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Structural Studies of Aliphatic Substituted Phthalocyanine-Lipid Multilayers

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A Langmuir–Blodgett film of aliphatic substituted phthalocyanines on a C18 silane supporting layer coupled onto a silicon substrate has been investigated using neutron reflectometry. This multilayer structure is seen as a possible candidate for phthalocyanine–lipid biosensor devices. The results show the suitability of the C18 ligands as an anchoring layer for the phthalocyanines. The scattering length density profiles demonstrate the effectiveness of a lipid monolayer in partitioning the composition of phthalocyanine layers from that of the bulk liquid. The effectiveness of this barrier is a critical factor in the efficiency of such devices.

Introduction

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Since the discovery of phthalocyanines (Greek word, rock oil 17 blue) in 1907¹ and their subsequent in-depth structural and 18 chemical study a number of years later,² these compounds have 19 shown potential in countless applications. Phthalocyanines are 20 thermally and chemically very stable. They also display intense 21 color and possess catalytic, electrical, as well as photophysical 22 properties.³ Phthalocyanine derivatives can form Langmuir 23 24 monolayers at the air-water interface. Their deposition on a 25 solid support using the Langmuir-Blodgett (LB) technique on hydrophobic surfaces has shown that the phthalocyanine ring will 26 exhibit an edge-on orientation.⁴ This is thought to be due to the 27 amphiphilic character of these compounds. 28

The design of integrated optical sensors using aliphatic sub-29 stituted phthalocyanines for continuous monitoring and spot 30 checks on water quality has been reported.⁵ These novel and 31 highly sensitive, integrated optical biosensors are based on detec-32 tion of change in the refractive index of a sensing layer saturated 33 with nitrous molecules encapsulated between a biomimic mem-34 brane (phospholipid) and a solid support. The change in refrac-35 tive index is measured using a stacked planar waveguide in a dual-36 polarization interferometer arrangement in these devices. The 37 38 diffusion of entrapped nitrous molecules through the biomimic encapsulation will change as a function of damage caused by 39 40 either bioagents (bacteria, viruses) or chemical agents (heavy metal contamination, pesticides) and is detected by change in the 41 refractive index of the sensing layer. 42

The focus of our project was to study the structure and hence
 optimization of the design of a disposable version of these sensors,
 removing the need for optical measurement and the reliance on

the waveguide as a means of detection. This design will involve 46 replacing the optical waveguide part of the sensor by a multilayer 47 of an aliphatic substituted phthalocyanine compound, which can 48 incorporate a color pigment, sensitive to nitrous oxide, which is in 49 turn deposited on a hydrophobically modified silicon substrate 50 using the Langmuir-Blodgett technique. A schematic diagram of 51 such a device is shown in Figure 1. The first step in the 52 F1 commercialization of these devices involves a better understand-53 ing of the structure of the interfaces, that is, layer uniformity, the 54 composition, and the effectiveness of a lipid layer as a membrane 55 separating the sensing region from that of the bulk water. We 56 report here a structural study of Langmuir-Blodgett deposited 57 aliphatic (C10) substituted phthalocyanines (Figure 2) and lipid 58 F2 multilayer structures on a hydrophobic solid support using 59 neutron reflectometry technique. We seek to determine the water 60 content of the phthalocyanine multilayers, as this plays an impor-61 tant role in the long term stability of these devices, and also to 62 investigate the interface between the aliphatic (substituted phtha-63 locyanine molecules) and the mimic layer (lipid monolayer). The 64 results will provide a better understanding of the interfaces and will 65 ultimately enable engineering of more robust and reproducible 66 commercial biosensors. 67

