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Contrast-Matched Isotropic Bicelles: A Versatile Tool to Specifically Probe the Solution Structure of Peripheral Membrane Proteins Using SANS

Raphael dos Santos Morais, Olivier Delalande, Javier Perez, Liza Mouret, Arnaud Bondon, Anne Martel, Marie-Sousai Appavou, Elisabeth Le Rumeur, Jean-François Hubert, Sophie Combet

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7 Contrast-matched isotropic bicelles: a versatile tool
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11 to specifically probe the solution structure of
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15 peripheral membrane proteins using SANS.
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22 *Bondon*⁵, *Anne Martel*⁶, *Marie-Sousai Appavou*⁷, *Elisabeth Le Rumeur*^{1,2}, *Jean-François*
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49 ABSTRACT
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53 Obtaining structural information on integral or peripheral membrane proteins is currently
54 arduous due to the difficulty of their solubilization, purification, and crystallization (for X-ray
55 crystallography (XRC) application). To overcome this challenge, bicelles are known to be a
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3 versatile tool for high-resolution structure determination, especially when using solution and/or
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5 solid state nuclear magnetic resonance (NMR) and, to a lesser extent, XRC. For proteins not
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7 compatible with these high-resolution methods, small-angle X-ray and neutron scattering (SAXS
8
9 and SANS, respectively) are powerful alternatives to obtain structural information directly in
10
11 solution. In particular, the SANS-based approach is a unique technique to obtain low-resolution
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13 structures of proteins in interactions with partners by contrast-matching the signal coming from
14
15 the latter. In the present study, isotropic bicelles are used as a membrane mimic model for
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17 SANS-based structural studies of bound peripheral membrane proteins. We emphasize that the
18
19 SANS signal coming from the deuterated isotropic bicelles can be contrast-matched in 100%
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21 D₂O-based buffer, allowing us to separately and specifically focus on the signal coming from the
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23 protein in interaction with membrane lipids. We applied this method to the DYS-R11-15 protein,
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25 a fragment of the central domain of human dystrophin known to interact with lipids, and we were
26
27 able to recover the signal from the protein alone. This approach gives rise to new perspectives to
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29 determine the solution structure of peripheral membrane proteins interacting with lipid
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31 membranes and might be extended to integral membrane proteins.
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40 INTRODUCTION

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43 While approximately 26% of the human proteome is predicted to be composed of
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45 membrane proteins¹, the structures of only 685 unique membrane proteins are available². This
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47 highlights the difficulty in handling membrane proteins during their solubilization, purification
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49 and crystallization³. For even more recalcitrant membrane proteins incompatible with high-
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51 resolution methods (nuclear magnetic resonance (NMR) and X-ray crystallography (XRC)),
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53 small-angle X-ray and neutron scattering (SAXS and SANS, respectively) are good alternative
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3 approaches to obtain the structural information of a protein not only alone in solution but also in
4 interaction with its partner(s)^{4,5}. In the SANS-based approach, the contrast matching (CM)
5 technique allows the signal coming from one component to be specifically probed by hiding the
6 signal coming from other partners in a multicomponent system. This is achieved by adjusting the
7 deuteration level of the components in an appropriate H₂O/D₂O ratio. The CM technique has
8 been successfully applied to the determination of the structure of one component in protein-
9 protein and protein-DNA complexes⁶. Protein-lipid complexes have also been successfully
10 investigated in the same way with different membrane mimic environments⁷. In an elegant study,
11 Gabel *et al.*⁸ investigated the conformation of the β -barrel transporter FhaC surrounded by a β -
12 octyl glucoside corona by coupling SANS and molecular modeling, obtaining a conformation of
13 the protein similar to that obtained by XRC. Although detergent micelles are considered suitable
14 for maintaining the tridimensional structure of membrane proteins, some authors⁹ proposed that
15 the loss of activity evidenced for some membrane proteins could be inherent to the lack of their
16 internal dynamics in detergent micelles compared with bilayered membrane mimics. Indeed,
17 membrane models other than micelles are available and the interest for reconstructing a
18 biological native lipid environment for membrane proteins is growing¹⁰⁻¹². Among them are the
19 classical liposome model but also relatively more recent ones, such as nanodiscs and bicelles,
20 which can be good alternatives¹⁰⁻¹². However, all of these membrane mimics have their
21 advantages and drawbacks¹⁰⁻¹².

