

Amyloid- β protein oligomerization and the importance of tetramers and dodecamers in the aetiology of Alzheimer's disease

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In recent years, small protein oligomers have been implicated in the aetiology of a number of important amyloid diseases, such as type 2 diabetes, Parkinson's disease and Alzheimer's disease. As a consequence, research efforts are being directed away from traditional targets, such as amyloid plaques, and towards characterization of early oligomer states. Here we present a new analysis method, ion mobility coupled with mass spectrometry, for this challenging problem, which allows determination of *in vitro* oligomer distributions and the qualitative structure of each of the aggregates. We applied these methods to a number of the amyloid- β protein isoforms of A β 40 and A β 42 and showed that their oligomer-size distributions are very different. Our results are consistent with previous observations that A β 40 and A β 42 self-assemble via different pathways and provide a candidate in the A β 42 dodecamer for the primary toxic species in Alzheimer's disease.

Many diseases share the common trait of peptide-protein misfolding that leads to oligomerization and, eventually, formation of plaques of β -sheet structure. Prominent among these are type 2 diabetes¹, Parkinson's disease² and Alzheimer's disease^{3,4}. Of these, Alzheimer's disease is the leading cause of late-life dementia and is the focus of this paper. An increasing body of evidence links oligomerization of a ubiquitous peptide, the amyloid- β protein, to disease causation³⁻⁶. For this reason, elucidation of pathways of oligomer formation may be critical for the identification of therapeutic targets.

Many types of oligomeric amyloid- β assemblies have been described (for a review, see Lazo *et al.*⁷). Recently, Bitan *et al.*⁸⁻¹⁰ used photoinduced cross-linking of unmodified proteins (PICUP) to reveal that the 42-residue form of amyloid- β , A β 42, formed (A β 42)₅ and (A β 42)₆ oligomers ('paranuclei') that could oligomerize to form structures of higher order. A β 40 did not form paranuclei, but instead existed as a mixture of monomers, dimers, trimers and tetramers. Chen and Glabe¹¹, in contrast, used fluorescence and gel electrophoresis to determine oligomer states of amyloid- β refolded from denaturing solutions. They observed only A β 42 monomer and trimer bands, and no oligomers of A β 40. Differences such as these may exist because of the diverse experimental systems used to monitor amyloid- β self-association. Also, it has been argued that, in addition to the intrinsic potential of amyloid- β to traverse different assembly pathways, flaws in experimental design may have misled researchers in their quest to elucidate fully the amyloid- β oligomerization process¹². Hence there is significant uncertainty about amyloid- β oligomer states and their position and relevance to amyloid- β aggregation.

Results and discussion

We used a different, more direct, method to probe the amyloid- β oligomerization process: ion mobility coupled with mass spectrometry¹³⁻¹⁵. Details are given in the Methods section. Here the results for A β 40 are given as an example. The mass spectrum of A β 40 is shown in Fig. 1a and arrival time distributions (ATDs) of the $z/n = -2$ and $-5/2$ peaks ($z =$ total charge, $n =$ oligomer order) are given in Fig. 1b,c, respectively, for both high and low injection energies. High-resolution ¹³C isotope distribution measurements (insets in Fig. 1a) on the $z/n = -3$ peak and $z/n = -2$ peak indicate that the $z/n = -3$ peak is caused by monomer (i.e. $n = 1$, $\Delta(m/z) = 0.33$), whereas the $z/n = -2$ peak results from both monomer ($z = -2$, $n = 1$, $\Delta(m/z) = 0.5$) and dimer ($z = -4$, $n = 2$, $\Delta(m/z) = 0.25$). The ATD of the $z/n = -2$ peak (Fig. 1b) is bimodal (peaks at ~ 520 and ~ 660 μ s), which indicates that at least two non-interconverting species exist on the timescale of the experiment, in agreement with the mass spectrum.

Injection-energy studies (Fig. 1b) unambiguously identified the 660 μ s peak as the monomer ($z = -2$, $n = 1$) because it occurred only at high injection-energies, and thus resulted from oligomer dissociation, and because no additional peaks were observed at longer times. Only two peaks were observed in the ATD at lowest injection energy, so the peak at 520 μ s can be assigned to the dimer ($z = -4$, $n = 2$). Similar injection energy studies on the $z/n = -5/2$ peak identified both a dimer with $z = -5$ and $n = 2$ at 100 eV injection energies and a tetramer with $z = -10$, $n = 4$ at 23 eV injection energies, although the resolution of the experiment was insufficient to observe a clean