Experimental Section

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Materials. *d*-1,2-Dipalmitoyl-*sn*-glycero-3-phosphocholine 69 (DPPC) and functionalized poly(ethylene glycol) (PEG) lipids 70 DSPE-PEG(2000) were purchased from Avanti (Avanti, Polar 71 Lipids, Inc.). The synthesis of the aliphatic substituted phthalo-72 cyanine is reported elsewhere.⁶ The D₂O was obtained from 73 Fluorochem (>99 at. D%), and ultrapure H₂O was produced 74 using an Elgastat water purification unit. The silicon blocks 75 (Crystran Ltd., Poole, U.K.), used as the substrate, were circular 76 in section with a diameter of 100 mm and a thickness of 10 mm and 77 were polished on one (111) face to ca. 6 A. The silicon surface was 78 initially cleaned using the RCA method.⁷ The substrates were then 79

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Figure 1. Schematic diagram of the phthalocyanine sensor device.



Figure 2. Chemical structure of the phthalocyanine ligand. The six R groups are $C_{10}H_{21}$.

chemically dried prior to treatment with a silane coupling agent
 octadecyltrichlorosilane (ACROS, 95%) using established
 methods.⁸ This has been shown to provide a chemically bound
 hydrophobic layer on the silica surface.⁹ In our previous work, we
 have characterized this layer by measuring the reflectivity from the
 Si-water interface using a series of contrasts.⁹

Neutron Reflectometery. Reflectivity measurements were 86 carried out using the reflectometer SURF9 at the ISIS Spallation 87 Neutron Source, Rutherford Appleton Laboratory, Didcot, U.K. 88 The neutron beam is polychromatic with wavelengths in the range 89 90 $0.53 < \lambda < 6.9$ Å. To obtain the widest amenable momentum 91 transfer, Q, range, reflectivity spectra were measured for a series of grazing incidence angles θ , where θ is the angle of incidence at 92 93 the Si-water interface and $Q = (4\pi \sin \theta)/\lambda$ is the momentum transfer at the interface. The nominal incidence angles used were 94 95 0.35, 0.8, and 1.8°. The actual angles of incidence were determined 96 by performing detector angle scans in reflection geometry once the 97 height alignment of the sample with respect to the neutron beam 98 has been optimized. The collimating slit settings were varied with 99 incidence angle in order to measure all reflectivities with a constant angular resolution ($\delta\theta/\theta = 3\%$). The sample was under-100 illuminated with an illuminated length of ~45 mm projected on 101 102 the sample.

The measured reflectivity profiles are normalized relative to the incidence beam monitor spectrum and corrected for detector efficiency as per standard reflectivity measurement on a time-offlight instrument,¹¹ and the data were subsequently corrected for the wavelength-dependent transmission through the silicon substrate. The data obtained were then overlapped in Q. 108

Neutron reflectivity is a technique sensitive to the average neutron refractive index, n, profile normal to an interface.⁸ The dispersive refractive index can be written as 111

$$n(\lambda) \approx 1 - \frac{\lambda^2}{2\pi} Nb + i \frac{\lambda}{4\pi} N\sigma$$

where λ is the neutron wavelength, $Nb = \sum_i N_i b_i$, and $N\sigma = \sum_i N_i \sigma_i$, 112 with N_i being the number density, b_i the coherent scattering 113 length, and σ_i the absorption and incoherent cross-section of 114 nucleus *i*. The multiple Nb is known as the scattering length 115 density of a medium with refractive index n. According to the 116 above equation, the large difference in the scattering lengths of ¹H 117 (b = -3.7406 fm) and ²H (b = 6.671 fm) can be exploited in 118 hydrogenous systems. Since Nb is linearly related to the volume 119 fraction composition $(Nb \approx \Sigma_i \phi_j \overline{Nb_i})$, where ϕ_i is the volume 120 fraction and Nb_i is the scattering length density of species j), a 121 layer model with discrete strata representing regions with different 122 chemical composition can be constructed and the reflectivity from 123 such a model can readily be calculated. Each layer, *i*, has a 124 thickness, d_i , refractive index, n_i , scattering length density, Nb_i , 125 and interfacial roughness, σ_i . The reflectivity can be calculated 126 using, among other methods, the standard optical-matrix method, 127 and the parameters of the proposed layer model can be optimized 128 using nonlinear least-squares fitting. 129