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49 Nanodiscs are phospholipid bilayers surrounded by either a membrane scaffolding protein
50 (MSP)¹³ or a styrene maleic acid (SMA) copolymer¹⁴. By using these membrane mimics, Maric
51 *et al.*¹⁵ proposed an interesting method to use SANS to analyze integral membrane proteins
52 incorporated into contrast-matched nanodiscs. In the case of peripheral membrane proteins, it is
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3 essential to maintain a phosphatidylcholine head all over the membrane mimic to avoid non-
4 specific interactions that could happen between the scaffold of the nanodisc and the protein of
5 interest. This can be achieved by handling peripheral proteins with fully phospholipid-based
6 bicelles. Moreover, the size of bicelles can be easily adjusted and does not require the use of
7 several MSP truncated forms^{16,17}. Last but not least, in addition to the intrinsic highly curved rim
8 region, few curvature-inducing molecules can be incorporated into bicelles in order to modify the
9 planarity of the bilayer part¹⁸. These properties reinforce the suitability of the bicelle model for
10 mimicking membrane bending, known to be essential for various biological processes¹⁹. These
11 three points constitute the most important advantages of phospholipid-based bicelles compared
12 to nanodiscs.
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28 Bicelle aggregates are made of long-chain phospholipids arranged in a bilayer surrounded
29 by a torus of short-chain phospholipids or detergents. Pioneering studies describing such objects
30 were performed on lecithin/bile-salt or analogues^{20,21} and
31 dipalmitoylphosphatidylcholine/diheptanoylphosphatidylcholine (DPPC/DH₇PC) solution^{22,23}.
32 To better mimic the membrane bilayer, unsaturated phospholipids, cholesterol^{24,25}, or anionic
33 phospholipids²⁶ were successfully incorporated into the bilayer part. Recently, bicelles were even
34 constructed from native *Escherichia coli* lipids²⁷, but the currently best-described system is made
35 of dimyristoylphosphatidylcholine/dihexanoylphosphatidylcholine (DMPC/DHPC) since its first
36 characterization by solid-state NMR²⁸. The molar ratio (DMPC/DHPC), denoted “*q*”, of the two
37 components forming the bicelles is the major parameter governing the bicelle size. This
38 parameter can be easily adjusted according to the desired final size for a required application. Its
39 value distinguishes two types of bicelles. The larger ones ($q \geq 2.5$) are able to align in a magnetic
40 field above the gel-to-fluid phase transition temperature (T_m). They are commonly used in solid
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3 state NMR^{29,30} and increasingly in XRC³¹⁻³³, while existing as perforated lamella rather than
4 disks in these conditions³⁴. The other type, small isotropic bicelles ($q < 2.5$), are commonly used
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6 in solution NMR^{29,30}. Despite their versatility, the morphology and the size of the bicelles are
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8 reported to be highly sensitive to q , temperature, and dilution^{35,36}. DMPC/DHPC bicelles were
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10 thoroughly investigated with cryo-electronic microscopy, dynamic light scattering (DLS) and
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12 NMR³⁵⁻³⁹, but only a few studies were performed with SANS⁴⁰⁻⁴⁴. The study by Luchette *et al.*⁴⁰
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14 is, to our knowledge, the only one devoted to SANS characterization of hydrogenated
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16 DMPC/DHPC isotropic bicelles.
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23 In the present work, we analyzed both hydrogenated and deuterated isotropic bicelles (h-
24 bicelles and d-bicelles, respectively) in the range of $1 \leq q \leq 1.3$. We demonstrate that the d-
25 bicelles can be contrast-matched in 100% D₂O-buffer regardless of their morphology and
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27 temperature. We applied this method to a protein fragment of the dystrophin central domain,
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29 DYS R11-15, known to interact with the lipid membranes⁴⁵⁻⁴⁷. We highlight that one can
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31 specifically probe the signal of a peripheral membrane protein bound to bicelles. In addition, we
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33 show that SANS analysis can be properly achieved on bicelles previously submitted to high-
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35 pressure size exclusion chromatography.
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43 MATERIALS AND METHODS

44 *Materials*

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49 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC), 1,2-dihexanoyl-sn-glycero-3-
50 phosphocholine (DHPC), 1,2-dimyristoyl-d54-sn-glycero-3-phosphocholine-1,1,2,2-d4-N,N,N-
51 trimethyl-d9 (DMPC-d67), and 1,2-dihexanoyl-d22-sn-glycero-3-phosphocholine-1,1,2,2-d4-
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3 N,N,N-trimethyl-d9 (DHPC-d35), conditioned in chloroform, are from Avanti Polar Lipids and
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5 were used without any further purification. D₂O, Tris-d11, and EDTA-d16 are from Eurisotop.
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8 9 *Bicelle preparation*

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11 A chloroform solution containing the appropriate amounts of DMPC/DHPC or DMPC-
12 d67/DHPC-d35, to obtain a ratio (mol/mol) $q = 1$, was dried overnight under vacuum. The lipid
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14 mixture was then rehydrated in d-TNE buffer solution (20 mM Tris-d11, 150 mM NaCl, and
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16 0.1 mM EDTA-d16, pD 7.5) or TNE buffer to reach a total lipid concentration of at least
17
18 200 mM. Once rehydrated, the solution was frozen (10 s in liquid N₂), thawed (10 min at 40°C),
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20 vigorously shaken with a vortex (1 min), and then centrifuged (1.5 min, 6,000 rpm, MiniSpin,
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22 Eppendorf). This procedure was repeated two more times to homogenize the solution. Stock
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24 solutions were diluted in d-TNE or TNE based on needs.
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31 32 *The ideal bicelle model (IBM)*

