspecific interaction between R3Aβ2m and αB-crystallin also involves β2m regions with a propensity toward protection from the aqueous environment to limit their hydrophobic exposure.

Assessing the Effect of α-Crystallin on Wild-type β2m Aggregation by NMR Diffusion Measurements—We utilized NMR DOSY experiments (55) to measure the translational diffusion coefficient ($D_T$) of wild-type β2m as a function of α-crystallin concentration (Fig. 4). At 310 K, a 0.44 mM solution of wild-type β2m reproducibly exhibited a $D_T$ value of $1.83 \times 10^{-10}$ m$^2$/s that did not change appreciably over 5 days. Assuming a spherical model and using a mass-dependent protein density estimate (62), this value is consistent with an apparent mass of 19.9 kDa. The molecular mass of β2m is 11.9 kDa, implying that the protein undergoes aggregation processes of variable stoichiometry, i.e. as per the association equilibria previously proposed (19) with a dimer being the most populated species. In this respect, the conclusion generated by NMR is different from that obtained by MS (see below), which we ascribe to the 4-fold β2m concentration difference between the two experiments. Consistent with this, NMR diffusion measurements with freshly prepared β2m solutions at a much lower concentration (~0.1 mM) generated a $D_T$ value consistent with a mass of 12 kDa in agreement with the MS data on similar samples that showed a predominance of monomeric species.

The diffusion coefficient of a 0.44 mM solution of β2m was also determined by NMR following sequential additions of α-crystallin. The $D_T$ value did not alter for an α-crystallin/β2m ratio of $1.49 \times 10^2$, but at ratios of 1:95, 1:60, and 1:35, a $D_T$ value of $1.89 \times 10^{-10}$ m$^2$/s was obtained (Fig. 4), with a slight increase observed over 7 months after the last addition of α-crystallin to the solution. For the $D_T$ value measured after this last addition, the corresponding apparent mass of β2m was determined to be 17.9 kDa, a value that suggests a small but significant shift of the oligomerization equilibria toward lower mass species in the presence of α-crystallin, a shift that occurred to a greater extent over the long term (Fig. 4). A control TOCSY spectrum confirmed the stability of the β2m sample whereby α-crystallin preserved the native conformation of β2m as was also observed by NMR spectroscopy for R3Aβ2m in the presence of αB-crystallin (Fig. 3A).

TOCSY spectrum confirmed the stability of the β2m sample whereby α-crystallin preserved the native conformation of β2m as was also observed by NMR spectroscopy for R3Aβ2m in the presence of αB-crystallin (Fig. 3A).

**Time Course Oligomerization of Wild-type β2m as Monitored by Mass Spectrometry**—The alteration over 20 days in the mass spectrum of wild-type β2m after dissolving the protein in solution at pH 6.8 is reported in Fig. 5. The spectra show time-dependent changes in peak identities and intensities, in particular as new (oligomeric) species are formed in solution. Immediately after dissolution, mainly monomeric and some dimeric β2m species were detected, and after 4 h, oligomers up to hexamers were readily recognizable (data not shown). Later, species with higher mass appeared with increasing intensities, but their unambiguous identification could only be determined up to the octameric species. In Fig. 5, the development with time of increasing peak intensities, corresponding to an increase in

![Figure 4](http://www.jbc.org/)

**Figure 4.** A, wild-type β2m NMR DOSY spectra as a function of α-crystallin concentration and time. A limited window of the aliphatic region is shown to highlight the small but significant alteration in diffusion coefficient. The contour traces of five different DOSY experiments are superimposed at α-crystalin/β2m molar ratios of 0:1 (black), 1:4.9 $\times 10^2$ (green), 1:95 (blue), 1:35 (pink), and 1:35 after 230 days from the last α-crystallin addition (red). The substantial overlap of the traces at the two lowest and highest α-crystallin concentrations reflects the increase in translational diffusion coefficient value, $D_T$, of β2m below some 100-fold molar excess with respect to α-crystallin. The effect persists with solution aging. B, corresponding $D_T$ values of β2m as a function of α-crystallin concentration. The plotted data are the center of gravity measured at 1.6 ppm in the DOSY traces in A, with errors representing the maximum excursions above and below the determined value in the same traces. The red point refers to the measurement repeated 230 days after the first determination on the freshly prepared solution of 1:35 α-crystallin/β2m.
concentration of dimer, trimer, tetramer, and the onset of larger aggregated species, is consistent with the formation of oligomers in solution that may be the forerunners of nucleation and fibrillogenesis (19).

