An ion channel containing model membrane: structural determination by magnetic contrast neutron reflectometry

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Summary

To many biophysical characterisation techniques biological membranes appear as two-dimensional structures with details of their third dimension hidden within a 5 nm profile. Probing this structure requires methods able to discriminate multiple layers a few Ångstroms thick. Given sufficient resolution, neutron methods can provide the required discrimination between different biochemical components especially when selective deuteration is employed. We have used state-of-the-art neutron reflection methods, with resolution enhancement via magnetic contrast variation to study an oriented model membrane system. The model is based on the *Escherichia coli* outer membrane protein OmpF fixed to a gold surface via an engineered cysteine residue. Below the gold is buried a magnetic metal layer which, in a magnetic field, displays different scattering strengths to spin-up and spin-down neutrons. This provides two independent datasets from a single biological sample. Simultaneous fitting of the two datasets significantly refines the resulting model. A β-mercaptoethanol (βME) passivating surface, applied to the gold to prevent protein denaturation, is resolved for the first time as a 7.6 ± 1.3 Å thick layer, demonstrating the improved resolution and confirming that this layer remains after OmpF assembly. The thiolipid monolayer (35.3 ± 0.5 Å), assembled around the OmpF is determined and finally a fluid DMPC layer is added (total lipid thickness 58.7 ± 0.9 Å). The dimensions of trimeric OmpF in isolation (53.6 ± 2.5 Å), lipid monolayer (57.5 ± 0.4 Å) and lipid bilayer (58.7 ± 0.9 Å) are precisely determined showing little variation.


Introduction

Currently, artificial planar layers of biological molecules on solid substrates are useful in two distinct research fields, the study of biological membranes\(^1\) and the formation of protein arrays.\(^2\) The biological membrane, a dynamic composition of lipid and protein which encloses most cells and organelles, is a structure still to yield its innermost secrets. The thirty five year old fluid mosaic model is the basis for our current thinking\(^3\) but the detailed information on the distribution of phospho-, glyco- and sphingolipids plus integral, surface and cytoskeletal proteins is still emerging.\(^4\) For example, lipid raft structures have recently revealed complex structure-function relationships within protein-lipid interactions.\(^4\) For many structural analysis tools both the living cell and its established model, the lipid vesicle, do not provide sufficiently oriented samples. Lipid monolayers at the air-water interface are useful\(^5, 6\) but flat substrates provide the possibility of assembling oriented bilayers.\(^1\) The substrate may be silicon, glass, gold, etc and particularly for the latter, there may be a chemical link to create a tethered lipid bilayer.\(^7, 8\) Detailed knowledge of immobilised biological layers also has biotechnological uses since the need for the immobilisation of protein arrays on substrates for use in drug discovery and diagnostics is growing strongly.\(^2, 9-11\) Furthermore, there is an increasing number of applications of nanoscale surfaces upon which cells will behave as though in complex tissues and thus provide a relevant \textit{in vitro} model of cellular processes.\(^12-14 15\)

Many physical methods are used to probe such layers. Microscopy, including electron, fluorescence\(^16\) and atomic force,\(^17, 18\) can provide detailed data on the structure and/or dynamics of the 2D distribution of membrane components, whilst spectroscopic techniques such as ATR-FTIR\(^19\) can probe structural and dynamic aspects of molecular organisation. Surface plasmon resonance,\(^20\)
quartz crystal microbalance,\textsuperscript{21} dual polarisation interferometry\textsuperscript{22} and ellipsometry\textsuperscript{23} are sensitive to mass or refractive index based effects but can also yield estimates of other parameters such as thickness and elasticity. Electrical impedance measurements can be used on conducting surfaces\textsuperscript{24} whilst solid state NMR can use stacked planar bilayers on glass plates to determine the structure of membrane proteins, particularly the orientation of transmembrane helices.\textsuperscript{19}

To determine the distribution of individual membrane components through the bilayer, methods that exploit the neutron’s ability to discriminate between hydrogen and deuterium are particularly powerful.\textsuperscript{25} Small angle neutron scattering (SANS) has been used to examine the molecular distribution of membrane proteins within lipid bilayers and detergent micelles\textsuperscript{26,27} whilst neutron crystallography has revealed detergent\textsuperscript{28} and glycolipid distributions in membrane protein crystals.\textsuperscript{29} Specular neutron reflection (NR) can probe planar membranes and is the only method capable of fully defining the distribution of individual components throughout the profile (z axis) of a single bilayer.\textsuperscript{30} We have recently determined the structure of a protein-reconstituted bilayer that contained the bacterial exotoxin, \(\alpha\)-hemolysin, in high concentrations and were able to locate the protein within the bilayer with a precision of \(~1 \text{ Å}\) along the surface normal.\textsuperscript{31} These approaches are providing increasingly detailed pictures of membranes and may allow investigation of other structures such as bacterial peptidoglycan.

The system investigated in this paper was initially developed as a model of the \textit{Escherichia coli} outer membrane for biological studies.\textsuperscript{32, 33} Later it was realised that self-assembled monolayers of gram-negative outer membrane proteins (Omp) on surfaces can also play a role in biotechnology.\textsuperscript{14} The protein used, OmpF from \textit{Escherichia coli}, is one of a large group of \(\beta\)-barrel membrane proteins and this one in particular forms extremely stable 16-stranded \(\beta\)-barrels which
are resistant to proteases, urea, guanidine hydrochloride, sodium dodecyl sulphate (SDS) and heat
denaturation.\textsuperscript{32} The high resolution crystal structure 2OMF\textsuperscript{34} shows it to have an asymmetric
shape, resembling a cylinder with a length and diameter \textit{circa} 60 and 30 Å, respectively. These
monomers always associate into trimers with a flat base, on the periplasmic face, approximated by
an equilateral triangle with a base length of 80 Å. The resilience of OmpF permits its
immobilisation on surfaces and subsequent characterisation without detergent in aqueous buffer
since exposure of the hydrophobic protein surface does not cause denaturation. This unusual
stability of the protein enables the step-by-step assembly of a stable, electrically sealing bilayer
membrane suitable for sensing applications.\textsuperscript{32, 35} Such layers need to be electrically tight\textsuperscript{36} and
measurement of water content within the hydrophobic centre of the membrane may be good metric
for successful fabrication as it may correlate with membrane resistance to ion transfer. Neutrons
are a prime tool to assess water content because of their isotopic sensitivity.