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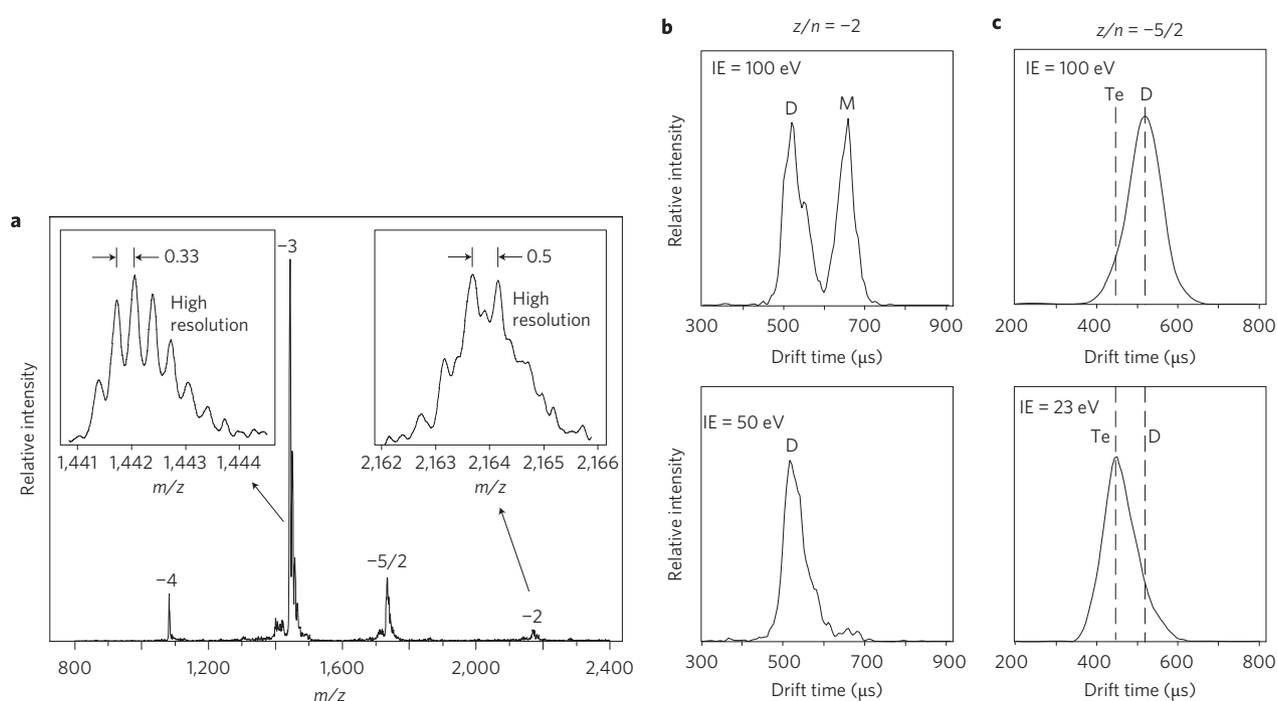


Figure 1 | Negative-ion mass spectrum and ATDs of A β 40. **a**, Mass spectrum from a 30 μ M solution at pH 7.4 labelled with z/n values (z = charge, n = oligomer order). The insets show high-resolution ^{13}C isotope patterns. The spacing of 0.33 for the $z/n = -3$ peak indicates that it is a monomer. The more complex pattern for $z/n = -2$ indicates that both monomer and dimer are present (dimer: $z = -4$, $n = 2$). **b,c**, ATDs at high and low injection energies (IE) for $z/n = -2$ (**b**) and $z/n = -5/2$ (**c**). The vertical dotted lines show the expected average experimental arrival times of the dimer and tetramer. M = monomer, D = dimer, Te = tetramer (see text).

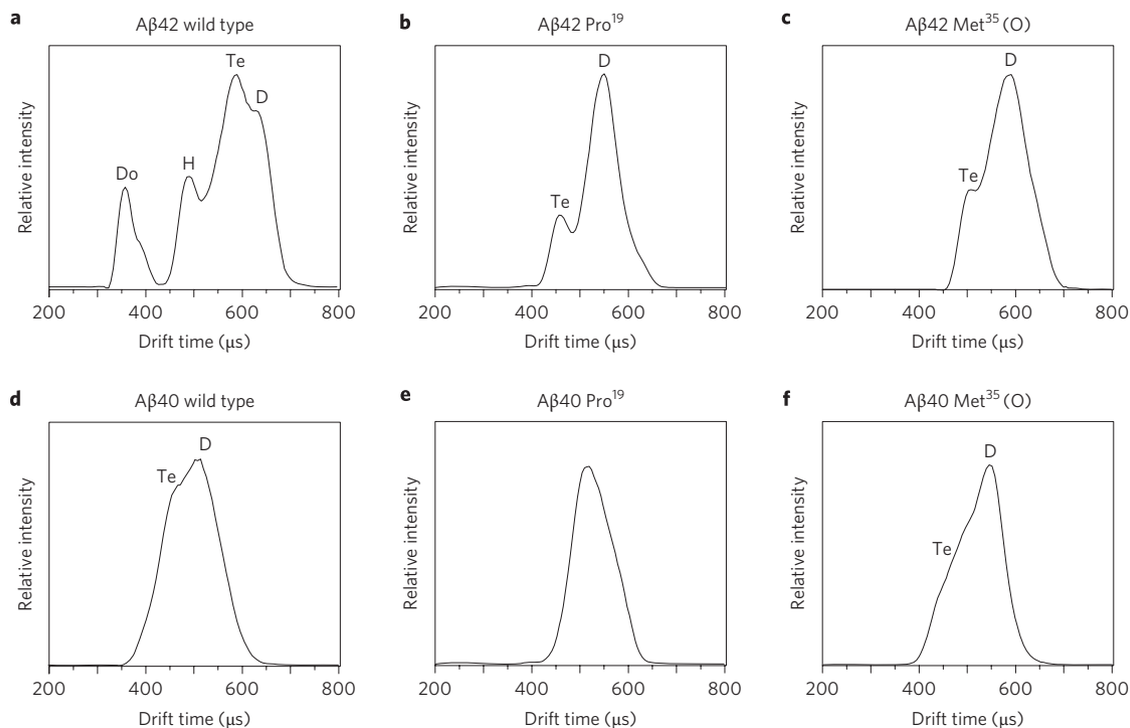


Figure 2 | Arrival time distributions. **a-f**, ATDs for $z/n = -5/2$ A β 42 (**a-c**) and A β 40 (**d-f**), showing the wild types (**a,d**), the alloforms Pro 19 (**b,e**) and Met $^{35}(\text{O})$ (**c,f**). The hexamer and dodecamer are only observed for A β 42 wild type and are thus implicated in the increased toxicity for this alloform. The distributions in **a** and **b** are from Bernstein *et al.*¹⁵ D = dimer, Te = tetramer, H = hexamer, Do = dodecamer.

