Interaction of partially denatured insulin with a DSPC floating lipid bilayer†

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The carefully controlled permeability of cellular membranes to biological molecules is key to life. In degenerative diseases associated with protein misfolding and aggregation, protein molecules or their aggregates are believed to permeate these barriers and threaten membrane integrity. We used neutron reflectivity to study the interaction of insulin, a model amyloidogenic protein, with a DSPC floating lipid bilayer. Structural changes consistent with protein partitioning to the membrane interior and adsorption to a gel phase model lipid bilayer were observed under conditions where the native fold of the protein is significantly destabilised. We propose that the perturbation of the membrane by misfolded proteins involves long term occupation of the membrane by these proteins, rather than transient perforation events.

I. Introduction

A wide range of incurable diseases (including Alzheimer’s, Parkinson’s and type II diabetes) are associated with the misfolding of proteins. Common to the pathology of each of these diseases are amyloid aggregates. The current consensus is that the toxicity mechanisms causing the onset and progression in these diseases arise from cell damage and death caused by the presence of oligomeric forms of amyloidogenic proteins.†,‡,†,‡ These toxic species carry out their activity at the cell surface and pathways as to how this leads to cell death are currently the focus of much investigation. Although a number of protein receptor systems have been postulated,§ studies using simple vesicular systems† have shown that only the phospholipid component is necessary for the proteins’ ability to traverse the membrane bilayer.

This membrane damage must therefore be an adsorption-driven process, with structural rearrangement either preceding or following the adsorption and incorporation of oligomers or monomers respectively. Either way, surface properties,§,† including those of lipids§ have been shown to play an active role in the unfolding and misfolding process, accelerating fibrillisation rates in comparison to those in the bulk. At interfaces there are significant physical and chemical differences in the local conditions to those in bulk solutions. FTIR§ and neutron reflectivity§ studies have previously shown how interfacial processes can alter conformation and structure in adsorbed protein films.†1,12

To study the nature of the interaction of proteins at a membrane surface, we carried out an experiment in which we directly probed the interaction of a fibrill-forming protein with a model lipid membrane using neutron reflectometry. The protein we used was insulin, which has frequently been used as a model system for protein fibrillisation due to its wide availability and textbook amyloid behaviour: well characterised insulin fibrils are readily formed by heating under acidic conditions. Insulin amyloids have also been isolated in vivo, although the extent of their involvement in pathology is probably limited to injection sites in diabetic patients, while that improper storage of insulin hormone can compound this problem and lead to reduced efficacy and shelf-life.

Neutron reflectometry (NR) is a structural probe which is extremely well suited to the study of model lipid membranes and biological processes occurring at these interfaces.16,17 This technique is capable of the high resolution determination of the layer thicknesses and composition, by exploiting the large differences in scattering between H2O and D2O, multiple measurements of the same sample can provide a unique solution to the interfacial structure for adsorbed films. In order to investigate the initial phase of penetration, conditions which accelerate the formation of intermediate states in the misfolding of insulin18 while still allowing the sample to remain unchanged during the long measurement times required for NR. A floating lipid membrane was chosen as a model membrane because a planar geometry is needed for NR and, with this type of membrane mimic, the bilayer is decoupled from the interactions with the underlying silicon substrate by a thin layer of water.
II. Experimental

Sample preparation

Insulin from bovine pancreas, 1-(trimethoxysilyl)-propylacrylate, guanidine hydrochloride, glycine, AAPH, triethylamine and all solvents used were obtained from Sigma-Aldrich. Lipids and modified deuterated lipids as shown in Fig. 1 were supplied by Avanti Polar Lipids, Alabaster, TN, USA. All materials were used without further purification except for toluene, which was dried using CaH₂, and distilled at reduced pressure to remove water, this was stored in glassware with a young’s tap to minimise moisture uptake. D₂O was obtained from Flurochem, Glossop, UK.

Grafted lipopolymer films on silicon were produced following the method described by Hughes et al.19 with some variation; firstly silicon blocks (Crystran, Poole, UK) were cleaned using the RCA-1 process (70 °C for 10 minutes in 5:1:1 H₂O:H₂O₂ 30% : NH₃ 30%) in order to remove any surface organic contaminants, then the blocks were dried using nitrogen gas and heated to 50 °C in the reaction vessel under vacuum, in order to remove any adsorbed water, an essential step before the deposition of the silane.