Here we use a modified OmpF containing a cysteine residue (OmpF-E183C) by which it
can bind to a gold surface. To cover the high-energy gold surface entirely after surface ligation of
the protein, we form contiguous membranes by back filling the area between the immobilised
protein with thio-lipids that also attach through gold-sulphur bonds.\textsuperscript{32} Subsequently, the resulting
OmpF/self-assembled monolayer (SAM) system is terminated by precipitation of a terminal layer
of 1,2-Dimyristoyl-sn-Glycero-3-Phosphocholine (DMPC) using the method previously reported
by McGillivray \textit{et al.}\textsuperscript{8}

The final architecture consists of a protein/lipid bilayer membrane that has its inner
monolayer leaflet bonded to the gold film and is arranged around OmpF-E183C trimers also
covalently bound to the gold surface (see Fig. 1). The self-assembly of the complex membrane is
monitored sequentially through the deposition process with the thickness and composition of the layers determined at each step. As we show, this general organisation of the membrane is confirmed by NR employing a combination of isotopic ($^2$H vs. $^1$H) and magnetic contrast neutron reflectometry (MCNR). This uses polarised (up or down spin) neutrons to provide two independent data sets from a single membrane (see Materials and Methods for details). Compared to conventional reflectometry,\textsuperscript{37} this method results in significantly improved structural resolution and moreover enables determination of structures with a high degree of compositional complexity. To quantify these advantages, we have assessed the performance of the method by a Monte-Carlo-based resampling technique\textsuperscript{15, 38, 39} The results demonstrate the application of MCNR for the determination of the molecular distribution across a complex biomembrane with Ångstrom resolution.

**Results**

**Resolving the β-mercaptoethanol layer**

Gold pre-treatment with small hydrophilic thiol containing molecules is a common method to passivate the high energy surface\textsuperscript{1, 40} and significantly reduces protein denaturation and non-specific binding to gold surfaces.\textsuperscript{32, 41} Figure 2a shows NR data for the ‘up’ and ‘down’ spin states after βME deposition onto a gold coated substrate (thickness, $d = 186.3 \pm 0.3$ Å) with an iron/nickel magnetic sub-layer ($d = 75.6 \pm 0.1$ Å) in contact with D$_2$O buffer. Because of the large difference of the neutron scattering length density (nSLD) values for the iron/nickel layer probed by the two spin states, the two NR spectra are different with the reflectivity for the spin ‘up’ state about an order of magnitude larger than that for the ‘down’ state. The derived nSLD profiles (Fig. 2b continuous lines) provide excellent fits to the data and are the result of the simultaneous
refinement of data from five different samples (3 hydrogenous and 2 deuterated βME) each with two spin states. Data was collected on one of the hydrogenous βME samples with two different H2O/D2O contrast conditions resulting in a total, using both spin states, of twelve different datasets. The data were fitted with the βME thickness constrained but with sub-layer (gold etc) properties and βME surface coverage allowed to vary between different samples. A βME thickness of 8.2 ± 0.6 Å with a nSLD of 3.07 ± 0.15 x 10⁻⁶ Å⁻² was obtained for the sample shown in Fig. 2, corresponding to a βME surface coverage of 57%. Table 1 lists the volume fraction (Vf) of the βME in the layer calculated for the various hydrogenous samples against a D2O buffer.

As gold is readily contaminated upon exposure to air it was necessary to confirm that the layer resulted from the βME treatment. This was confirmed by repeating a set of experiments using deuterated βME (d-βME), inset Fig. 2b, where the dashed line represents the profile for d-βME when the buffer is contrast matched to gold. This clearly demonstrates that deuterated material is present on the surface. We note that the calculated nSLD for a filled layer of d-βME is 4.8 x 10⁻⁶ Å⁻², very similar to that of gold (4.5 x 10⁻⁶ Å⁻²). This value is very different from that for a filled layer of hydrogenous βME (0.62 x 10⁻⁶ Å⁻²). Model calculations demonstrate that the disappearance of fine features observed in the h-βME NR datasets (details in supplementary material) is due to the replacement of hydrogenous material with deuterated material.

**OmpF-E183C on surface-passivated gold**

It is well-known that thiol-gold bonds are relatively weak. Even after the gold surface has been fully passivated, incubation with other thiolated species leads to spontaneous replacement of
However, the extent of this spontaneous exchange in the OmpF system was not known. There are therefore two approaches that can be taken to fitting the OmpF data. One may either include a βME layer or assume that the βME has been completely displaced by the OmpF. Far superior fitting results were always obtained when a layer corresponding to βME was included in the fit. Figure 3a shows the results of the Monte Carlo resampling method applied to a fit from four different samples where the OmpF thickness was constrained to fit to the same value for all datasets (fitted value 53.6 ± 2.5 Å). A recent high resolution AFM study\textsuperscript{17} determined the protein thickness as 55 ± 13 Å from measurements of individual trimers. The thickness of the βME layer was fixed at 8 Å with the nSLD of the βME and OmpF layers allowed to vary between samples allowing for variations in surface coverage. Figure 3b displays the resampling results for the protein nSLD for sample #5 (Table 1) with the data and fit shown in Fig. 4a (filled squares) and 4b(solid line) respectively.

The βME, OmpF and D\textsubscript{2}O \(V_f\) in the mixed layer near the gold surface can also be calculated (Table 1). Firstly the OmpF proportion is fixed to that determined in the upper OmpF/D\textsubscript{2}O layer where there are only two components and the proportion of βME and D\textsubscript{2}O required to produce the observed nSLD in the mixed layer is calculated. This approach may underestimate the amount of βME in this layer as the base of the protein is not completely flat; making it likely that there is some reduced OmpF density immediately adjacent to the surface.