To characterize quantitatively the time course of $\beta_2$m oligomerization, peak areas in the mass spectra were evaluated. Because of the absence of an internal time-independent reference, raw data of areas corresponding to dimers or the sum of higher oligomers, i.e. from trimers to octamers, were normalized in each spectrum to the area corresponding to the overall sum of the $\beta_2$m peaks. This normalization procedure can be considered an internal scaling of the experimental data sets, and thus the intensity data obtained thereof are referred to as autoscaled areas. It should be noted that, despite mass conservation, comparisons among autoscaled areas may be affected by systematic errors due to the failure to detect very large oligomeric species (i.e. larger than octamers) in the mass spectra, especially for samples that were aged for a long period.

The time course of signal intensity for mass spectra of the $\beta_2$m dimer species and the higher oligomer pool, from trimers up to octamers, normalized over the total area sum, was quantified (Fig. 6). The percentage of dimer species present in solution increased steadily with time reaching 32% of the total 8 days after preparation. A similar trend was seen also for the sum of the other oligomers larger than dimers. In all cases, oligomer autoscaled areas were more intense in the unfiltered than the filtered samples with a 0.02-$\mu$m threshold (20-nm-filtered samples). The difference between 20-nm-filtered and unfiltered samples was greater for the oligomer pool compared with the dimeric species (Fig. 6).

Effect of $\alpha$B-crystallin on Wild-type $\beta_2$m Oligomerization as Monitored by Mass Spectrometry—To examine the effect of $\alpha$B-crystallin on $\beta_2$m aggregation, mixtures of wild-type $\beta_2$m and $\alpha$B-crystallin at molar subunit ratios of 30:1, 60:1, and 90:1 $\beta_2$m/$\alpha$B-crystallin were prepared and examined by MS. Two sets of experiments were performed. The first set involved the preparation of a fresh mixture of $\beta_2$m and $\alpha$B-crystallin in solution, followed by its monitoring over time by MS in tandem with a control solution that contained $\beta_2$m only. In the second set of experiments, $\alpha$B-crystallin was added to an aged (20-day-old) solution of $\beta_2$m, at the same molar ratios listed above.

Fig. 7 shows the mass spectrum with time of the solution containing simultaneously dissolved $\beta_2$m and $\alpha$B-crystallin (at a subunit molar ratio of 60:1) as monitored over 20 days (A–E)
**β₂-Microglobulin and αB-crystallin Interaction**

**FIGURE 6.** Time course of the autoscaled peak areas, *i.e.* expressed as percentage ratio with respect to the sum of all β2m signals, corresponding to the MS-detected β2m oligomers larger than dimers (A) and dimers (B). In particular, the bars report the sum of the autoscaled areas of all detected charge state signals for the higher oligomers, *i.e.* trimers, tetramers, etc., up to octamers (A), and the autoscaled areas for the dimers (B). The results for the ordinary (unfiltered) samples are indicated with solid bars, and those from the nonordinary samples (the samples filtered with a 0.02-μm threshold) are indicated with empty bars. The percentage of dimer species present in solution increases steadily with time reaching 32% of the total 8 days after preparation (B). A similar trend is seen also for the sum of the other oligomers larger than dimers (A). In all cases, oligomer autoscaled areas are more intense in the unfiltered than in the 20-nm-filtered samples. The difference between 20-nm-filtered and unfiltered samples is greater for the oligomer pool compared with the dimeric species. Because of the lack of an internal reference species with time-independent area, the raw area data corresponding to oligomers were normalized in each spectrum using the internal reference species with time-independent area, the raw area data (Fig. 9) or lysozyme, as representative of a typical medium-sized protein that has no chaperone ability. Independently of the molar subunit ratio (90 or 60 or 30:1 β2m/αB-crystallin to break down β2m oligomers in solution. In doing so, αB-crystallin converts β2m oligomers to monomers.

Similar MS experiments were conducted with two unrelated control proteins, GroEL, the well characterized molecular chaperone involved intimately with the folding of target proteins (for MS see ref. 63), and lysozyme, as representative of a typical medium-sized protein that has no chaperone ability. Independently of the molar subunit ratio (90 or 60 or 30:1 β2m/αB-crystallin) and preliminary solution filtering, none of the effects observed for αB-crystallin, *i.e.* prevention of β2m oligomerization and dissociation of β2m oligomers, was observed with GroEL (Fig. 9) or lysozyme (data not shown).

**DISCUSSION**

In this study, we have performed complementary NMR and MS experiments, addressing the very first stages of the β2m aggregation pathway and its inhibition due to interaction with the molecular chaperone αB-crystallin. We have previously shown that R3Aβ2m accumulates into fibrillar deposits in dialysis-related amyloidosis and exhibits a convenient aggregation time scale, while preserving all the features of the parent protein (19). Here, we have shown that R3Aβ2m forms fibrils at neutral pH upon agitation, a process that is inhibited by substoichiometric amounts of αB-crystallin.