The blocks were then sealed in a PTFE reaction dish with a 25 mM solution of 1-(trimethoxysilyl)-propylacrylate (Sigma-Aldrich, UK) in dry toluene (Sigma-Aldrich, UK; HPLC grade, further dried using calcium hydride and distilled) with 21.5 mM of triethylamine (Sigma-Aldrich, UK). A period of 18 hours incubation in the solution was used for the formation of the self-assembled monolayer (SAM) after which the blocks were cleaned using the method described by Hughes et al.19

For the final Langmuir–Schaeffer dip the pressure-jump measured for the samples is presented in Fig. S6 (ESI†) for the 0 M, 1.5 M and 3 M Gdn-HCl. Using the software Sasview a Guinier fit was used to extract the radius of gyration of Insulin.

Small angle X-ray scattering (SAXS)

SAXS data was collected on beamline B21 at the Diamond light source, the solution scattering was measured using the automated sampling system available at B21, this system fills a cuvette, collects the data and then automatically cleans the cuvette prior to the next solution. The solutions were made in the same way as the for the NR experiments insulin at 1 mg ml⁻¹ with 50 mM glycine buffer acidified to pH 2.3 using concentrated hydrochloric acid. The data for the samples is presented in Fig. S6 (ESI†) for the 0 M, 1.5 M and 3 M Gdn-HCl. Using the software Sasview a Guinier fit was used to extract the radius of gyration of Insulin.

Dynamic light scattering

Dynamic light scattering (DLS) measurements were performed on a Brookhaven Instruments 200SM laser light scattering goniometer using a HeNe (125 mW, 633 nm) laser. Single five minute exposure scans over a period of 13 hours were performed and particle sizes were estimated using the CONTIN multiple pass method of data analysis at an angle of 90°.

Neutron reflectometry and data fitting

Neutron reflectivity measurements were performed on the D17 instrument at the Institut Laue-Langevin (ILL) and it was used...
in its standard time-of-flight mode using a 4% resolution in Q. Two angles (0.7° and 2.5°) were used to cover the Q-range to 0.2 Å⁻¹. Normalisation of the detector was performed by running a cuvette of light water (H₂O) and the data sets from the two angles were joined using the Cosmos software. Light water contrasts were run for significantly longer than those in D₂O buffers, but due to the lower scattering in water it was not possible to obtain the same quality of statistics due to time constraints. Bilayers were prepared 3 days before the experiment and stored in sealed sample cells, which had been cleaned in Decon-90 (Decon Laboratories Ltd) surfactant and washed thoroughly with ultra pure water (18 MΩ cm) and alcohol in water at room temperature, until the beginning of the experiment this ensured that the possibility of bacterial contamination was greatly reduced.

All buffers and protein solutions were prepared freshly using Millipore ultra pure water and freshly opened D₂O (Flurochem). All buffers were acidified using concentrated HCl (Sigma Aldrich) to pH 2.3. Bovine insulin (Sigma Aldrich, I5500) solutions were made at a concentration of 1 mg ml⁻¹ and were mixed gently using a pipette so as not to cause bubbles (as bubbling can cause interfacial aggregation).

Different contrasts for each measurement were prepared using D₂O and H₂O solutions of insulin at 1 mg ml⁻¹ with 50 mM glycine buffer acidified to pH 2.3 using conc. HCl and the specified Gdn·HCl concentration. The sample was allowed an incubation period of 5 hours to equilibrate with protein or change of contrast with the protein present before the reflectivity measurement.

The data were model fitted and where possible simultaneously co-refined with both contrasts using the Motofit reflectivity fitting package for Igor Pro. The final fitting of the sample at 3.5 M concentration used models where the substrate and the covalently attached monolayer were totally constrained from the initial structure. Remaining layer parameters then employed the simultaneous co-refinement used for solving the initial structure. Mutation and recombination values for the fit search were optimized after.

In addition to the fitting of the structure of the bilayer in 3.5 M Gdn·HCl, null hypothesis fitting was also performed. For this, reflectivity curves for the membrane both before and after exposure to the protein were simultaneously fit. This fitting procedure resulted in a significantly increased value of χ².

Reflectivity curves were fit by two different methods: simultaneously fitting in both contrasts using fairly unconstrained models initially and a genetic algorithm to find solutions. This was performed alongside fitting of individual contrasts using much tighter physical constraints. Both methods gave fairly similar results with some interesting differences due to which regions each contrast was most sensitive to. The a priori knowledge of the molecular detail of the bilayer allows some of these differences to be explained. All fits were derived from using the values from Hughes et al. as an initial condition but with a modification to headgroup SLD to account for isotopic substitution in this region.