**Thiolipid layer: Backfilling of the OmpF layer and termination with DMPC**

In a first test experiment, NR data revealed that the deposition of the thiolipid 1,2-Dipalmitoyl-sn-Glycero-3-Phosphothioethanol (DPPTE) was incomplete at room temperature, i.e.
below its phase transition temperature (data not shown), producing layers with high water content and lipid tails that were not fully extended. This was overcome by undertaking the DPPTE deposition at 50 °C with the solution in contact with the surface for two hours. Longer exposure times ran the risk of the DPPTE displacing OmpF trimers. Figure 4 displays exemplary data, i.e. the “up” polarisation reflectivity of a sequence of sample preparation states, all measured under D$_2$O-based buffer. The “down” polarisation state and samples in contact with various contrast buffers have also been measured and were included in the constrained fit shown. The sample preparation steps involved sequential adsorptions of OmpF, DPPTE, and finally DMPC. The large change in signal between the OmpF and DPPTE datasets in Fig. 4a is due to the displacement of D$_2$O (positive nSLD) from around the OmpF by hydrogenous thiolipid (negative nSLD). The extra fringes observed in the data are due to this hydrogenous layer. Subsequent addition of DMPC then produced a much smaller change, with ‘new’ fringes shifted to slightly lower Q values signifying a thickening of the hydrogenous lipid layer. While the adsorbed OmpF on the βME was fitted in a two-layer model and the final sample (OmpF/DPPTE/DMPC) was well described by a three-layer model, a four-layer model was required to satisfactorily fit the data obtained from the intermediate step.

The interpretation of the models (Tables 1 & 2) is straightforward. Upon chemisorption, the protein partially displaces βME, and as a result, the surface film consists of a layer in which βME fills area in between surface-ligated protein, while at distances greater than ~ 8 Å from the interface solvent fills the corresponding volume, which results in a change in (average) nSLD. In the completed sample, the three subsequent layers contain OmpF and DPPTE headgroup (innermost layer), OmpF and DPPTE or DMPC acyl chains (central layer) and OmpF and DMPC
headgroups (outermost layer). Interpretation of the interface film structure at the intermediate stage of the preparation (OmpF and DPPTE chemisorption) is more complex. We have not been able to realistically model the data with less than 4 layers. These layers are interpreted as the DPPTE headgroups (plus OmpF), the aliphatic DPPTE chains (plus OmpF), a diffuse hydrocarbon layer (plus OmpF), and finally OmpF with some associated lipid or detergent protruding from these hydrocarbon layers (plus buffer). The inclusion of an intermediate layer, conceivably comprised of detergent adsorbed from the washing solution to the hydrophobic DPPTE chains, resulted in sensible layer thicknesses and nSLDs. With such a layer included in the model, the overall OmpF thickness, $57.5 \pm 0.9 \text{ Å}$, is in excellent agreement with AFM results. This protein layer thickness is in fact slightly larger than the thickness determined prior to DPPTE adsorption. Again, this is in excellent quantitative agreement with previous AFM data that suggested that surface-immobilised OmpF is stabilised by DPPTE backfilling which leads to a slightly longer projection of the protein’s crystallographic axis on the surface normal, resulting in a slightly larger layer thickness after backfilling. The result for the intermediate sample preparation suggests that detergent remains so tightly bound to the hydrophobically terminated DPPTE chains that it is resistant to copious rinsing with buffer.

The final self-assembly step adds a single layer of DMPC to the substrate to complete the bilayer structure, resulting in a hydrophilic outer surface. The overall thickness of the acyl chain layer ($31.6 \pm 0.8 \text{ Å}$) is as reasonably expected from the apposition of a palmitoyl and a myristoyl layer (combined length in the all-trans state: $\sim 35 \text{ Å}$). Whether any detergent is retained in the final preparation and if so, how much, cannot be determined, without performing dedicated experiments using deuterated detergent or DMPC. Splitting the lipid chains region into DPPTE and DMPC tails
did not improve the fit. The quantitative interpretation of the surface structure at different solvent contrasts shows that the addition of DMPC displaces nearly all of the buffer that was resident in the DPPTE layer during the intermediate sample preparation stage \( (V_f = 0.06) \). In the final sample, the buffer \( V_f \sim 0.01 \).

**Discussion**

The application of magnetic contrast neutron reflection (MCNR) has enabled us to define the model membrane system with significantly improved accuracy compared to earlier work and the Monte Carlo parameter resampling enables a rigorous determination of the experimental resolution. The improved resolution reveals that the OmpF protein only partially displaces the βME passivation layer. In distinction, the thiolipid back fill displaces βME quantitatively and leads to the formation of a dense layer with little water penetration. The assembly of the fluid DMPC layer around the OmpF is shown to be complete. The OmpF height which was difficult to measure by AFM in the absence of a rigid thiolipid layer\(^{17}\) is shown to be remarkably constant in the different stages of membrane formation. The improvement of resolution that underlies this interpretation of the surface structure has been achieved without lipid or protein deuteration, the use of which will further extend the complexity of structures that can be determined.

**Resolving the β-mercaptoethanol layer**

MCNR enables the structural characterisation of the coherent layer formed by βME. Adsorption of d-βME leads to a slight increase in nSLD near the gold interface (inset in Fig. 2b). Contamination by adsorbed organic species would have produced a large drop in nSLD near the gold surface similar to that seen for h-βME (Fig. 2b). That such a drop is not observed after d-βME
is adsorbed indicates that the adsorbate layer is βME. Moreover, even after this passivation layer is penetrated and partially replaced by OmpF, the remainder of this thin βME film can be structurally quantified by neutrons, as any model without a βME layer leads to a significant reduction in the fit quality (for example, an increase in $\chi^2$ from 1.67 to 2.56 for the data shown in Fig. 2). The data further indicates that the βME layer is finally replaced during DPPTE deposition.