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34 Isotropic bicelles are considered to be disk-shaped objects made of long-chain phospholipids
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36 forming a bilayer, surrounded by short-chain phospholipids located at the rim (Figure 1A). The
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38 molar ratio of the two phospholipids, denoted “ q ” (not to be confused with “ Q ” used here for the
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40 scattering vector in SANS and DLS experiments), is the main parameter governing the size of
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42 the bicelles. The effective molar ratio is denoted “ q_{eff} ”^{35,37} when the proportion of free DHPC is
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44 considered, and is defined by
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$$50 \quad q_{eff} = \frac{[DMPC]}{[DHPC]_{total} - [DHPC]_{free}} \quad \text{Eq 1}$$

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55 Several IBMs are proposed to correlate q_{eff} to the radius of the bicelle. These models are based
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57 either on head group area^{30,43} or on phospholipid volume⁴⁹. We consider the latter model to be
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3 the most reliable one since the molecular volumes of phospholipids are barely different below
4 and above T_m ^{50,51} and the equation does not depend on the thickness of the bilayer. The radius of
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6 IBM can be expressed as^{38,49} (Figure 1)
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$$R = r_{\perp} + \frac{r_{\perp} q_{eff}}{4\Lambda} \left[\pi + \left(\pi^2 + \frac{32\Lambda}{3q_{eff}} \right)^{1/2} \right] \quad \text{Eq 2}$$

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15 where R is the radius of the bicelle, r_{\perp} is the thickness of the rim (11 Å, the length of a DHPC
16 molecule), Λ is the volume ratio of DHPC to DMPC (0.61), and t is the thickness of the bilayer
17 (assumed to be 40 Å)⁴⁹. According to Small⁵², the volumes of DMPC and DHPC are 1090 and
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19 660 Å³, respectively.
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25 26 *³¹P NMR spectroscopy*

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29 ³¹P NMR spectra were recorded on a Bruker spectrometer Avance 500 equipped with a 5-mm
30 BBO probe operating at 202.46 MHz. Spectra of 10 kHz spectral width and 32 K data points
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32 were acquired with proton decoupling using 128 scans, a 30° flip angle, and a 0.5 s relaxation
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34 delay. The data were processed with the TopSpin3.2 software (Bruker). Before applying the
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36 Fourier transform, free induction decays were treated with an exponential broadening of 2 Hz.
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39 85 % H₃PO₄ was used as an external standard for the ³¹P chemical shift.
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44 45 *Dynamic light scattering*

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47 The monodispersity and the hydrodynamic radius (R_h) of the bicelles were estimated by DLS
48 using a Zetasizer instrument (Nano ZS, Malvern Instruments). Measurements were done in a
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50 temperature range from 15 to 37°C in low volume Hellma cells (100 μL). The data were
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52 processed with the Zetasizer software v7.11 with default parameters and characterized by the
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54 size distribution by intensity. The decay rate, Γ , of the autocorrelation function is linked to the
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3 diffusion coefficient D by $\Gamma = DQ^2$, where Q , the momentum transfer, is defined as $Q =$
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5 $\frac{4\pi n \sin \theta}{\lambda}$, where n is the refractive index of the medium, λ the wavelength of the laser, and 2θ the
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7 scattering angle. R_h is determined with the Stokes-Einstein equation: $R_h = \frac{k_B T}{6\pi\eta D}$, where k_B is the
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9 Boltzmann constant, T the temperature, and η the viscosity of the medium. Data were processed
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11 using the viscosity of heavy water according to the temperature⁵³.
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17 *DYS R11-15 protein purification*

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20 The DYS R11-15 protein was expressed and purified as previously described⁴⁵. The purity was
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22 assessed by SDS-PAGE (Figure S3A) and Coomassie blue staining (InstantBlue, Expedeon).
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24 TNE to d-TNE buffer exchange was performed with Amicon Ultra-15 (MWCO 10 kDa). Exactly
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26 the same buffer was used for both bicelle rehydration and SANS acquisition to ensure a perfect
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28 buffer subtraction in all cases.
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32 *Intrinsic tryptophan fluorescence*

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35 Fluorescence measurements were obtained on a Fluorolog spectrofluorometer (Jobin-Yvon).
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37 Tryptophan fluorescence emission spectra were recorded at 20°C in low volume quartz cuvettes
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39 (120 μ L) between 310 and 420 nm using an excitation wavelength of 295 nm (bandwidth 2 nm)
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41 in TNE buffer. The protein concentration was 10 μ M and the lipid concentration was 50 mM.
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46 *Chromatographic co-elution*

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49 Data were collected using a Bio SEC-5 500 Å column (5 μ m, 4.6 mm x 300 mm, Agilent)
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51 mounted on an ÄKTA Explorer HPLC system (GE Healthcare). The column was equilibrated
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53 with TNE buffer supplemented with 6 mM of DHPC to avoid bicelle deformation under diluted
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55 concentration. A sample volume of 50 μ L was loaded onto the column. For DYS R11-15 alone,
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3 the protein concentration was 60 μM and for the bicelles alone, the lipid concentration was
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5 50 mM. For the lipid-protein complex, the concentrations were 94 μM for the protein and
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7 50 mM for the lipids. The flow rate was 0.2 mL/min.
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10 11 *Small angle neutron scattering experiments*

