Experimental report

| Proposal: | TEST | -2358 | | | Council: 4/2014 | | |
|---------------------------------|--------|---|----------------|----------------|------------------------|------------|--|
| Title: | Natura | latural and synthetic lipid bilayers: interactions with bio-molecules(AmB and IAPP) | | | | | |
| Research area: | | | | | | | |
| This proposal is a new proposal | | | | | | | |
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| Samples: IAPF AmE | 3 | | | | | | |
| Instrument | | | Requested days | Allocated days | From | То | |
| D17 | | | 3 | 3 | 21/07/2014 | 24/07/2014 | |
| Abstract: | | | | | | | |

Experimental report for proposals TEST2358, TEST2335, 8-02-742 and 8-02-688

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These beam times were part of a large study including Neutron reflectometry (NR), SANS and laboratory techniques to investigate the interaction of IAPP fragments with POPC bilayers. IAPP is the peptide involved in β -cell depletion leading to the development of Diabetes Mellitus type 2. In order to understand the interaction of the peptide with cell membrane and its consequences, we used SANS an NR. The peptides we choose are depicted table 1: IAPP human peptide (hIAPP₁₋₃₇) has but splitted in two halves: the N-terminal half (hIAPP₁₋₂₀) is suspected to interact with lipids and the C-terminal half (hIAPP₂₃₋₃₇) is very aggregation-prone and has been shown to fold into a beta-sipper and form the spine of the amyloid fibers. The rat homologs (rIAPP₁₋₃₇, rIAPP₁₋₂₀ and rIAPP₂₃₋₃₇) have been used as non-amyloidogenic and non-toxic controls. Human and rat peptides differ by 6 positions which only one is in the C-terminal half. The chimeric peptide rIAPP₁₋₃₇R18H associating the human N-terminal half and the rat C-terminal half has also been analysed.



The ability of each peptide to permeate a POPC bilayer has been measured on the PSCM set up for Electrophysiology on DIB. Figure 1 shows the results of these measurements. They show that hIAPP₁₋₃₇ is able to permeate the model membrane but hIAPP₁₋₂₀ and rIAPP₁₋₃₇R18H are much more efficient than the full length human peptide. An interpretation of these results is that the N-terminal half of the human peptide is responsible for lipid bilayer disturbance and that the amyloidogenicity of the C-terminal half inhibits it. The other peptides: the 3 rat peptides and the human C-terminal half, does not have any effect on the POPC bilayer conductance.



Structural characterization of the POPC bilayer disturbed by the peptides has been performed by NR with the POPC membrane deposited on the surface of a Silicon crystal on top of the buffer chamber for eventual aggregates to be discarded from the interface by gravity. Buffers with various D₂O content were used for contrast variation and the curves were simultaneously fitted using Aurora software. The SLD profiles resulting from these fittings are shown by figure 2. We observed a dramatic lipid depletion following the injection of hIAPP₁₋₂₀ nearby the bilayer. A less pronounced effect was observed with hIAPP₁₋₃₇ which apparently inserts through the first leaflet of the bilayer, mainly amongst the headgroups, as well as rIAPP₁₋₃₇R18H which insertion in the first leaflet is accompanied with lipid depletion. These results confirm that the N-terminal half, in particular the hitidine18, is crucial for the interaction of the peptide with a POPC bilayer. They also strongly suggest that lipid depletion could be the mechanism of bilayer permeation.



This hypothesis raises the question of the fate of the lipids removed from the bilayer. The structural analysis of aggregates of hIAPP₁₋₃₇ formed in presence or absence of POPC vesicles have been carried out by contrast-variation SANS (D22). The results are presented on figure 3 as the square root of the intensity extrapolated at Q=0 angle as a function of D₂O content of the buffer. This analysis shows that the presence of POPC vesicles during hIAPP₁₋₃₇ aggregation changes the aggregates match point from 39% to 26% D2O, which would correspond to an incorporation of three lipid molecules per peptide within the aggregates. Furthermore, the Intensity scattered by the lipid-peptides co-aggregates never reaches 0, which suggest that within the aggregates, the lipids and the peptides segregate in two phases rather than mixing intimately.



All together, these results enable us to propose a cytotoxicity mechanism in which the N-terminal half of $hIAPP_{1-37}$ would insert in the lipid bilayer and extract a large amount of lipids from it leading to membrane permeation. The amyloid aggregation could be an inhibitory mechanism selected as a rescue system to delay the onset of the disease.

A publication presenting these results and interpretation is in preparation.