**OmpF-E183C on surface-passivated gold**

The production and analysis of OmpF layers is highly dependent upon the use of high quality substrates. Compared with our previous low resolution NR study\(^3^7\) where about 10% of the surface was covered with protein we are consistently achieving OmpF surface coverages of 30% or greater. To put this into context, the volume fraction of protein in a 2D OmpF crystal is about 0.50 – so we are achieving 60% of this maximum packing. The precision obtained for the protein thickness and density is far in excess of that previously obtained. In view of recent data on monomeric OmpG and OmpA porins in detergent micelles, the height of the protein array is an especially interesting parameter. As reported by NMR, the non-crystalline porins in micelles display shorter β strands and a lower overall height compared to their respective x-ray crystal structures\(^4^3, 4^4\). No NMR data exist yet for the larger trimeric OmpF but AFM data\(^1^7\) on gold immobilised OmpF showed that the proteins were too flexible to observe clearly before thiolipid addition. However, after backfilling the protein array with thiolipid, the trimers were clearly imaged by AFM but the average height was the same both cases. The result from NR confirms the unchanging dimensions of surface-bound OmpF, suggesting that the height of the trimeric OmpF may be less sensitive to the change from detergent stabilised to tight packing than monomeric porins. The AFM work could not measure the absolute OmpF height in thiolipid and assumed a
thiolipid thickness through which the OmpF penetrates. Neutrons on the other hand penetrate the whole layer, are non-destructive, and provide an average over the whole surface. Therefore, agreement between these very different but complementary techniques is strong evidence for the accuracy of the models.

**Thiolipid Addition**

Deposition of the DPPTE below its phase transition was unsuccessful with poor surface packing and very little extension of the lipid tails. In AFM studies\(^\text{17}\) of this system the thiolipid was deposited at 45 °C, above the phase transition temperature of DPPC. A similar protocol (at 50 °C) led to a dense, dry (6 % water content) layer corresponding to the expected thickness of DPPTE (Fig. 4), confirming that assembly of a dense ordered DPPTE monolayer only occurs above the lipid phase transition temperature. To avoid this temperature requirement in future work with possibly less robust proteins, thiolipids using diphytanoyl chains with lower phase transition temperatures may be used.

The expected result at this stage of the assembly is a half bilayer facing the water, through which the proteins protrude. Due to the thinner cross section of the *E. coli* outer membrane, compared to more usual fluid lipid bilayers, the outer membrane protein’s hydrophobic surface is largely covered by the single thio-DPPTE layer.\(^\text{17}\) This leaves the thiolipid’s hydrophobic surface exposed to the aqueous phase. Such surfaces have a high free energy but, due to the covalent attachment of the thiolipid to the gold, are stable.\(^{45, 46}\) It has been demonstrated that adsorbed detergents will dissociate from hydrophobic SAMS when adsorbed from solution concentrations up to about 8 times greater than the CMC.\(^\text{47}\) In this work the SDS washing was undertaken at about concentrations about seventy times greater than the CMC with the rinsing undertaken on an open
surface with very different shear forces than may be experienced in a confined SPR cell. It also needs to be kept in mind that CMC is usually determined in pure buffer (i.e. in the absence of lipid aggregates) and that the effective CMC of a particular detergent may be affected by the presence of such aggregates. In the AFM study extensive washing was used and no evidence for an additional surface layer was obtained. MCNR reveals that there is a highly disordered hydrocarbon layer adsorbed onto the hydrophobic outer DPPTE surface. This layer is about 5 Å thick with a $V_f$ of circa 0.6, most likely consisting of detergent molecules lying flat on the surface. As with the AFM study, the samples were prepared ex situ which allowed copious washing of the surfaces so if the same layer was present in the AFM study it was either removed by the AFM tip or was incorporated into the measurement baseline. This residual layer corresponds to a coverage of ca 100 pmol/cm$^2$, compared to the total surface coverage of 280 pmol/cm$^2$ determined by Sigal et al [REF 45] prior to washing.

**Membrane Completion with DMPC**

The final step of the process used DMPC to complete the bilayer surrounding the OmpF. The fits to the experimental data (Fig. 4) indicate that this was successfully achieved. The OmpF thicknesses in the samples comprised of (thiolipid + OmpF) and (thiolipid + DMPC + OmpF) were within 1.5 Å of each other (Table 2). The DMPC displaced both the disordered hydrocarbon and some of the water from the DPPTE during deposition. Both effects may have been aided by the fact that the rapid solvent exchange method is undertaken with the lipid dissolved in ethanol. The final bilayer with OmpF incorporated ($V_f = 0.26$) can obtain a maximum $V_f$ of 0.74 for the phospholipid layer. Therefore the calculated $V_f = 0.73$ indicates the DMPC coverage of the available surface exceeds 98%, consistent with results with DMPC (and other lipids) on tethered bilayer lipid membranes.
The versatility of the rapid solvent exchange method is demonstrated by the finding that the surface coverage in a layer formed around an existing protein protruding from a lipid monolayer was equal to or exceeded that obtained for lipids deposited onto a stable well formed self assembled lipid monolayers.