We monitored the effect of αB-crystallin on R3Aβ2m in detail by NMR spectroscopy. First, we used ordinary solutions prepared from lyophilized R3Aβ2m, either unfiltered, centrifuged, or filtered through 0.22-μm filters. In the absence of α-crystallin, these samples exhibited a wide range of R3Aβ2m aggregation, precipitation, and unfolding behavior. Second, we used nonordinary R3Aβ2m solutions, *i.e.* solutions that were prepared under carefully controlled conditions, with un-lyophilized protein and filtered through 0.02-μm filters.

In ordinary samples of R3Aβ2m, aggregation was preceded by a lag phase that is consistent with the initial steps of the nucleated conformational conversion mechanism that R3Aβ2m undergoes during its aggregation (19). In the presence of αB-crystallin, the unfolding and aggregation of R3Aβ2m compared with the corresponding control spectrum of β2m after 20 days in the absence of αB-crystallin (F). As is readily evident from inspection of Fig. 7, the presence of αB-crystallin potently inhibits the development of oligomeric β2m species. After 1 day (Fig. 7B), the mass spectrum of the mixture was essentially dominated by peaks from the monomer and dimer species and was highly comparable with the corresponding “time 0” spectrum of β2m on its own. In the absence of αB-crystallin, the corresponding spectrum of a β2m solution contained peaks arising from oligomeric species up to hexamers (data not shown). Furthermore, contrary to the spectrum of a 20-day-old solution of β2m, the spectrum of a 20-day-old mixture of β2m and αB-crystallin did not exhibit any indication of conversion to large oligomeric β2m species.

Similar results were obtained using β2m/αB-crystallin molar subunit ratios of 30:1 and 90:1, with either unfiltered and filtered solutions. At a 90:1 β2m/αB-crystallin ratio, the sHsp was similarly effective at inhibiting the formation of β2m oligomers. Thus, MS demonstrates the impressive capacity of αB-crystallin, even at very low stoichiometric ratios, to completely inhibit the aggregation of wild-type β2m.

Fig. 8, A and B, shows the mass spectra of an aged (20-day-old) β2m solution at 1 and 20 days, respectively, after addition of αB-crystallin (at a 60:1 β2m/αB-crystallin ratio). The peaks corresponding to the oligomeric species that occur in aged β2m samples in the absence of αB-crystallin (Fig. 8C) are significantly decreased in its presence so that the mass spectrum of the aged β2m solution after 20 days of equilibration with αB-crystallin displays no oligomeric species and even a net decrease of the residual dimer peaks that were detected after a single day of equilibration with αB-crystallin. Importantly, the data in Fig. 8 provide compelling evidence for the capability of αB-crystallin to break down β2m oligomers in solution. In doing so, αB-crystallin converts β2m oligomers to monomers.

In this study, we have performed complementary NMR and MS experiments, addressing the very first stages of the β2m aggregation pathway and its inhibition due to interaction with the molecular chaperone αB-crystallin. We have previously shown that R3Aβ2m accumulates into fibrillar deposits in dialysis-related amyloidosis and exhibits a convenient aggregation time scale, while preserving all the features of the parent protein (19). Here, we have shown that R3Aβ2m forms fibrils at neutral pH upon agitation, a process that is inhibited by substoichiometric amounts of αB-crystallin.

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were always substantially reduced implying that αB-crystallin interacts with R3Aβ2m at the earliest stages of its aggregation pathway, as we have observed for the interaction of αB-crystallin with other fibril-forming proteins, e.g. apolipoprotein C-II (42), α-synuclein (43, 50), and ataxin-3 (47).

In ordinary samples of R3Aβ2m, the lag phase prior to precipitation was 1 to several days, depending on the sample's concentration and previous history. Removing solute heterogeneity required filtered, un-lyophilized samples. These nonordinary samples enabled reproducible data to be obtained on the very early stages of aggregation.