bimodal ATD. (This issue is dealt with in more detail in Supplementary Information for the $z/n = -5/2$ ATD of both A β 40 and A β 42. A peak-width analysis is given based on the

diffusion equations for ions drifting in gases.) Hence, the A β 40 solution dominantly contained monomer, dimer and tetramer, with no trimer and larger oligomers observed.

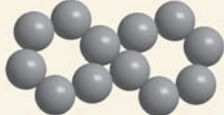
Similar experiments were carried out on the Pro¹⁹ and Met³⁵(O) alloforms of A β 40 and on the Met³⁵(O) alloform of A β 42 (details are given in Supplementary Information). These systems were chosen because PICUP experiments have shown that both substitutions are known to either reduce or eliminate fibril formation^{16,17} and restrict oligomer production to relatively small sizes according to Klein *et al.*¹⁸ and Bitan *et al.*¹⁹ In all cases the largest oligomer observed was the tetramer. The ATDs for the $z/n = -5/2$ peaks sprayed from solutions under identical conditions of all the systems considered here are summarized in Fig. 2. These data indicate that the ATD for A β 42 is much more complex than that for the other five systems. Injection-energy studies¹⁵ indicate that the peak at the shortest times in the A β 42 ATD is the dodecamer (A β 42)₁₂, and that the complex multipeak feature at longer times results from the dimer, tetramer and hexamer (see detailed discussion in Supplementary Information). We discuss this ATD shortly.

Ion mobility allows collision cross-sections (σ) of each oligomer to be measured accurately²⁰ (see Supplementary Information for the measured cross-sections). A plot of σ/n versus n (Supplementary Information) has two important features. Firstly, for all systems, as n increases σ/n decreases. This indicates that the apparent 'size' of each monomer unit within an oligomer decreases as oligomerization proceeds. This result is not unexpected, but emphasizes that oligomerization induces a certain amount of structural accommodation. One cause of this accommodation could be the self-selection of favoured structures as oligomerization proceeds. For example, both experiment and molecular dynamics modelling are consistent with a reasonably diverse ensemble of A β 42 monomer structures in solution²¹. Detailed analysis of the $-5/2$ ATD peak shapes of A β 42 (Supplementary Fig. S4) indicates that multiple structures are also present for dimers and tetramers of this alloform. However, the experimental linewidth significantly narrows for the A β 42 hexamer and dodecamer, which suggests a single structure for these two important species.

The second feature in the σ/n versus n graph is that A β 42 remains relatively larger (that is, accommodates less) than all other alloforms. These data are consistent with the results of intrinsic fluorescence studies²², which showed that the structure of the centre (Leu¹⁷–Ala²¹) of A β 42 was relatively stable compared with that of A β 40. In addition, these data offer insight into why A β 42 aggregation states and, by implication, the contribution of A β 42 to the aetiology of Alzheimer's disease are so different from those of other amyloid- β peptides. We obtained this insight by comparing measured cross-sections with those predicted by a simple structural model (for model details see Supplementary Information)²³.

In the model, the centre-to-centre distance (diameter) of an amyloid- β dimer constructed from quasispherical monomer units was adjusted to obtain the experimental dimer cross-section. This fitting accounts for a significant fraction of the monomer accommodation as n increases. Larger structures were then assembled using these calibrated dimer centre-to-centre distances. However, as mentioned above, a number of structures contribute to the experimental dimer cross-section, which indicates that the model cross-sections will be a bit too high when compared to oligomers of higher order and hence should be upper limits because no further accommodation is included. The results for A β 42 given in Table 1 are revealing. For example, the experimental tetramer cross-section falls between the model linear tetramer and a tetramer segment of the planar hexagon, and is much larger than the model planar square cross-section (more on this point shortly). The experimental hexamer cross-section is significantly smaller than the model linear cross-section and substantially larger than the closest-packed model cross-section, which rules these structures out. However, it is only moderately smaller than the model planar hexamer cross-section, a result consistent with the assumptions of the model and the tetramer structure, and so it supports this structure for (A β 42)₆.

Table 1 | Experimental and model cross-sections (Å²) for A β 42 oligomers.

Oligomer	Structure [†]	Cross-section	
		Model	Experiment
Dimer*		1,256	1,256
Tetramer		2,358	
		2,277	2,332
		2,135	
Hexamer		3,450	
		3,100	2,898
Dodecamer		2,578	
		5,824	
		4,562	4,308
			

*Model cross-section fit to experiment. [†]Limiting structures of high symmetry.