**Magnetic Contrast**

To quantify the benefit of MCNR one can fit the data in an unconstrained manner thereby examining each spin state in isolation. In this procedure, the fit often converges to a lower $\chi^2$ value as there are fewer boundary conditions to be satisfied. To demonstrate the impact of the MCNR we have considered the case of βME adsorption where there is data available with varying βME and subphase contrasts. The βME layer thickness is near the resolution limits of the technique but the range of contrasts available mean that the test is carried out under conditions most favourable to the ‘standard’ NR approach. Figure 5 displays the MC resampling analysis for the βME layer from fits to datasets relying on magnetic or subphase contrast fitting. The total thickness obtained is very similar 220.6 versus 221.2 for the magnetic and subphase contrasts respectively. There is a variation in both the βME thickness and nSLD values obtained. For the magnetic contrast datasets the βME layer is thicker (7.6 ± 1.3 versus 3.7 ± 0.6 Å) with a lower $V_r$ 0.66 ± 0.07 versus 0.80 ± 0.14. The changes in βME parameters are compensated for by the gold layer being thicker for the subphase contrast fits. The βME is poorly determined in the subphase contrast fits, with a narrow distribution centred very close to the lower limit allowed in the fit, furthermore when this thickness is convoluted with the interfacial roughness required in for a realistic model the layer becomes less than the combined width of the adjacent interfaces. This is not the case for the magnetic contrast fit
where the layer is clearly visible when adjacent interfacial roughnesses are taken into account. Relying on subphase contrast alone it was not possible to clearly distinguish the \( \beta \text{ME} \) layer.

Time did not permit data collection on all samples with both \( \text{D}_2\text{O} \) and \( \text{H}_2\text{O} \) buffers (or combinations thereof) so it is not possible to similarly demonstrate the impact of magnetic contrast for each assembly step. Nevertheless, when one considers the nSLD profiles for samples to which the thiolipid had been added (Fig. 4) it is clear that \( \text{H}_2\text{O} \) would have provided very poor contrast especially since the thiolipid is only available in hydrogenous form. In fact the calculated nSLD in \( \text{H}_2\text{O} \) is almost indistinguishable before and after thiolipid adsorption (Fig. 6). Therefore magnetic contrast combined with a standard \( \text{D}_2\text{O} \) buffer is the best approach available to readily access contrast variation in this system. Furthermore the approach does not complicate the sample preparation as a metallic binder layer, such as titanium or chrome, is always required between the gold and silicon layers.

**Conclusions**

Our previous study of this system by neutron reflection, while producing useful data about the buried proteins layers *in situ* and confirming OmpF orientation,\(^{37}\) did not demonstrate sufficient resolution to disentangle the scattering from all components. Structural and compositional resolution of this system has only been achieved through the production of high quality samples and the application of magnetic contrast neutron reflectometry. The presence of the \( \beta \text{ME} \) layer has been observed with the expected structural parameters. Such layers are important in the formation of tethered lipid bilayers \(^1,\ 35,\ 40,\ 41\) and this is the first report which reveals the layer *in situ*. Furthermore it indicates the layer thickness resolution available from this method and it is clear that after OmpF adsorption the \( \beta \text{ME} \) is still present on the surface between the OmpF trimers. It is
not until the thiolid is added to the surface that the \( \beta \)ME is displaced. Through the use of magnetic contrast and some minimal buffer contrast variation we have been able to accurately determine the relative proportions of water, lipid and protein in the biomolecular layer. The height of the OmpF has been measurable throughout with a small initial extension (3.9 Å) upon the assembly of the TL. The final assessment of the structure indicates that the original aims of producing a stable robust protein scaffold, with solution accessible surface, embedded within a model membrane lipid bilayer have been achieved. The method outlined is especially suited to the study of biotechnical devices that employ gold substrates for charge carrying purposes. More broadly it would be applicable to any layered soft matter system that incorporates a magnetisable layer or substrate. This application is especially relevant when the selective deuteration of components is problematic.

**Materials and Methods**

**Magnetic scattering of polarised neutrons**

The neutron is scattered from the atom’s nucleus and the application of neutron scattering techniques allows the spatial distribution of nuclear species to be determined. Biological systems are hydrogen rich, and the two stable isotopes \( ^1\text{H} \) (protium) and \( ^2\text{H} \) (deuterium), although chemically very similar, interact very differently with thermal neutrons, as defined by their nuclear scattering lengths \((b(\text{H}) = -3.74 \times 10^{-15} \text{ m}, b(\text{D}) = +6.67 \times 10^{-15} \text{ m})\) related to a characteristic refractive index, analogous to that of a photon's. The combination of \( b \) with the physico-chemical composition of a material (isotope density) defines its characteristic nSLD (equation 2) and regions of different nSLD provide the contrast variation needed to solve structures by neutron methods. Hydrogen contrast variation therefore provides a range of possibilities within a multi component
sample and has been extensively applied to studies of small molecules such as surfactants at interfaces, for a review see Lu et al.\textsuperscript{48} For example the CH\textsubscript{2} groups of cetyl trimethylammonium bromide have been selectively deuterated as a function of position along the surfactant tail enabling high spatial resolution information on the self assembled layer to be attained.\textsuperscript{49-51} As the molecules involved increase in size, selective deuteration becomes more labour intensive and ultimately chemically intractable. Deuteration of proteins has been achieved in a limited number of systems\textsuperscript{52-54} and the use of aqueous phase deuteration is also somewhat limited. There is, therefore, a need to find alternative means of achieving increased contrast variation in a manner that neither complicates sample preparation nor alters the biological sample.

As in X-ray or neutron diffraction measurements, only the intensity of the scattered radiation is measured in a NR experiment. Without the corresponding phase information a unique solution to the density profile that produced the scattering cannot be guaranteed and ambiguities in the compositional depth profiles arise. This is especially the case when a single specular reflection dataset is considered since there are usually multiple mathematically acceptable solutions. In certain cases, independent knowledge of parts of the nSLD profile can suffice to identify which of a number of model profiles, each an equally good mathematical solution to the reflectivity, corresponds to physical reality. Reference layers, incorporated into the sample, may be used to retrieve the missing phase information which is the cause of the inherent ambiguity in nSLD.\textsuperscript{55} X-ray protein crystallography employs an analogous approach where a heavy metal ion is incorporated into the protein crystal providing a reference point that enables phase information to be retrieved from the data. Similarly reflectivity curves from a sufficient number of samples, each
consisting of an invariant unknown region plus a varying but known reference part, would enable
the determination of the complex reflection amplitude of the unknown.