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14 SANS data were recorded on either the PACE⁵⁴ (LLB, Saclay), KWS1^{55,56} (MLZ, Garching), or
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16 D22⁵⁷ (ILL, Grenoble) SANS instruments. According to the predicted size of the bicelles, two or
17
18 three sample-to-detector distances were used with a wavelength varying from 4.7 to 6 \AA , to
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20 cover a Q -range from 0.005 to 0.5 \AA^{-1} for the largest one, where $Q = \frac{4\pi \sin \theta}{\lambda}$ is the momentum
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22 transfer, λ is the wavelength, and 2θ is the scattering angle. All measurements were done in
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24 1 mm Hellma Suprasil quartz (QS) cells.
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28 29 30 *High-pressure size exclusion chromatography-SANS*

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33 High-pressure size exclusion chromatography (HPSEC)-SANS data were acquired on the D22
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35 instrument (ILL, Grenoble) with the single configuration of 8 m collimation and sample-detector
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37 distance, at $\lambda = 6 \text{\AA} \pm 10\%$, covering a Q -range from 0.008 to 0.17 \AA^{-1} . The sample environment
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39 was described elsewhere⁵⁸. Data were collected using a Bio SEC-5 500 \AA column (5 μm , 4.6 mm
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41 x 300 mm, Agilent) equilibrated with the d-TNE buffer supplemented with 6 mM of. A volume
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43 of 100 μL of bicelle solution at 50 mM was loaded onto the column. The flow rate was 0.2
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45 mL/min. The SANS signal of the buffer was collected in-line before the void volume, allowing a
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47 perfect subtraction to be done.
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51 52 *SANS data analysis*

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55 SANS data were analyzed with the ATSAS suite⁵⁹, following its guidelines unless otherwise
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3 indicated. The PRIMUS software⁶⁰ was used to estimate the dimensions of the bicelles. At small
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5 Q values, the Guinier approximation (for $QR_g < 1.3$) was used to determine the forward intensity
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7 $I(0)$ and the radius of gyration R_g and is defined by⁶¹
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$$I(Q) = I(0) \exp\left(-\frac{Q^2 R_g^2}{3}\right) \quad \text{Eq 3}$$

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15 At intermediate Q values, for a disk-shaped object, the Guinier approximation allows the cross-
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17 sectional radius of gyration R_t to be approximated and is defined by⁶¹
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$$I(Q) = \frac{I(0)}{Q^2} \exp(-Q^2 R_t^2) \quad \text{Eq 4}$$

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22 For a disk-shaped object of homogenous SLD contrast, R_t and R_g are related to the thickness t
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24 and the radius R of the disk by⁶¹
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$$R_t^2 = \frac{t^2}{12} \quad \text{and} \quad R_g^2 = \frac{R^2}{2} + \frac{t^2}{12} \quad \text{Eq 5 and 6}$$

30 31 32 33 34 35 *SANS data fitting parameters*

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38 For our calculations, we used the volumes of DMPC and DHPC from Small⁵² (1090 and 660 Å³
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40 respectively). The volume of the tail, excluding the carbons belonging to the carbonyl functions,
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42 was estimated using the empirical rule of Tanford⁶². Then, the volume of the lipid head was
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44 obtained simply by subtracting the volume of the tail from the total volume. The densities were
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46 estimated from the molecular formula and the volumes. The scattering length density (SLD) was
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48 calculated using the NIST website⁶³. A summary table is available (Table S1). Data were fitted
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50 with the “core-shell cylinder” model⁶⁴ (Figure 1B), described in the supporting information (Eq.
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60 S1, S2, and S3). We fixed the following parameters to fit SANS data: the thicknesses of the shell

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3 (t_s) and of the core (t_c) were 6 and 28 Å, respectively. The SLD of the solvent was $6.34 \cdot 10^{-6} \text{ \AA}^{-2}$.
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5 The thickness of the core had to be released in order to fit the data of d-bicelles at 37°C. All data
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7 fitting procedures for form factor determination were done using the SASview v3.1.0 software⁶⁵.
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10 RESULTS AND DISCUSSION