Two silicon blocks, S1 and S2, were used in these experiments. 130 The surfaces of both these substrates were first cleaned and 131 then rendered hydrophobic by means of silane coupling. A C-18 132 layer was deposited. A series of 2, 4, and 6 layers of substituted 133 phthalocyanine were subsequently deposited onto the C18 134 coated silicon substrate S1, using Langmuir-Blodgett dipping 135 technique.¹² The deposition was performed using a standard 136 Teflon Langmuir trough. The trough was filled with ultrapure 137 water (Milli-Q). A solution of phthalocyanine in chloroform was 138 prepared and then carefully spread at the air-water interface. An 139 ordered Langmuir film was then obtained by moving the poly-140 (tetrafluoroethylene) (PTFE) barriers of the trough until a surface 141 pressure of 25 mN m⁻¹ was reached. The surface tension was 142 monitored by the Wilhelmy plate method. A sample holder was 143 manufactured to dip the silicon block into the water, and while the 144 block was immersed, the surface pressure was maintained at 145 25 mN m^{-1} by computer controlled motion of the PTFE barriers. 146 The sample holder allowed dipping/removing of the block from 147 the aqueous phase at a constant speed of 5 mm/min. First, a bilayer 148 was deposited. The block with a bilayer was then characterized 149 using neutron reflecotmetery with D₂O forming the aqueous 150

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Detector

Figure 3. Neutron reflectometery experimental arrangement.



Figure 4. Reflectivity profiles for $1 \times$ bilayer (\bigcirc), $2 \times$ bilayer (\triangle), and $3 \times$ bilayer (\Box) of phthalocyanine at the silicon/ D_2O interface. The solid lines show the best fit to the data. The profiles are shifted by a factor of 10 for the purpose of clarity.

subphase. After this first measurement, the block was carefully 151 removed from the cell and a second bilayer was deposited using the 152 same procedure described above. Now those two bilayers of 153 phthalocyanine were again characterized using neutron reflecto-154 155 metery. The same procedure was followed for the deposition of the 156 third bilayer.

In a second experiment, a mixture of DPPC and DPPE carrying 157 158 a PEG group (3% w/w) monolayer were transferred on top of the 159 second C18 coated second silicon substrate, S2, as a control experiment. The deposition of the lipid mixture was also per-160 formed at a constant surface pressure of 25 mN m^{-1} and a dipping 161 speed of 5 mm/min. As only a monolayer had to be deposited, the 162 block was immersed in the aqueous phase, and then the air-water 163 interface was carefully cleaned by vacuum suction. Only then the 164 block was slowly removed from the trough. The structure of this 165 controlled monolayer was then determined. The lipid monolayer 166 was then washed away, and a bilayer of phthalocynaine followed 167 by a lipid monolayer was then deposited. In actual devices, the 168 169 nitrous oxide would be entrapped below this lipids monolayer. 170A series of neutron reflectivity profiles were then obtained to characterize these layers. 171

The C-18 layer has already been characterized using a series of 172 solvent contrasts on numerous occasions.9 The silicon oxide layer 173 174 and the C-18 hydrophobe layer were also characterized using the technique of ellipsometry (see the Supporting Information). The 175 176 layer thickness and density of the oxide layer and C18 layer were then used as a starting point in our subsequent fitting procedure. 177 178 The neutron reflectometery's experimental arrangement is shown **F3** 179 in Figure 3.

Results and Discussion

The measured reflectivity data for the C18 coated silicon block 181S1 with 1, 2, and 3 bilayers of aliphatic substituted phthalocyanine 182 with D_2O forming the aqueous subphase are shown in Figure 4. F4 183 The solid lines correspond to modeled reflectivity based on the 184



Figure 5. Scattering length density profiles for 2 (solid), 4 (long dash), and 6 (short dash) layers of phthalocyanine at the silicon-D₂O interface.