One possible reference is a buried, saturated ferromagnetic layer that results in two
different nSLD profiles, depending on the spin eigenstate of the neutrons in a polarised beam. By
measuring two reflectivity datasets from a composite system, consisting of a ferromagnetic
reference layer plus unknown, one with a beam of neutrons in the "+" spin state and the other with
spin "-" neutrons, the complex reflection amplitude for the unknown segment alone can be exactly
obtained and directly inverted to provide its corresponding scattering length density depth profile.\(^{55}\)

In practice, however, statistical uncertainty and limited range of scattering vector can make an
essential part of this calculation process — the selection of the physical root of a quadratic
equation — problematic.\(^{56}\) A minimum of two magnetic contrast datasets were simultaneously
refined with appropriate constraints to retrieve the unknown nSLD of the region of interest; an
approach that implicitly takes into account the phase information distributed over the multiple
datasets. Without magnetic contrast the external reference approach to phase retrieval would
require the impossible assembly of two identical biological samples (with identical thickness,
surface coverage) upon different reference layers with identical surface roughness and if present
oxide layer thickness. Future developments in data collection strategies may enable directly
inversion of the MCNR data to obtain the correct model independent nSLD profile.\(^{57}\)

If possible selective deuteration of the synthetic and protein components would enable
different contrasts for the alkane and protein components of the model membrane in different
samples with assembly on a range of different solid substrates. This would provide independent
sets of data with different contrasts but with inherent variability between each. Not only is it far
easier to apply the magnetic contrast approach to a single sample but also both contrasts can be achieved at each self-assembly step on a single sample. Furthermore, this approach is entirely complementary to deuteration and has therefore the potential to greatly enhance the information content of data obtained from deuterated samples. Data collection on the NG1 instrument involved repeated scans over the entire scattering vector (Q) range. The equality of subsequent scans then demonstrates that samples are unchanged throughout data acquisition (12 – 24 hours). Our experimental programme has also enabled us to develop an assembly protocol that produces highly reproducible samples and it may therefore be profitable to pursue the synthetic route for deuterated DPPTE to achieve even higher resolution compositional data.

The metallic layers required a thickness homogeneity and surface roughness of better than 10 Å over the entire surface (area > 4.5 cm²). To ensure appropriate smoothness, flatness and stability a metallic binder layer was first deposited on the silicon, followed by the gold layer. Correct selection of the binder enabled a magnetic reference layer to be incorporated into the standard substrate. Application of an external magnetic field controlled the direction of the magnetisation in this binder layer. The incoming neutron beam was polarised so that the neutron spin was either parallel or anti-parallel to the direction of magnetisation in the layer, resulting in two different datasets with different reflectivity signals. During data fitting all model parameters were kept exactly the same for the two datasets except for the nSLD values of the magnetic layer.

**Materials**

Unless otherwise stated all materials were obtained from Sigma and used without further purification. Thiolipid (1,2-dipalmitoyl-sn-glycero-3-phosphothioethanol, DPPTE) and DMPC (1,2-dimyristoyl-sn-glycero-3-phosphocholine) were purchased from Avanti Polar Lipids. Sample
preparation entailed two distinct procedures; firstly there was the deposition of the gold and magnetic layers in advance of the NR experiments. The second stage (Fig. 1) was solution based with the protein and lipid multilayer self-assembled onto the gold surface. NR experiments were undertaken at National Institute of Standards and Technology (NIST), Gaithersburg MD, USA and at ISIS, Rutherford Appleton Laboratories, Oxfordshire, UK. The magnetic (40 – 80 Å) and gold (150 – 250 Å) layers were deposited sequentially in the same chamber. Three different magnetic layers were employed during the experiments, pure iron, permalloy (80 % nickel, 20 % iron) and mu metal (75 % nickel, 15 % iron, trace copper and molybdenum). Experiments at NIST were undertaken on silicon disks 75 mm diameter by 5 mm thick which were coated in a DC Magnetron Sputtering chamber (Auto A306; BOC Edwards, UK). Experiments at ISIS were carried out on silicon discs 100 mm diameter by 10 mm thick with the metal layers produced by INESC Microsystems & Nanotechnologies, Lisbon Portugal, in a Nordiko 3000 Ion Beam system.

The self-assembly from solution and washing steps were carried out in buffer A (1% OG, 20 mM Tris-HCl pH 7.5, 1 mM TCEP) and B (5 mM Na$_2$HPO$_4$, 100 mM NaCl, pH 7.4). The OmpF was prepared and purified as has been previously described$^{17, 32}$. The assembly of all components except for the DMPC were carried out \textit{ex situ}.

**Neutron Reflection**

The NIST Center for Neutron Research NG1 instrument is located on a reactor source and operates with a neutron beam of fixed wavelength incident on the sample whereas CRISP at ISIS is a time of flight reflectometer on a spallation source supplying a multiwavelength neutron beam. The different instrument configurations and neutron beam characteristics resulted in very different data
collection approaches for the two instruments. The reflectivity (ratio of the reflected to incident beam intensity) was measured as a function of $Q$, the scattering vector. $Q$ is defined as follows;

$$Q = \frac{4\pi \sin\theta}{\lambda}$$  \[1\]

where $\theta$ is the angle the incident beam makes with the interface and $\lambda$ is the wavelength of the neutrons. It is therefore possible to vary $Q$ by altering either the wavelength (CRISP) or the angle (NG1) during a measurement. Data were collected over $Q$ values from 0.011 to 0.3 Å$^{-1}$. Specular reflection occurs where the angle the incoming beam makes with the surface is equal to that of the reflected beam.