Similarly, the side-by-side planar hexagons can be ruled out as a structure for the dodecamer because their model cross-section is far larger than the experimental one, but the cross-section of the stacked hexagon model structure is in very good agreement with experiment. Hence, the experimental cross-sections are consistent with a quasiplanar hexagonal structure for the six-member 'paranucleus' and with stacked hexamers for the dodecamer.

An expanded version of the ATD for the $z/n = -5/2$ of A β 42 wild type is given in Fig. 3a with the 'structural' assignments shown for the various peaks. There are two additional important features. Firstly, the ATD goes to the baseline where the octamer would appear. Hence, A β 42 does not oligomerize via monomer or dimer addition past formation of the paranucleus (A β 42)₆, which suggests the mechanism in Fig. 3b occurs. Apparently, the structural stability of the paranucleus ring greatly reduces the monomer (dimer) addition rate as it accommodates homotypical growth. This critical feature indicates the potential importance of paranuclei in the aetiology of Alzheimer's disease¹⁰.

The second important feature in the A β 42 ATD in Fig. 3 relates to the complete absence of any signal at arrival times prior to that of dodecamer. This is unusual because, in most instances, a sloping ATD that extends to earlier times is observed, which indicates the presence of larger oligomers that are not well resolved. The absence of ion signal at these earlier times is consistent with the absence of octadecamers (18-mers) and indicates a lack of monomer or dimer addition to the dodecamer. The absence of an 18-mer that corresponds to ((A β 42)₆)₃ strongly suggests that addition of the third paranucleus is slow, and leads to accumulation of ((A β 42)₆)₂. The reason aggregation of A β 42 stops at the ((A β 42)₆)₂ dodecamer is not clear. Two plausible mechanisms are discussed in the Supplementary

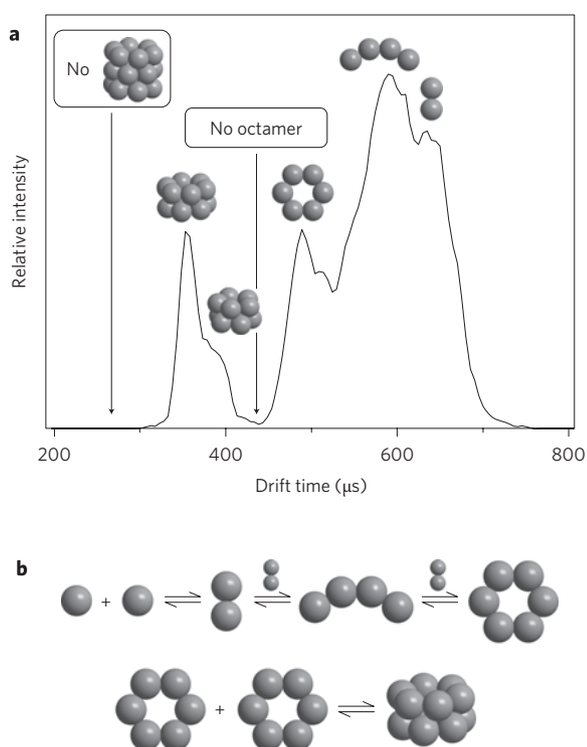


Figure 3 | A β 42 oligomer distributions. **a**, The ATDs of $z/n = -5/2$ with structural designations in place. **b**, A plausible mechanism for forming the oligomer distribution given by the ATD in **a**.

Information^{24,25}. Both mechanisms rationalize why the addition of a third hexamer is energetically unlikely and leads to a long-lived, but metastable, dodecamer. Apparently, this dodecamer eventually rearranges and rapidly adds monomer and forms fibrils. That A β 42 clogged the nanospray tips used to spray the sample solution (that is, rapidly formed very large oligomers and/or fibrils), whereas each of the other five systems reported here (see Fig. 2) sprayed beautifully for a minimum of several days up to more than a week without the tip clogging supports the rearrangement hypothesis. Aggregates of higher order (A β 42)_{*n*} would rapidly move out of the mobility–mass range of the ion mass spectroscopy instrument and their dispersal would make them difficult to detect with the limited sensitivity of this instrument.

To test this aggregation hypothesis we looked at the A β 42 positive-ion mass spectrum on a Q-TOF instrument specially built to observe high-mass aggregates under conditions in which A β 42 aggregation is maximized (see Supplementary Fig. S6)^{26,27}. A mound of ion intensity was observed that stretches from less than m/z 1,000 to m/z 8,000 and reflects a nearly continuous array of highly charged oligomers. Two sharp features at low m/z were identified as +4 and +3 residual monomers. This assignment was confirmed by tandem mass spectroscopy (MS/MS) (Supplementary Fig. S6 inset). The MS/MS result indicated that monomer loss dominated the dissociation process, which is common for collisional dissociation of large aggregates²⁸. When Pro¹⁹A β 42 was sprayed under identical conditions, a very clean monomer-dominated spectrum was observed (Supplementary Fig. S6), consistent with the fact that this alloform does not form fibrils¹⁷.