11 *Characterization of the h-bicelles*

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18 Beaugrand *et al.*³⁷, in analogy with the critical micellar concentration for the micelles, introduced
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20 the critical bicellar concentration (CBC), which corresponds to the concentration of free DHPC
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22 in a bicelle solution³⁵. They also defined the concentration boundaries to maintain the bicelle size
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24 for a defined q ratio. Mineev *et al.*³⁸ recently proposed to determine the concentration of free
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26 DHPC by NMR diffusion, as previously done by Chou *et al.*⁶⁶, and to dilute the sample in a
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28 buffer containing the same DHPC concentration in order to maintain the expected q ratio and
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30 thus the bicelle size under diluted conditions. We decided to take advantage of the free DHPC in
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32 solution to obtain bicelles with a molar ratio, q_{eff} , ranging from 1 to 1.3 by assuming a CBC
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34 value of 6 mM throughout this study³⁷. This value is known to be almost constant in our q_{eff} -
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36 range and in a wide range of temperatures³⁷. Note that all concentrations hereafter indicate the
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38 total lipid concentration where the CBC is not subtracted. The q_{eff} -range ($1 \leq q_{eff} \leq 1.3$) was
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40 chosen since the morphology of small isotropic bicelles ($q_{eff} < 1$) is still largely debated. Below
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42 this threshold, the objects would be present as mixed-micelles³⁷ or as real disk-shaped
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44 objects^{38,39}. First, we checked the specific organization in bicelles and the correct molar ratio of
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46 our preparation by analyzing a concentrated sample (200 mM, $q = 1$) by ³¹P NMR (Figure 2A).
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48 The spectrum is characteristic of isotropic bicelles with a narrow peak and a broader one
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50 attributed to DHPC and DMPC, respectively²⁸. The molar ratio of 1, calculated from the
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3 integrals of the two peaks, is in line with the $q = 1$ expected value. Since the morphology and the
4 size of the bicelles may be affected by q_{eff} ^{36,67}, less concentrated samples diluted with the d-TNE
5 buffer, at 100, 75, and 50 mM of lipids (Figure 2B), were analyzed by SANS at 20°C, below the
6 gel-to-fluid phase transition temperature of DMPC (24°C)⁶⁸. For these three concentrations, a
7 plateau is reached, which is characteristic of objects with a finite size. The radii of gyration (R_g)
8 obtained for the three concentrations are 25.5 ± 0.2 , 28.0 ± 0.3 , and 33.5 ± 0.4 Å. As expected,
9 the bicelles size increases with q_{eff} (Figure 2B). On the other hand, the intermediate radius of
10 gyration (R_t) remains constant at 12.0 ± 0.1 Å. For a disk-shaped object of homogeneous SLD
11 contrast, the R_g values correspond to geometric radii of approximately 32, 36, and 44 Å, while
12 the R_t value corresponds to a thickness (t) of approximately 41.5 Å, in line with the dimension of
13 a DMPC bilayer in gel phase⁶⁹. We also analyzed the stock solution (265 mM) and observed a
14 structure peak (Figure S1) associated with an interparticle spacing previously observed by
15 Luchette *et al.*⁴⁰ in similar conditions. This peak is no longer observable for the more diluted
16 samples, down to 50 mM. More diluted samples were also analyzed (25 and 10 mM) and, as
17 expected, the samples are no longer disk-shaped objects at these concentrations since q_{eff}
18 increases strongly (data not shown). Finally, to confirm that the bicelles at 50 mM are effectively
19 discoidal objects, data obtained at this concentration were fitted with the “core-shell cylinder”
20 form factor model⁶⁴. This two-step SLD cylinder model allows the tails and the heads of the
21 phospholipids to be distinguished. We assume that the SLD for DMPC and DHPC heads are
22 identical since they have the same chemical function. This affirmation might not be absolutely
23 true because differences in the level of hydration between the bilayer and the rim could modify
24 the SLD. Indeed, the areas occupied by the heads of both phospholipids are known to be
25 different in the case of a DMPC bilayer in gel (47 Å²)⁶⁹ or in fluid phase (60 Å²)⁷⁰ or for DHPC
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3 micelles (100 \AA^2)⁷¹. We also fixed the thickness of the core and of the shell to 28 and 6 \AA ,
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5 respectively, to reduce the number of free parameters during the fitting procedure. Despite these
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7 approximations, good fits were obtained (Figure 2B) using this simplified model for h-bicelles,
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9 giving overall dimensions of 40 \AA for the thickness (t) and 42 \AA for the total radius (R). These
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11 results are in line with the first Guinier approximations and the IBM proposed by Tryba *et al.*⁴⁹.
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13 Altogether, these data show that the experimental conditions used for the h-bicelles led to disk-
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15 shaped objects. Therefore, the same experimental conditions were used in the following
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17 experiments with the d-bicelles.
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22 *Deuterated bicelles are contrast-matched in 100% D₂O buffer*