On NG1 data is collected with the neutron wavelength fixed at 4.75 Å and the incident beam angle scanned from 0.23 to 7.2 $^\circ$ with increasing slit openings for each data point such that a fixed sample area is illuminated. For CRISP a neutron beam, wavelength range 1.2 – 6.5 Å, is incident on the sample at three different incident angles also with a fixed illumination area producing an overall similar $Q$ range on both instruments. The reflectometers were operated in polarised beam mode, achieved by the same method on both devices. The incoming neutron beam is first incident on a polarising supermirror, guide fields are then used to maintain the neutron polarisation and finally the beam passes through a spin flipper which sets the direction of the neutron beam polarisation onto the sample. The sample geometry is such that the polarisation is parallel to the silicon surface and the magnetisation of the buried layer is induced by the application of an external magnetic field. The spin flipper flips the direction of the neutron polarisation so that it is either parallel (‘up’ spin) or anti-parallel (‘down’ spin) to the direction of the reference layer magnetisation. It is possible to analyse the polarisation of the neutron beam.
after the sample but this was not undertaken as the biological layer used did not contain any magnetic structure. Tests were performed on NG1 confirming the complete absence of spin-flip scattering (data not shown) which would have indicated non-uniform magnetic structure in the buried layer. As indicated in Fig. 1 the neutron beam is transmitted through the silicon substrate prior to reflecting from the interface.

**Data Fitting**

The processing of the raw data to produce absolute reflectivity was different for each instrument. On NG1 the main correction to be applied was that related to the changing slit settings across the angular range. A slit scan with the beam transmitted through silicon was performed at the beginning of each experimental campaign to correct for this. On CRISP, corrections for silicon transmission and detector efficiency, which varies as a function of wavelength, were required. Nevertheless once the corrections were applied and the data correctly scaled there was no difference in the data fitting programmes or routines used to treat the data. Fitting of the data was undertaken using the GA_refl programme [P.A. Kienzle, M. Doucet, D.J. McGillivray, K.V. O'Donovan, N.F. Berk, C.F. Majkrzak; http://www.ncnr.nist.gov/reflpak. 2000-2006]. This approach is based upon the optical transfer method which assumes that the interface can be described as a series of slabs where there is a change in refractive index for neutrons at the interface of each slab. If protiated and deuterated material are mixed correctly it is possible to produce an interface which is contrast matched and becomes invisible to the neutron beam. There are three parameters that describe each slab or layer, the thickness (d, Å), a Gaussian interfacial roughness (σ, Å) and the nSLD (Å⁻²) of the layer. It was possible to simultaneously co-refine multiple datasets (from both NG1 and CRISP) with a range of constraints applied to the fit. For the
simplest case, where two magnetic contrast datasets were co-refined, all parameters were set to refine to the same value except for the nSLD of the magnetic layer. If magnetic contrast plus a variation in buffer contrast datasets were refined together then all thickness and roughness values were equivalent between datasets with only the appropriate nSLDs (reference, buffer) allowed to vary.

The nSLD of a material is a function of the chemical composition (atomic number and density) and the nuclear scattering length as shown in equation 2.

\[ nSLD = N_A \sum_i \frac{p_i b_i}{A_i} \]  \hspace{1cm} \text{[2]}

Where \( N_A \) is Avagadro’s number, \( p_i \) the physical density, \( A_i \) the atomic number and \( b_i \) the nuclear scattering length of component \( i \). From comparisons of the nSLD from a fitted layer with the theoretical value for a complete layer it is quite straightforward to calculate the volume fraction or, if we assume that within the resolution of the experiment the nSLD profile of a component is homogenous in all directions, we can calculate the surface coverage via projection of the volume fraction onto a planar surface.

The data fits were evaluated by the application of a Monte Carlo resampling procedure\textsuperscript{59} using the best fit to the dataset as a starting point. At least \( N=1000 \) synthetic datasets were produced by applying random Gaussian weighted deviations from the data based upon the counting statistics of each data point. These synthetic datasets were analysed in the same manner as ‘real data’, outputting \( N \) variations of each parameter. The fits to the synthetic data were analysed producing a frequency plot of fitted values. These parameter distributions were statistically
analysed with the parameter value reported as the midpoint of the 95 % confidence interval. The standard deviation of the distribution is reported as the error. The frequency distribution obtained from this procedure is histogrammed (bin width = 3.49*standard deviation*N-1/3) before presentation.

Acknowledgements.

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References


Tables
Table 1. Layer thicknesses and Volume Fraction of components from constrained fits to multiple datasets, numbers in brackets are the errors assessed via the Monte Carlo resampling method. Where data was collected for βME adsorption and then OmpF adsorption volume fractions are reported for both steps. Where only data on samples with βME or OmpF adsorption were collected volume fractions are only reported for one adsorption. For example sample 1 had a βME volume fraction of 0.63, after OmpF adsorption the sample surface showed βME, OmpF and Buffer volume fractions of 0.59, 0.27 and 0.14.

<table>
<thead>
<tr>
<th>Sample Number</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>βME Thickness$^1$ (Å)</td>
<td>8.2 (0.6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>βME V$_f$</td>
<td>0.63</td>
<td>-</td>
<td>0.83</td>
<td>0.54</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>(0.05)</td>
<td></td>
<td>(0.17)</td>
<td>(0.03)</td>
<td></td>
</tr>
<tr>
<td>OmpF Thickness$^2$ (Å)</td>
<td>53.6 (2.5)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>βME V$_f$</td>
<td>0.59</td>
<td>0.35</td>
<td>0.31</td>
<td>-</td>
<td>0.43</td>
</tr>
<tr>
<td></td>
<td>(0.02)</td>
<td>(0.08)</td>
<td>(0.01)</td>
<td></td>
<td>(0.01)</td>
</tr>
<tr>
<td>OmpF V$_f$</td>
<td>0.27</td>
<td>0.39</td>
<td>0.34</td>
<td>-</td>
<td>0.26</td>
</tr>
<tr>
<td></td>
<td>(0.01)</td>
<td>(0.02)</td>
<td>(0.01)</td>
<td></td>
<td>(0.01)</td>
</tr>
<tr>
<td>Buffer V$_f$</td>
<td>0.14</td>
<td>0.26</td>
<td>0.35</td>
<td>-</td>
<td>0.31</td>
</tr>
<tr>
<td></td>
<td>(0.02)</td>
<td>(0.08)</td>
<td>(0.02)</td>
<td></td>
<td>(0.02)</td>
</tr>
</tbody>
</table>

1. βME thickness from fits to βME adsorption step only.