A clue as to why A β 42, but none of the other five systems, forms paranuclei is found in the relative cross-sections of the tetramers. This effect can be quantified using the model developed here. Each of the systems studied terminates at the tetramer, except A β 42. A normalized plot of the size of each tetramer experimental cross-section relative to several possible tetramer model structures is given in Fig. 4a. The model cross-section for the linear tetramer was taken as the benchmark because it is the largest possible structure

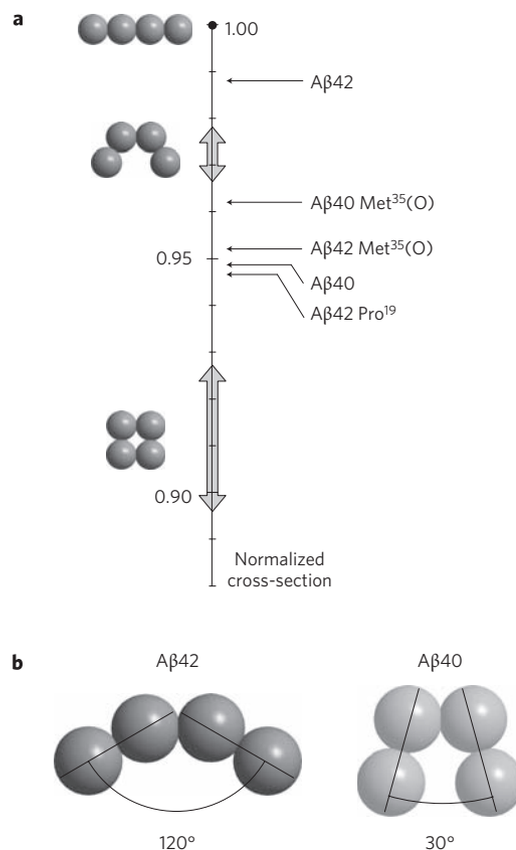


Figure 4 | The normalized cross-sections for the tetramers. **a**, Relative sizes of all tetramers reported herein (shown to right of vertical scale). The vertical scale is the relative cross-section of the model tetramers normalized to the linear structure. The broad arrows on the vertical axis show the spread of values for the modelled structures. A β 42 is a much more open tetramer than the other alloforms. **b**, The structures of the A β 42 and A β 40 tetramers taken from their location on the relative cross-section scale given in **a**, assuming the angle between dimer components is the determining parameter.

and all system sizes were normalized to that value. The results indicate that the A β 42 tetramer is much more ‘open’ than A β 40 or the other three alloforms shown. If the experimental cross-sections are fitted to a dimer–dimer angle the structures in Fig. 4b are obtained.

Another dimer could easily add to the ‘open’ A β 42 tetramer structure (Fig. 4b) to form the quasiplanar hexamer paranucleus (or a monomer could add to form a pentamer, which explains the stacked dipentamer shown in Fig. 3). However, the ‘closed’ A β 40 tetramers in Fig. 4b (and other alloforms) are not structurally disposed to the addition of either a monomer or a dimer and hence they form stable, long-lived tetramers. Our experimental data show the structure of the tetramer to be crucial in determining whether an amyloid- β alloform goes on to form paranuclei and dodecamers and subsequently contributes to the aetiology of Alzheimer’s disease. Hence the (perhaps small) folding differences in the various monomers are amplified by aggregation, and at the tetramer dramatic differences occur. From a different perspective, the ‘dimer–dimer’ interface is different from the ‘monomer–monomer’ interface, and perhaps the A β 42 dimer is uniquely suited to form an open tetramer and the other systems are not. The tetramer is small enough to hope that serious modelling can be done to extract the factors at the molecular level in these systems that drive such crucial structural differences. These studies are underway.

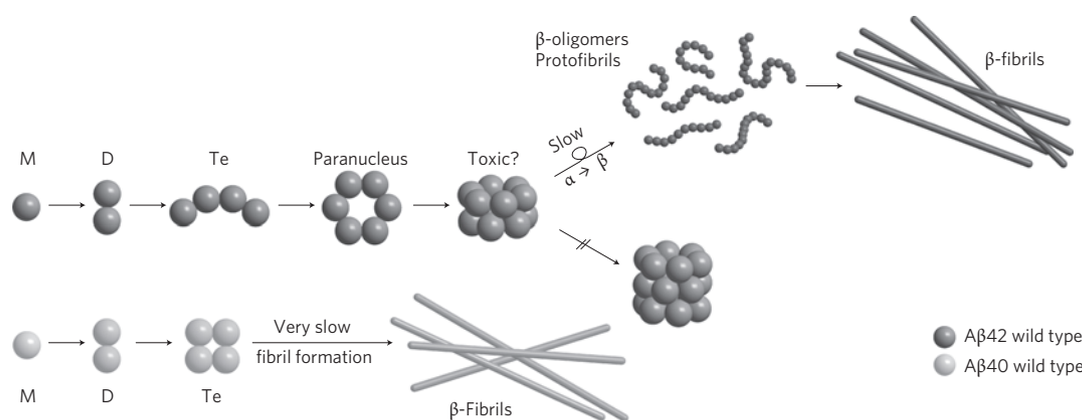


Figure 5 | Mechanism of oligomerization and eventual fibril formation for A β 42 and for A β 40. For A β 40 the key structure is the tetramer that resists further monomer or dimer addition. In A β 42 an ‘open’ tetramer promotes the formation of the planar hexamer (paranucleus) and the stacked dodecamer, which resists further reaction. For A β 40 the tetramer apparently eventually forms fibrils (observed by others), but these were not observed in our experiments. For A β 42 a rate-limiting slow α - to β -sheet transformation may occur for the dodecamer, but this was not explicitly observed in our experiments. Fibril formation was indirectly observed through macroscopic clogging of the spray tips used for A β 42.