NMR studies did not detect any protein aggregation or unfolding in the nonordinary samples of R3Aβ2m, even over extended time intervals. There was no specific interaction site of R3Aβ2m with αB-crystallin because a general dipolar broadening, associated with dipole-dipole interactions between closely spaced 1H nuclei, was observed that affected the relaxation and increased the linewidths of R3Aβ2m resonances. It is therefore concluded that a large region of R3Aβ2m interacts with αB-crystallin. By evaluation of the backbone R3Aβ2m resonance attenuations, broadening was quantified, and the contacts between the two proteins were mapped (Fig. 3). β2m is a protein that in vivo, apart from the fibrillar deposits in pathological conditions, normally occurs in complex with the polymorphic moiety of MHC-I. The finding that αB-crystallin is in contact with the regions that are involved in MHC-I assembly suggests the involvement of hydrophobic interactions in driving the association between R3Aβ2m and αB-crystallin. In our NMR examination of the interaction between α-synuclein and αB-crystallin, κ-casein and α-crystallin, and ataxin-3 and αB-crystallin, we also observed nonspecific association between the two proteins and a large region of interaction on the target protein (43, 47, 50). Thus, it would seem that αB-crystallin has a general, essentially entropy-driven mechanism of interaction with fibril-forming target proteins that involves transient, low affinity interactions with a large solvent-exposed region of the latter proteins. Another important feature of this interaction is that as a consequence of the weak transient contacts, the turnover efficiency leads to an extensive sampling of the target protein surface by the sHsp. This prerogative of sHsps is crucial for preventing the aggregation of the target proteins and appears to be a specific feature of sHsps when compared with the interaction of other molecular chaperones with target proteins. Thus, the failure of GroEL to prevent β2m oligomerization suggests
that it is not as versatile as sHsps in interacting with target proteins under stress conditions, i.e. it only interacts with target proteins along their folding pathway (9). The same inability of lysozyme ensures that the efficient, albeit nonspecific, chaperone action of sHsps cannot be attributed to a generic protein interaction.

The low affinity of αB-crystallin for β2m is an advantage in terms of preventing β2m aggregation. Most likely, the very potent chaperone ability of αB-crystallin, i.e. its ability to inhibit β2m aggregation at very low ratios of αB-crystallin to β2m, arises from the combination of the fast rate of transient binding and dissociation of αB-crystallin to β2m coupled with the slow process of unfolding of β2m toward an amyloidogenic conformation. Furthermore, at any one time, there are very few β2m molecules undergoing this unfolding process. As a result of these factors, a single αB-crystallin molecule can potentially contact multiple β2m molecules to stabilize the native β2m conformation.

An important observation from the MS studies presented herein is that αB-crystallin is capable of dissociating soluble wild-type β2m oligomers into monomers in aged β2m solutions incubated in the absence of αB-crystallin. Thus, αB-crystallin can dissociate soluble β2m oligomers into smaller species and thereby remove them from the fibril-forming, off-folding

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6 C. Brockwell, H. Ecroyd, and J. A. Carver, unpublished results.
FIGURE 9. Mass spectra of a β2m and GroEL mixture (molar ratio of 60:1 β2m/GroEL), 20 min (A), 1 day (B), and 20 days (C) after mixing the proteins together are shown. The mass spectrum of the 20-day-old β2m control solution is also reported (D). The data show clearly that GroEL has no effect on the β2m oligomerization process, i.e. the spectra at 20 days from solution preparation in the presence (C) and in the absence (D) of GroEL are highly comparable. Similar results were observed also at molar ratios 30:1 and 90:1 β2m/GroEL (data not shown).

FIGURE 10. Schematic mechanism for the inhibition of amyloid fibril formation by αB-crystallin. During the formation of amyloid fibrils, target proteins (e.g. β2m), either folded or unfolded in their native state, can adopt a partially folded intermediate that is prone to aggregation to form prefibrillar oligomers. These can then sequester other intermediates to form the fibrillar species. αB-crystallin can interact with any of these species. Early along the aggregation pathway, i.e. at the intermediate or early oligomerization stage, αB-crystallin interacts transiently with the target protein and pushes it back into its monomeric, native conformation. It can also bind to the amyloid fibril to stabilize it and prevent its fragmentation and elongation.
pathway (9, 66, 67). As these oligomeric species are proposed to be cytotoxic, the ability of αB-crystallin to remove them is potentially of great importance in mitigating against their toxicity in diseases such as Alzheimer and Parkinson. Our results showing that αB-crystallin prevents the cellular toxicity of fibril-forming species, e.g. the Aβ peptide (41), is consistent with the ability of αB-crystallin to dissociate β2m oligomers.

Furthermore, our recent work has shown that αB-crystallin binds along the length and to the ends of fully formed amyloid fibrils of α-synuclein (50, 68), Aβ (69), and apolipoprotein C-II (70) to prevent fibril elongation and fragmentation. Fragmentation is recognized to be a major factor in fibril propagation due to it leading to secondary nucleation events and hence enhanced fibril formation (71).

The ability of αB-crystallin to interact with species at the various stages of the amyloid fibril-forming pathway is schematically represented in Fig. 10. The interaction at the early stages (with monomers and oligomers) is transient while binding of αB-crystallin to amyloid fibrils occurs (68–70). Thus, αB-crystallin is multifaceted in its ability to prevent fibril formation. Clearly, this multiplicity and plasticity in functionality provide αB-crystallin with enhanced capabilities in vivo in preventing the generation of toxic species arising from amyloid fibril formation and is consistent with the important role that αB-crystallin (and other sHsps) plays in minimizing protein aggregation.

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