2. Fitting from samples with βME + OmpF adsorbed where the βME thickness was constrained to that determined above.
Table 2. Fitted thickness and layer compositions after thiolipid and then DMPC addition to sample 5 (Table 1).

<table>
<thead>
<tr>
<th></th>
<th>DPPETE Addition</th>
<th>DMPC Addition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner Headgroup Thickness (Å)</td>
<td>13.4 (0.1)</td>
<td>14.4 (0.4)</td>
</tr>
<tr>
<td>Lipid Tails Thickness (Å)</td>
<td>21.9 (0.5)</td>
<td>31.6 (0.8)</td>
</tr>
<tr>
<td>Hydrocarbon Thickness (Å)</td>
<td>4.6 (0.7)</td>
<td>-</td>
</tr>
<tr>
<td>Protruding OmpF Thickness (Å)</td>
<td>17.6 (0.4)</td>
<td>-</td>
</tr>
<tr>
<td>Outer Headgroup Thickness (Å)</td>
<td>-</td>
<td>12.7 (0.3)</td>
</tr>
<tr>
<td>Layer total (Å)</td>
<td>57.5 (0.9)</td>
<td>58.7 (0.9)</td>
</tr>
<tr>
<td>OmpF $V_f$</td>
<td>0.26</td>
<td>0.26</td>
</tr>
<tr>
<td>Lipid Tails $V_f$</td>
<td>0.68</td>
<td>0.73</td>
</tr>
<tr>
<td>Buffer $V_f$</td>
<td>0.06</td>
<td>0.01</td>
</tr>
</tbody>
</table>

Figure Legends.

Figure 1. Schematic of the sample configuration for the neutron reflection studies. The substrate was exposed to a 1 % (v:v) βME in ethanol solution. OmpF-E183C (300 µg/ml in buffer A) was incubated on the gold surface for at least 3 hours at room temperature. After incubation, the surface was washed, with a 2 % (w:v) SDS solution and then with distilled water, to remove any non-specifically bound protein. Thiolipid DPPETE (1,2-dipalmitoyl-Sn-glyero-3-phosphothioethanol), (1.0 mg/ml in buffer A), was then deposited to infill around the OmpF trimers (right hand side of figure). Finally DMPC (10 mg/ml in ethanol) was added to the assembled surface (left hand side of figure) and incubated for 5 minutes. The DMPC solution was removed by washing the cell quickly with 50 ml of buffer B. The figure shows an OmpF trimer (PDB 2OMPF) attached to the gold via cysteine residues surrounded by DPPETE (pdb file 870160.mol from Avanti Polar Lipids]. DMPC molecules used to represent the upper layer are
taken from a simulated DMPC bilayer structure ([http://moose.bio.ucalgary.ca/files/dmpc_npat.pdb](http://moose.bio.ucalgary.ca/files/dmpc_npat.pdb)]. The cysteine residues are yellow and space filled and the belt of tyrosine residues (wire frame, red) delineates the membrane region. The incident neutron beam (red arrow labelled \(k_i\)) was directed to reflect (\(k_r\)) from behind the membrane layer. The blue arrow labelled \(Q\) shows the scattering vector.

Figure 2. a) Reflectivity data (symbols) and fit (line) after \(\beta\)ME adsorption onto the surface, \(D_2O\) buffer. Below the critical edge (CE) the neutrons are totally externally reflected and the reflectivity is defined as unity. \(Q\) is the scattering vector defined in equation 1. The data have not been offset, the separation results from the different contrast of the magnetic layer for spin up and spin down neutrons. b) Real space nSLD profile corresponding to the fit shown in a). The zero point has been set at the interface between the gold and the \(\beta\)ME. The silicon substrate is on the left and \(D_2O\) buffer on the right. The 2 different nSLD values for the magnetic layer are clearly seen. The hydrogenous \(\beta\)ME layer is clearly evident between the gold and the buffer. The inset shows the \(\beta\)ME region of the sample expanded with the dashed line the fit obtained with d-\(\beta\)ME next to a gold matched water buffer. NOTE: Data presented in Figs 2 & 4 are all from successive depositions on the same substrate.

Figure 3. Results from the Monte Carlo resampling of magnetic contrast data for self-assembled OmpF layers where the line is a simple Gaussian fit intended to provide a guide to the eye. The
frequency axis represents the number of times a particular result was obtained in the 1000 trial fits. 

A) OmpF layer thickness, and b) nSLD of the OmpF layer (see Figure 4).

Figure 4. a) Data, ‘up polarisation’ state (symbols) and fit (lines) after OmpF adsorption (x), DPPTE adsorption (+) and DMPC adsorption (o). The OmpF and DMPC datasets have been offset for clarity. B) the real space nSLD profiles corresponding to the fits of all polarisation data, solid line - OmpF adsorption, dots – DPPTE adsorption and dashes – DMPC adsorption. The figure has been labelled to illustrate the main constituent of each region, HGs indicates regions of lipid headgroup.

Figure 5. Monte Carlo resampling analysis of the use of magnetic contrast upon the parameters for the βME layer. a) layer thickness and b) nSLD values where circles are from magnetic contrast fits with squares corresponding to fits relying solely on buffer contrast. The lines are simple Gaussian fits to guide the eye.

Figure 6. nSLD variation with OmpF surface coverage for D$_2$O and H$_2$O buffers before and after thiolipid adsorption. Pure H$_2$O and D$_2$O nSLD are shown as solid horizontal lines (vertical dashed line represents maximal OmpF surface coverage from this work). The available contrast against the aqueous buffer is the difference from the contrast of interest, at the appropriate surface coverage, to either the D$_2$O or H$_2$O lines.
Figure 1.

Figure 2
Figure 3.
Figure 4.
Figure 5.
Figure 6.