A global scheme of amyloid- β oligomerization can be deduced by combining the data presented here with previous published data^{9,10} (Fig. 5). For A β 42 a receptive structure in the tetramer allows further dimer addition and paranucleus formation with a planar hexagonal structure. Two paranuclei combine to form a stacked paranuclear dimer, which resists further transformation. Eventually, ((A β 42)₆)₂ rearranges to form a new structure to which monomer rapidly adds to form large β -sheet oligomer assemblies. A β 40, however, forms a more compact tetramer that firmly resists further addition and hence forms no paranuclei. A similar fate to that of A β 40 befalls the Pro¹⁹ and Met³⁵(O) alloforms of both A β 42 and A β 40. Although some of these systems eventually do form fibrils, they appear to do so by a mechanism that bypasses paranucleus formation¹⁰.

In related work Barghorn *et al.*²⁹ isolated a species they called ‘globulomer’ from A β 42 using a quite complex protocol. This species is soluble and has a molecular weight between 60 and 100 kDa. It is stable against dissociation for several days and only slowly self-aggregates to form larger oligomers in the absence of A β 42 monomer. Application of proteases truncated the N-terminus of the globulomer A β 42 monomer constituents up to about residue F20, which indicates that the C-terminal hydrophobic residues are in the centre of the globulomer. This result is consistent with our view of both the hexamer and the dodecamer complexes, in which the hydrophobic tails are envisioned as occupying the centre of the structures (see Supplementary Fig. S4 and the cartoon in Baumketner *et al.*²¹). Barghorn *et al.* suggest the globulomer is made up of 12 A β 42 molecules, but their evidence for this is not conclusive²⁹. They also assume that the globulomer is formed by an independent mechanism than through A β 42 fibrils, a suggestion at odds with the mechanism we give in Fig. 5 and one that would be difficult to rationalize with a kinetic model (which they did not present).

One additional important point is worth making. Lesné *et al.*⁶ used amyloid precursor protein transgenic Tg2576 mice to demonstrate that memory defects occur in ‘middle-aged’ mice (6–14 months) without neuronal loss, but in older mice abundant neuritic plaques that contain A β 42 were observed. What they observed in these middle-aged mice was an extracellular accumulation of a ~56 kDa soluble amyloid assembly, A β *56. These authors proposed that this 56 kDa assembly impairs memory by an unknown mechanism independent of either plaque formation or neuronal loss. Their observations are supported by recent results of Cheng *et al.*⁵, who also used transgenic mice and the ‘Arctic’ (E22G) mutant alloform of A β 42, which implicate A β *56 rather

than fibrils as a determinant of Alzheimer’s disease. Here we show that A β 42 forms *in vitro* a dodecamer composed of stacked hexamer paranuclei. This dodecamer is the terminal species observed in our experiments, and has a mass of ~55.2 kDa, which suggests it is the soluble assembly that Lesné *et al.* observed⁶. This assembly has not yet rearranged into a β -sheet structure and may possibly be the key neurotoxic assembly in Alzheimer’s disease.

Methods

Materials. All peptides were synthesized on an automated peptide synthesizer (Applied Biosystems model 430A) using methods based on 9-fluorenylmethoxycarbonyl (Fmoc) essentially as described by Lomakin *et al.*³⁰ For A β 42 an Fmoc-L-Ala-2-chlorotrityl resin was used (0.51 mmol g⁻¹, Biosearch Technologies). Lyophilized peptide was dissolved at a concentration of 4 mg ml⁻¹ in deionized water. To this solution, 0.006 times the volume of 1 N NaOH was added, followed by filtered 50 mM ammonium acetate solution (pH 7.4) to reduce the peptide concentration to about 2 mg ml⁻¹. The solution was sonicated for one minute and then 100 μ l was filtered through a 10,000 AMU Gel Filtration G-10 Macro Spin Column purchased from Nest Group. The filters were prehydrated and washed in 25 mM ammonium acetate pH 7.4 according to the manufacturer’s suggestions for desalting and buffer exchange to ensure the pH of the peptide solutions was maintained at 7.4, and cleaned of interfering salts for mass-spectrometry purposes. The sample was centrifuged for four minutes at 2,000g and the filtrate was collected and used immediately. The final concentration of peptide ranged from 30 to 40 μ M (determined by amino-acid analysis) in 25 mM ammonium acetate and adjusted to pH 7.4.

For A β 42 it was necessary to carry out the experiments as quickly as possible. Although the procedure outlined here led to stable solutions, the signal still rapidly diminished over a period of about a day because of aggregation of the peptide (eventual clogging of the spray tips confirmed this). No new features were observed in either the mass spectrum or ATD for A β 42 for as long as observations were possible.

The other peptides were much better behaved with solutions stable up to a week (several weeks for Pro¹⁹A β 40 and Pro¹⁹A β 42). The mass spectra and ATDs did not change over this time period for any of these peptides, which indicates that hexamers and dodecamers were not formed in measurable quantities. Hexamers and dodecamers were exclusively formed in A β 42 wild type. All experiments were completely reproducible over many trials over a period of more than a year. Circular dichroism spectra were taken of all samples to ensure consistency from batch to batch. In all cases the spectra were identical for a given alloform over the complete timescale of the project, which indicates excellent sample consistency.