Table 1. Fitted Structural Parameters Obtained from the Fits to the **Reflectivity Profiles Shown in Figure 4**

Si (S1) substrate						
	layer	layer thickness (Å)	$NB \\ (\times 10^{-6} \text{\AA}^{-2})$	roughness (Å)		
		1× bilayer phtha	llocyanine			
1 2	C18 phthalocyanine phthalocyanine	$\begin{array}{c} 30\pm2\\ 15\pm2\\ 28\pm2 \end{array}$	0.84 4.50 5.60	2 2 7		
		$2 \times$ bilayer phtha	locyanine			
1 2 3	C18 phthalocyanine phthalocyanine phthalocyanine	30 ± 2 15 ± 1 30 ± 2 30 ± 2	-0.43 2.83 4.50 5.60	2 2 2 7		
		$3 \times$ bilayer phtha	alocyanine			
1 2 3 4	C18 phthalocyanine phthalocyanine phthalocyanine phthalocyanine	$ \begin{array}{r} 30 \pm 2 \\ 15 \pm 2 \\ 30 \pm 2 \\ 30 \pm 2 \\ 30 \pm 2 \end{array} $	$-0.43 \\ 1.46 \\ 2.83 \\ 4.50 \\ 5.60$	2 2 2 2 7		

scattering length density profiles that are shown in Figure 5. In the 185 F5 modeling of the data, the thickness and scattering length density 186 of the oxide film was found to be $d = 13 \pm 2$ Å, $Nb = 3.6 \times$ 187 10^{-6} Å⁻². The fitted structural parameters obtained from these 188 fits are given in Table 1. 189 T1

The C18 layer was found to be 30 ± 2 Å, which is slightly 190 thicker than the expected value of 24 Å reported previously9 and a 191 value of 17 ± 2 Å for a dry C18 layer coupled silicon sample 192 deduced from the ellipsometry data. This indicates an entangle-193 ment of the C18 chain with the aliphatic tail of the phthalocyanine 194 layer, hence the good anchoring characteristic of the C18 Layer. 195 Neutron reflectometery cannot distinguish between the C18 chain 196 and the aliphatic tail of the phthalocyanine because of the lack of 197 contrast between the protonated C18 and the aliphatic part of the 198 phthalocyanine which is also protonted (similar scattering length 199 density value). One striking feature of these results is the fact that 200 as the number of phthalocyanine bilayers increases, the water 201 (D_2O) content of the underlying structures decreases. This can be 202 seen in the scattering length density profiles (Figure 5). The total 203 layer thickness for the aliphatic substituted phthalocyanine for 204 the 1 bilayer sample was found to be 43 Å, for the 2 bilayers 75 Å, 205 and for the 3 bilayers 105 Å. This indicates a 13% and an 18% 206 reduction in the overall ideal layer thickness for 2 and 3 bilayer 207 samples which results from the D₂O expulsion as more layers are 208



Figure 6. (a) Reflectivity profiles for DPPC-DPPE+PEG layer and (b) 2 layers of phthalocyanine covered by DPPC-DPPE+PEG at the silicon $-D_2O$ interface. The best fits to the data are shown by solid lines.

 Table 2. Fitted Structural Parameters Obtained from the Fits to the Reflectivity Profiles Shown in Figure 6

	Si (S2) substrate				
	layer	layer thickness (Å)	$\frac{\text{NB}}{(\times 10^{-6}\text{\AA}^{-2})}$	roughness (Å)	
		C18 + DPPC m	nonolayer		
	C18 + d-lipid	19 ± 2	0.55	4	
1	lipid-chain	16 ± 2	3.30	3	
2	lipid head-PEG	60 ± 5	5.83	14	
	$C18 + 1 \times bila$	ayer phthalocyar	ine + DPPC me	onolayer	
	C18	30 ± 2	-0.12	2	
1	phthalocyanine	15 ± 1	1.22	2	
2	phthalocyanine	30 ± 1	2.32	2	
2	lipid-chain	16 ± 1	3.48	5	
3	lipid head-PEG	60 ± 5	5.83	20	

deposited. This is also supported by the scattering length densityprofiles (Figure 5).