The fitting process requires physical constraints both to make sense and to limit available parameter space. It should be noted that in the case of lipids the splitting of the molecule into head and tail regions requires further assumptions on where the neutrons ‘see’ the head to finish and tail to begin. This however is an unavoidable simplification due to the slab fitting of molecules, which are not actually slabs of homogeneous scattering material.

Values for the SLD limits of buffers containing Gdn·HCl were calculated by weighing a 100 ml volumetric flask, adding the necessary mass of the salt, weighing again and then filling to 100 ml before a further weighing. Using the known density of water it was possible to obtain measured values for the density of the salt and from this calculate accurately the SLD for both the dry salt and the buffer.

III. Results and discussion

We investigated whether the adsorption of a model amyloidogenic protein to a lipid membrane is altered by destabilisation of the native state as the sampling of many molecular conformations by non-native monomers means there is considerably more flexibility for conformational rearrangement. During these measurements we have focused on the unfolding event immediately prior to formation of oligomers by using the denaturant guanidine hydrochloride (Gdn·HCl). We used three solution conditions in the measurement – insulin in pH 2.3 glycine buffer without guanidine, 1 M and 3.5 M Gdn·HCl. The choice of pH was made so that our results are comparable with the acidic conditions used by others to study insulin fibrilisation. The choice of the other two conditions were 1 M as a high-salt, natively folded condition and the 3.5 M point because Gdn·HCl titration of insulin shows significant changes between around 2.75 M and 3.25 M as revealed by circular dichroism. This transition to an expanded monomeric state is an on-pathway step in the formation of the amyloid state. Another effect of Gdn·HCl at lower concentrations is to significantly reduce the Debye length in solution, as it is a salt. This is important to highlight as at low concentrations the enhanced screening of electrostatic forces can actually stabilize proteins rather than destabilize them.

DLS measurements confirmed that under these solution/temperature conditions and timescales no detectable bulk protein aggregation occurs. This provides evidence that there are no aggregates larger than the instrument resolution limit of 30 nm (see ESI†). While this does not rule out the presence of oligomeric species it is clear that formation of fibrils does not occur over the experimentally relevant timescale. The radius of gyration of the protein as measured by SAXS at 0 M Gdn·HCl was shown to be 11.7 Å ± 0.5 Å, sadly this was only possible for the 0 M dataset, as the change in electron density upon the addition of Gdn·HCl reduces the overall effective contrast in the system. This result is in good agreement to the size of the insulin monomer measured in 20% acetic acid solution. This previous study found that the effect of guanidine hydrochloride to the solution caused expansion of the $R_g$ under solution conditions of 20% acetic acid and disassociation of hexameric insulin as a function of Gdn·HCl concentration therefore we expect that insulin was monomeric under all solution conditions used. The lack of detectable signal in the SAXS results for
the samples containing Gdn-HCl can be seen as a secondary validation of the result seen by DLS – that initially there are no aggregates present as with rising molecular weight the signal from SAXS is increased. It should be noted that the previous study by Ahmad was conducted at higher concentration and this facilitated their measurement.

It has been suggested that line defects in bilayers have the potential to act as sites which allow partitioning of the protein into the membrane structure from solution through the interaction of exposed hydrophobic residues facilitating irreversible adsorption events. This should be accelerated when defects are long-lived and the protein is in a state that allows significant freedom to explore conformational space. For this reason lipid bilayers in the gel phase were used to provide a model system for this.

Initial reflectivity measurements for the measured bilayers in 1 M and 3.5 M Gdn-HCl at pH 2.3 showed profiles similar to those shown in ref. 19, however the initial structure for the sample without the salt present was significantly different to the rest. A satisfactory fit for the sample without salt was not possible using simultaneous co-refinement, however separate fits of the individual contrasts converged on structural solutions with features consistent with a lipid bilayer with a significantly larger reservoir next to the grafted substrate (not shown). In this case the usefulness of the data as a control is therefore of limited use to the study.