Mass spectrometry. Standard mass spectra and ion mobility experiments were accomplished on an instrument built in-house and composed of a nano-electrospray ionization (N-ESI) source, an ion funnel, a temperature-controlled drift cell and a quadrupole mass filter followed by an electron multiplier for ion detection³¹. The high-resolution ¹³C isotope distributions for each peak in the mass spectra were obtained on a Q-TOF mass spectrometer (Micromass UK) equipped with a N-ESI source.

In the ion-mobility measurements, the ions were stored at the end of the ion funnel and then pulsed into the drift cell filled with 5 torr of helium gas and drawn through the cell under the influence of a weak electric field. The ions’ injection

energy into the drift cell was varied from 0 to 150 eV. At low injection voltages, the ions were gently pulsed into the mobility cell and only needed a few 'cooling' collisions to reach thermal equilibrium with the buffer gas. At high injection voltages, the larger collision energy led to internal excitation of the ions before cooling and equilibrium occurred. This transient internal excitation can lead to annealing, that is partial or complete isomerization, to give the most stable conformers or dissociation of dimers and oligomers of higher order¹⁵. The ions exited the drift cell, passed through a quadrupole mass filter and were detected as a function of time to produce an ATD. The arrival time is simply related to the mobility, which in turn is inversely proportional to the collision cross-section³². All factors in these equations²⁰ are either universal constants or parameters that were accurately measured in the experiment. Hence accurate ($\pm 1\%$) collision cross-sections were obtained.

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Author contributions

S.L.B., N.D.L., G.B., D.B.T., J.-E.S. and M.T.B. conceived and designed the experiments, S.L.B. and T.W. carried out the experiments, N.F.D. and T.W. designed and performed the modelling, S.L.B., B.T.F. and C.V.R. measured the highly aggregated mass spectra, M.M.C. synthesized the peptides and M.T.B. wrote the paper.

Additional information

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BIOANALYTICAL CHEMISTRY

Protein oligomers frozen in time

The size and shape of amyloid- β protein assemblies have been studied using electrospray-ionization ion-mobility mass spectrometry, and the protein tetramers and dodecamers have been identified as an important oligomerization state in the development of neurodegenerative disease.

David E. Clemmer and Stephen J. Valentine

When Fenn and co-workers recorded early electrospray ionization (ESI) mass spectra of biopolymers, they observed what are now characteristic charge-state distributions as well as the formation of non-covalent complexes that persisted into the gas phase¹. Writing in *Nature Chemistry*², Bernstein *et al.* describe the remarkable differences in the distributions of oligomers formed by different protein isoforms — different forms of the same protein that may be produced by small changes in the encoding gene — of amyloid- β (A β) protein using electrospray-ionization ion-mobility mass spectrometry. Their results address the first steps of β -fibril assembly — a phenomenon that may provide a physical explanation for the first steps in the formation of large fibrils, which are found as signatures in patients with neurodegenerative diseases such as Alzheimer's.

Bernstein *et al.* suggest that oligomerization of these proteins proceeds through key oligomer sizes and structures that regulate further growth and assembly. Some polypeptides form relatively small complexes with stable, closed structures that resist further growth. One sequence, the wild-type A β 42, is dramatically different. The small complexes that are formed are not self-terminating and thus can coalesce, ultimately leading to fibril formation.

The ability to characterize protein structures in the gas phase is quite remarkable in and of itself. In the mid 1990s, MacLafferty and co-workers trapped different charge states of cytochrome *c* ions in a vacuum for extended times (up to 30 minutes)³. On exposure to very dilute deuterated solvents, variations in the exchange rates and levels for the different charge states were found. The fact that heteroatom hydrogens within the proteins are protected from deuterium exchange to different degrees requires that the proteins have defined structures, even in the complete absence of solvent. Moreover, some individual charge states exchanged at different rates, such that a single peak in the mass spectrum would split into multiple