In part one of the second experiment, a mixture of DPPC and 211 DPPE carrying a PEG group (3%w/w) was deposited as a 212 monolayer on a hydrophobic Si (C18 coated) surface (S2). The 213 F6 214 reflectivity profile is shown in Figure 6a. The fit to the data 215 indicated a mixing of the deuterated tail of DPPC and the 216 protonated C18 chain over a 19 Å region. This confirms the earlier results for the S1 sample (i.e., the interpenetrations of the 217 aliphatic chains) and can now be seen because of better contrast. 218 A three layer model was required to fit this data: a 19 Å layer 219 consisting of a mixture of C18 and DPPC tail, a 16 Å layer 220 representing the DPPC tail region, followed by a 60 Å layer 221

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Figure 7. Scattering length density profiles for DPPC-DPPE+PEG layer (solid) and 2 layers of phthalocyanine covered by DPPC-DPPE+PEG (long dash) at the silicon $-D_2O$ interface. The box represents the phthalocyanine layer contribution to the data.

representing the large PEG and DPPC headgroup region. The 222 fitted parameters are given in Table 2. 223 T2

The lipid monolayer was then washed from the S2, and a 224 bilayer of phthalocvanine was deposited followed by a lipid mono-225 nlayer. The reflectivity data obtained are shown in Figure 6b. This 226 deposited layer sequence represents an ideal device structure. The 227 water penetration into the phthalocyanine bilayers was found to 228 be less than that in the case of the $1 \times$ bilyaer phthalocyanine 229 (S1 sample) with the same overall layer thickness. The water content 230 of the phthalocyanine bilayer is an indication of the ability of the 231 lipid monolayer to form an effective barrier between the D_2O bulk 232 and the phthalocyanine bilayer beneath. These are ideal character-233 istics required for these types of devices. The full scattering length 234 density profile for the fit is shown in Figure 7. The fitted structural 235 F7 parameters are given in Table 2. 236

Conclusions

In this experiment, we have shown the suitability of the C18 238 ligands as an anchoring layer for phthalocyanine on a solid 239 support. The thicknesses of multilayers of phthalocyanine at the 240 silicon-air interface measured with ellipsometry (see the Sup-241 porting Information) prior to the solid-liquid neutron experi-242 ment were 41, 82, and 124 Å, respectively, for 2, 4, and 6 layers, in 243 overall good agreement with the neutron data and those reported 244 in the literature.¹³ Although the layer thicknesses reported here 245 for neutron are slightly lower than these because of the inter-246 penetrations of C18 and the aliphatic substituted phthalocyanine 247 and the lack of scattering contrast between these layers. 248

The data for the phthalocyanine bilayer with the deposited lipid 249 monolayer showed almost a complete depletion of water from the 250 C18 layer in addition to a depletion of water from the phthalo-251 cyanine bilayer. This result demonstrates the effectiveness of the 252 lipid layer in partitioning (sealing) the deposited phthalocyanine 253 layers from the bulk water. This is crucial for viability of these 254 systems as the biosensors' operation relies on the entrapment and 255 subsequent release of nitrous oxide gas depending on the integrity 256 of the lipid membrane. However, some trapped water content of 257 these layers is required to maintain the sensing layer saturated 258 with nitrous molecules for effective functioning of these devices. 259 In a future experiment, we plan to use polished quartz as a 260 substrate with color incorporated into the phthalocyanine bilayer. 261 The transparent quartz will allows us to observe the color changes 262

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in the sensing layer when the lipid membrane is attacked, resulting
in the escape of the nitrous oxide gas, while simultaneously
measuring the changes to the layer structures using neutron
reflectometery.

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Supporting Information Available: Ellipsometry data obtained at the air-water interface. This material is available free of charge via the Internet at http://pubs.acs.org. 274