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26 We used almost fully deuterated DMPC and DHPC lipids, commercially available, to form the d-
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28 bicelles (five hydrogen atoms remain near the glycerol group). The d-bicelles at 50 mM were
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30 analyzed at three contrasts: 42, 70, and 100% D₂O/H₂O ratios (Figure 3). First, to probe the
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32 expected disk-shaped morphology, data from the d-bicelles at 42% D₂O (corresponding to the
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34 classical contrast-matching point of hydrogenated proteins) was used for further analysis after
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36 incoherent subtraction (Figure 3). We used the Guinier approximation to estimate the overall
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38 dimensions of these disk-shaped objects. We obtained R_g and R_t values of 31.5 ± 0.1 and 12.5
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40 $\pm 0.1 \text{ \AA}$, respectively, corresponding to a radius (R) of approximately 41 \AA and a thickness (t) of
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42 43 \AA . Then, the data for d-bicelles were fitted with the same model and assumptions as for the h-
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44 bicelles. A good fit was obtained (Figure 3) by using the “core-shell cylinder” model, giving
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46 overall dimensions of 40 \AA for the thickness (t) and 44 \AA for the total radius (R), in-line with the
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48 Guinier approximation and the theoretical IBM.
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3 According to our calculations, the SLD for DMPC-d67 and DHPC-d35 are 6.65 and $6.20 \cdot 10^{-6} \text{ \AA}^{-2}$,
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According to our calculations, the SLD for DMPC-d67 and DHPC-d35 are 6.65 and $6.20 \cdot 10^{-6} \text{ \AA}^{-2}$, respectively, and are close to the SLD of 100% D₂O-based buffer of $6.34 \cdot 10^{-6} \text{ \AA}^{-2}$. Therefore, considering SANS resolution, d-bicelles should be nearly contrast-matched in 100% d-TNE buffer. As expected, d-bicelles at 100% D₂O are virtually contrast-matched in SANS, with a small residual signal at small angles, which is negligible (Figure 3, insert). The signal of the d-bicelles in 70% D₂O is more than 20-fold higher and almost two orders of magnitude higher in 42% than in 100% D₂O. By plotting the square-root of SANS intensity as a function of the percentage of D₂O, we obtained for the data above a match-point of 104% D₂O (Figure S1A). To compare with our experimental data, theoretical curves were generated for a Q -range from $2.5 \cdot 10^{-2}$ to 0.5 \AA^{-1} with the SASview software, using the above parameters obtained for the dataset at 42% D₂O, in 0, 20, 42, 70, 90, and 100% D₂O (Figure S1C). The theoretical match-point of the d-bicelles was determined with the same plot as above (Figure S1B) and is around 99% D₂O, in-line with the experimental determination. Considering experimental and theoretical data, 100% D₂O is assumed to be a reliable match-point value. We concluded that the d-bicelles are virtually contrast-matched in 100% d-TNE buffer, whose salt concentration and pH are relevant for biological applications⁴⁶. These first analyses were performed at 20°C. The gel-to-fluid transition temperatures of DMPC and DMPC-d54 are 24 and 20°C^{72,73}, respectively, and the acute control of the bicelle size and morphology is a key parameter in these studies. Therefore, we decided to explore structural modification of the isotropic bicelles depending on temperature since this was previously achieved in other systems^{34,36}.

Thermal effects on bicelle morphology and contrast-matching

To ensure that the studied membrane mimic objects were disk-shaped, we performed a screening of temperature effects on both the h- and d-bicelles using DLS. Measurements were performed

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3 from 15 to 37°C (Figure 4A). The figure highlights the hydrodynamic radius (R_h) variation of the
4 bicelles according to temperature. For the h-bicelles, R_h decreases from 4.9 to 3.7 nm when the
5 temperature rises from 15 to 24°C (T_m) and then progressively increases up to a plateau value of
6 6.2 nm at temperatures above 30°C. The d-bicelles exhibited the same qualitative behavior, with
7 R_h values of 5.1, 4.4, and 7.0 nm at 15, 20 (T_m), and above 30°C. These results clearly highlight
8 that structural modifications occur above T_m , as previously reported in other studies^{34,36,39}. Next,
9 we explored by SANS the effect of the temperature on d-bicelle morphology in 42% d-TNE at
10 22°C, two degrees above the gel-to-fluid transition temperature, and at 37°C, which is the
11 physiological temperature (Figure 4B). In correlation with our DLS data, the morphology of d-
12 bicelles is clearly modified with temperature. Not surprisingly, this change occurs close to T_m . At
13 22°C, good fits were obtained with the “core-shell cylinder” model with a radius (R) of 42 Å and
14 a thickness (t) of 40 Å (Figure 4B) and we can conclude that disk-shaped objects remain present
15 at least two degrees above T_m . On the other hand, no suitable fits with the same parameters were
16 obtained for data acquired at 37°C. Instead, at 37°C, good fits were obtained by keeping the
17 same model but by changing the initial parameters and unfixing the thickness of the core in order
18 to design an elongated cylinder (Figure 4B). The calculated values for the cylinder were 26 Å for
19 the total radius (R) and 192 Å in thickness (t). The same observations were done for the h-
20 bicelles (data not shown). We conclude that elongated mixed-micelles do appear at this
21 temperature, as reported in previous studies^{36,43}.