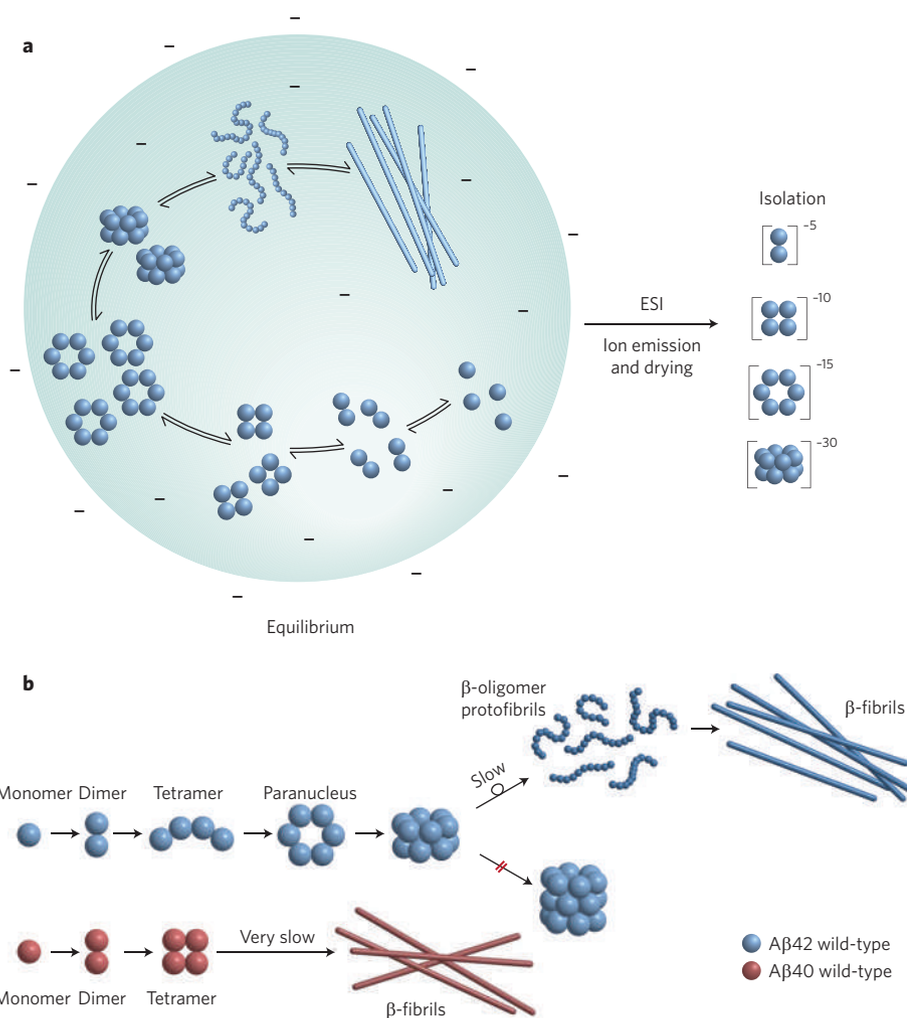


Figure 1 | Electrospray ionization allows protein oligomers in solution to be frozen out and their size and shape to be determined by ion-mobility spectrometry. **a**, Cartoon of an electrospray droplet that contains a distribution of the A β 42-protein aggregation states that exist in equilibrium in solution. Once electrosprayed into the gas phase the various oligomer species are isolated from one another and the population of states is fixed. Separations of oligomers based on mass-to-charge (which is similar for all the oligomers) can be further refined by shape-to-charge. **b**, All the amyloid- β proteins studied (for example, the wild-type A β 40) assemble to form dimers, tetramers and ultimately fibrils, but the open geometry of the wild-type A β 42 tetramer allows it to also form hexamers (paranucleus) and dodecamers.

peaks over time — with differing degrees of deuteration. This result indicates that the nature of populations of conformations

in the gas phase is very different than in solution. That is, in the gas phase, the different conformations do not seem to be

in equilibrium with one another, even after extended trapping times. This work led to a flurry of interest in the structures of entirely 'naked' proteins^{4,5}. Is it possible that different conformational states become trapped as the distribution of ions emerges from the ESI droplets — freezing the distribution in time? To the extent that this is true (Fig. 1a) studies of these structures in the gas phase can provide insight about populations of conformations that are impossible to observe in solution because of the rapid equilibrium between states.

Bernstein *et al.* inject short pulses of the long-lived gas-phase structures that are formed from the ESI droplet into a helium buffer gas and expose the ions to a weak electric field. As the 'packet' of ions drifts through the gas, different species separate according to their mobilities. For example, oligomers with compact conformations undergo fewer collisions than those with extended conformations, and thus will have higher mobilities. Similarly, oligomers with higher charge states experience a greater drift force than those with lower charge states, and thus will also have higher mobilities. Therefore, it is possible to distinguish between the different oligomers based on variations in shape-to-charge even when they have the same mass-to-charge ratio. Moreover, because the interactions with the simple gas are well understood it is possible to calculate ion mobilities for trial geometries and by comparison with experiment, obtain insight into the shapes of the oligomers^{6,7}.

When this approach is applied to the *in vitro* distributions of oligomers of the A β 40 and A β 42 isoforms the results are remarkable. The ion-mobility distribution

for the wild-type A β 42 shows features that are assigned to the dimer, tetramer, hexamer and dodecamer, whereas the A β 40 isoform forms only the dimer and tetramer (Fig. 1b). Similar studies of other A β 42 alloforms — in which single amino acid residues have been changed — also show oligomer formation; however, the growth again seems to be restricted at the tetramer. The abrupt absence of larger oligomers for the A β 40 isoform and other A β 42 alloforms suggests that the tetrameric form has a key role in the inability of these systems to assemble into larger complexes — and thus is an important species in the development of disease. Bernstein *et al.* propose a compact tetrameric form for these less toxic proteins. It is the closed nature of this structure that resists rapid accumulation of protein units. On the other hand, the wild-type A β 42 tetramer favours an open conformation that can easily incorporate another dimer to form a hexamer ring, and these rings can stack together to form dodecamers. The authors speculate that over time the dodecamer may rearrange such that further assembly can occur, leading to the large fibrils seen in the advanced stages of neurodegenerative diseases.

The potential understanding of the early steps of fibril formation provided by this study are profound and a number of interesting issues are raised. For example, one can imagine using this insight to develop interventions that are designed to arrest oligomer growth at a specific size. Although the present application of these technologies is tremendously inspiring, there is a clear need to improve

the techniques. For example, a method of selecting and trapping the different oligomers based on differences in shape would make it possible to carry out dissociation studies that could address the stability of different oligomer sizes and also corroborate assignments. Furthermore, one imagines that within each of the oligomer sizes there are likely to be populations of stable structures. As the ability to resolve species with this technology improves it will become possible to examine more of the details of how different conformations of assemblies grow. Finally, these results present an extraordinary opportunity for theory. With the additional constraint of the measured mobility it will be interesting to see what calculations can say about the details of structures that terminate or allow fibril growth. □

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