The reason for bilayer instability in the 0 M Gdn-HCl case is most likely due to electrostatic effects caused by the low pH. The pK_a of the phosphatidylcholine headgroup is quoted as being between 1 and 2 in ref. 23 and as this pH is approached a fraction of the lipids will become charged. This charging will not only cause like-charge repulsive effects but also possibly affect headgroup hydrogen bonding. The combination of repulsion between headgroups, possible perturbation of H-bonding and also layer–layer repulsion due to the grafted layer becoming similarly charged. The importance of the headgroup–headgroup hydrogen bonding network is illustrated by phase-separation observed in PC membranes under conditions of moderate pH perturbation. The salt effect on the membrane is consistent with the effect of monovalent salts on membrane properties being to increase membrane rigidity with rising salt concentration. This increased membrane rigidity, combined with enhanced screening of electrostatic effects is what we would attribute this stability to, although the exact contribution of each is impossible to determine based on these measurements. The estimated Debye screening length in the salt-free conditions was 13.6 Å in the 50 mM buffer, compared to 1.6 Å and 3 Å for the samples with Gdn-HCl present. This increase in the Debye screening length in solution means that while the samples in guanidine have screening lengths smaller than the headgroup size, the sample without would therefore not screen nearest neighbour electrostatic effects.

After incubation of the floating lipid bilayer with insulin in 1 M and 3.5 M Gdn-HCl buffers for 5 hours no change was observable in the 1 M sample with identical reflectivities for both contrasts, the 3.5 M sample showed significant change in reflectivity together with a shift of the fringe minimum to lower Q. The comparison of the lack of change in conditions where the main effect of the added guanidine is that of charge screening (1 M) to those where the protein is partially denatured (3.5 M) shows that the protein stability in solution has a strong effect on its propensity to adsorb to a lipid bilayer.

Fig. 3 highlights the lack of any structural changes for the sample in 1 M Gdn-HCl, where the main action of the low denaturant concentration is that of Debye screening. This is juxtaposed against what we observe for the sample in 3.5 M guanidine (Fig. 4); clear changes in both contrasts are visible. The shift in the position of critical reflection for the sample incubated in the D_2O solution was due to an incomplete
exchange of the sample cell. The adsorption occurring at the outer solution–lipid interface has an increase in roughness consistent with the radius of gyration of monomeric insulin plus the original roughness of the bilayer. A significant change in the SLD of the bilayer is also visible in these conditions. This could be explained by incorporation of protein into the bilayer core, probably in a non-native conformation. Similar observations were recently made by van Maarschalkerweerd\textsuperscript{26} where α-synuclein was found to partition to a lipidic phase resulting in lipid–protein co-aggregates in the bulk. Interestingly, this study also found that the lipids in these aggregates also had a “gel like character”.

An upper limit for protein concentration in the lipid bilayer for the sample in 3.5 M Gdn\textsubscript{HCl} was estimated to be 35% at the outer leaflet and 11.6% in the membrane interior using the approach described in ref. 27 and the value from the change in scattering length density. This method properly accounts for the changes in scattering length density for the protein due to the effects of H–D exchange in both side-groups and backbone due to the solution conditions. Comparing the two conditions where Gdn\textsubscript{HCl} is present, it is clear that in these conditions, where the Debye length is comparable, partitioning to the membrane is favoured at higher Gdn\textsubscript{HCl} concentration where the protein has greater conformational flexibility. This is favourable because the protein has increased disorder and flexibility to explore the different conformational arrangements, allowing exposure of hydrophobic residues, which stabilise the membrane-bound state.

As the negative result from the DLS rules out bulk aggregation the presence of an adsorbed protein film is consistent with the results from ref. 26 and 28 and has additional significance in the context of phosphatidylcholine derived stabilisers for insulin\textsuperscript{29} where binding in the headgroup region has also been observed.

### IV. Conclusions

A comparison between the behaviour of insulin in the 1 M and 3.5 M Gdn\textsubscript{HCl} shows that, in conditions which partially unfold the protein, there appears to be evidence that the protein inserts itself in the bilayer and adsorbs at the outer leaflet of the membrane. This change occurs throughout the membrane in both the tail and head regions there is also an increased interfacial roughness at the top sample surface. This accumulation of material could explain in part the acceleration of fibrillation at membrane surfaces observed by others. There is no evidence as to whether this is caused by individual proteins or aggregates, although complementary measurements found no evidence for aggregates in the bulk solution phase.

The alternative hypothesis of adsorption to the membrane surface and a destabilisation of the membrane leading to an increased amount of membrane edges would also lead to a similar rise in scattering length density in the aliphatic region. However, both of these results would be consistent with significant alterations of membrane properties due to incubation with the partially folded protein.

The lack of adsorption in 1 M Gdn\textsubscript{HCl} is consistent with what would be expected based on previous results such as the X-ray reflectivity study by Perriman\textsuperscript{30} and QCM-D measurements by Glasmastar.\textsuperscript{31}

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