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49 As shown in Figure 4B, the SANS signal observed for the d-bicelles in 100% d-TNE is at least
50 two orders of magnitude lower than that in 42% TNE. Thus, despite the change in morphology,
51 deduced from data of d-bicelles in 42% d-TNE, we conclude that full contrast-matching is
52 obtained for d-bicelles in 100% d-TNE.
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3 To sum up, whatever the form adopted by the bicelles, we show that they can be contrast-
4 matched. This novel approach may be designed to specifically probe the SANS signal of a
5 protein in interaction with membrane lipids. The morphological modifications should, however,
6 be considered for any further investigations, particularly if one wants to use molecular dynamics
7 simulations to model the entire protein/bicelle complex.
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16 *Experimental application: DYS-R11-15 in interaction with d-bicelles*
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19 To experimentally validate this approach and to specifically probe the signal coming from a
20 protein bound to membrane lipids, we applied the method to a protein-lipid complex. This
21 complex is made of an amphipathic peripheral protein bound to a bicelle. For the protein part, we
22 used a protein fragment belonging to the dystrophin central domain DYS R11-15 (from the 11th
23 to the 15th spectrin-like repeats) for which both stability alone in solution and strong interaction
24 with lipids have been thoroughly described⁴⁵⁻⁴⁷. Although the three-dimensional structure of
25 R11-15 alone is not accessible by NMR and XRC due to its size (60 kDa) and its flexibility, an
26 all-atom model was recently proposed by coupling SAXS and molecular modeling⁷⁴. For the
27 bicellar part of the complex, we used DHPC/DMPC h-bicelles ($q_{eff} = 1.3$, 50 mM). We examined
28 the interaction of the protein with the h-bicelles, in the same experimental conditions as for
29 further SANS analysis, by performing intrinsic tryptophan fluorescence measurements and
30 chromatographic co-elution (Figure S3B and S3C). Then, we analyzed the DYS R11-15 protein
31 alone or in interaction with the d-bicelles by SANS (Figure 5). We used exactly the same buffer
32 and the same batch of bicelles in the samples of bicelles alone and DYS R11-15/bicelle
33 complexes. Data were recorded at 93 μ M (5.6 g/L) of DYS R11-15 in 100% d-TNE buffer at
34 18°C. The specific SANS data were obtained following buffer or d-bicelles signal subtraction for
35 the protein alone or in the presence of d-bicelles, respectively, and after incoherent background
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3 subtraction. In experiments designed to probe three-dimensional modifications of peripheral
4 proteins bound to lipids, it is fundamental to avoid any signal coming from the peripheral
5 proteins alone in solution. Within the concentrations used in our experiments presented in Figure
6 5, each protein in solution is bound to at least one bicelle, as concluded from chromatographic
7 co-elution (Figure S3C). SANS raw data of the dystrophin in absence and presence of bicelles
8 are perfectly superimposed, as shown in the insert on Figure 5. From the Guinier approximation,
9 the radii of gyration are 56.2 ± 0.8 and 55.9 ± 0.8 Å for the protein alone and for the protein in
10 the presence of contrast-matched d-bicelles, respectively. In the case of large conformational
11 changes due to lipid binding, R_g would have been affected and the scattering curves would be
12 different. Under the conditions of this experiment, we conclude that the native tertiary structure
13 of the protein is maintained when bound to bicelles or that no conformational modification is
14 detectable by SANS. A study of conformational modification of dystrophin fragments bound to
15 bicelles in other conditions is beyond the scope of the present paper. To determine whether the
16 method would be suitable for experiments requiring higher amounts of bicelles, in additional
17 experiments, more concentrated samples of d-bicelles alone were also analyzed. We observed
18 that their residual signal remains negligible (Figure S1). According to the IBM and using a CBC
19 of 6 mM, bicelles with $q_{eff} = 1.3$ at 50 mM, $q_{eff} = 1.1$ at 100 mM, and $q_{eff} = 1$ at 225 mM of lipids
20 correspond approximately to 130, 350, and 920 μM of bicellar objects, respectively. These
21 concentrations of bicellar objects seem large enough to ensure that all proteins are bound to at
22 least one contrast-matched bicelle in the classical protein concentration range used in SANS (5-
23 10 g/L).

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54 *Going further in bicelle analysis: bicelles in HPSEC-SANS*
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3 All the SANS data presented above were acquired in a standard sample environment (Hellma
4 cell) since the DYS-R11-15 is a quite stable protein. Some less stable membrane proteins tend to
5 form aggregates that would dramatically affect the SANS signal. To overcome this, we
6 attempted to explore the behavior of bicelles in a size exclusion chromatography system, with
7 the aim of targeting the SANS acquisition on the appropriate membrane mimic system, either
8 alone or putatively in interaction with a partner. The results in Figure 6 show that the
9 DMPC/DHPC h-bicelles are compatible with an on-line HPLC system such as the one available
10 on D22⁵⁸ (ILL, Grenoble) since they are eluted as a single peak. The elution of bicelles can be
11 followed thank to their absorbance at 210 nm and does not need the incorporation of a probe as
12 previously described with LC experiments performed on DMPC/DPC (dodecylphosphocholine)
13 bicelles¹⁸ (the new ILL-D22 SEC-SANS set up enables to record absorbance at up to 4 different
14 wavelengths). Interestingly, SANS data acquired on samples from the top of the elution peak
15 were perfectly fitted with the “core-shell cylinder” model⁶⁴, highlighting that bicelles maintain
16 their disk-shaped conformation after travelling through the HPSEC column. We could not
17 analyze the d-bicelles in the same way because of the prohibitive cost of the d-DHPC-
18 supplemented elution buffer. Using h-DHPC instead does not allow the d-bicelles to remain
19 contrast-matched due to the rapid exchange between the d-DHPC from the bicelle rim and the
20 free h-DHPC present in the buffer (data not shown). Nevertheless, supplementing the buffer with
21 d-DHPC should allow the d-bicelles to remain contrast-matched throughout an HPSEC-SANS
22 measurement. Thus, this system could be applied to the characterization of other peripheral
23 proteins and even integral membrane proteins in a bicellar environment, as previously done in
24 NMR and XRC.
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56 CONCLUSION
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3 In the present paper, we show that isotropic hydrogenated and deuterated bicelles can be
4 characterized by SANS and finely controlled in size for biological applications. We demonstrate
5 that the isotropic d-bicelles can be contrast-matched in SANS experiments using a 100% D₂O-
6 based buffer at physiological pH and salt concentrations. Moreover, we highlight that the signal
7 coming from a peripheral protein bound to the d-bicelles can be separately and specifically
8 probed by SANS. As a reminder, deuterated phospholipids used in the present study are
9 commercially available enabling an easy preparation of d-bicelles with a determined size.
10 Thereby, our method may represent an advantage compared to the deuterated nanodiscs, for
11 which MSP purification in deuterated form might be tricky. These contrast-matched d-bicelles
12 give rise to a wide range of biological applications to separately and specifically probe the
13 solution structure of peripheral proteins and even fully integral membrane proteins in interaction
14 with membrane lipids.
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32 ASSOCIATED CONTENT

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36 SANS data of h- and d-bicelles at several concentrations; SDS-PAGE analysis of DYS R11-15,
37 tryptophan fluorescence spectra of the protein with or without h-bicelles and chromatographic
38 co-elution analysis. This material is available free of charge via the Internet at
39 <http://pubs.acs.org>.
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3 The authors declare no competing financial interest. Schemes of the proteins in the graphical
4 abstract were drawn from [MyDomains - Image Creator](#). The bicelle models were generated by
5 molecular modeling^{75,76}.
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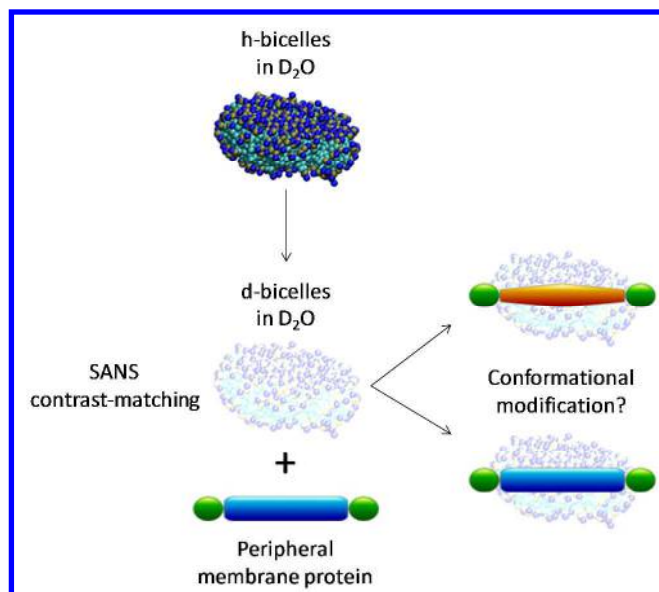
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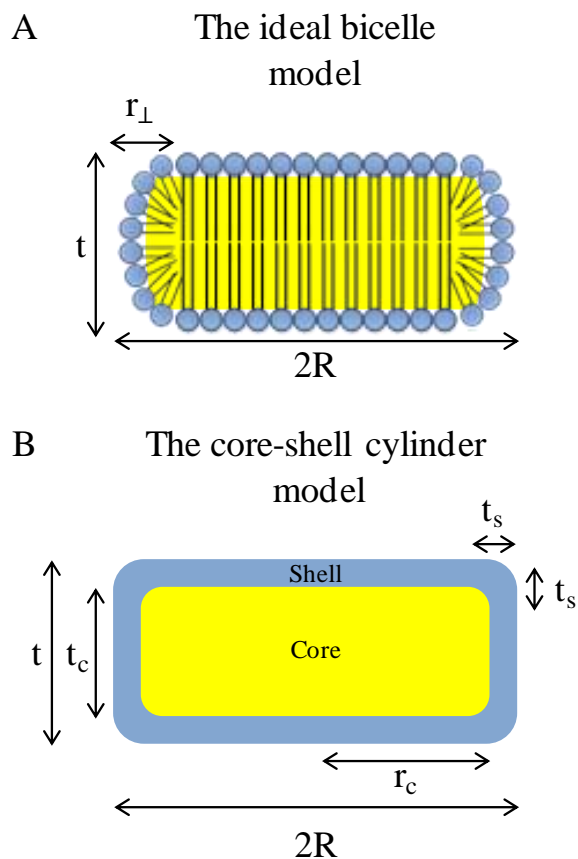
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33 Figure 1: (A) Schematic representation of the ideal bicelle model. For DMPC/DHPC bicelles,
34 R is the radius of the bicelle, r_{\perp} is the thickness of the rim (11 \AA , the length of a DHPC
35 molecule), and t is the thickness of the DMPC bilayer (assumed to be 40 \AA)⁴⁹. (B) Schematic
36 representation of the “core-shell cylinder” model, where t_c is the thickness of the core, r_c is
37 the radius of the core, and t_s is the thickness of the shell. As described in the materials and
38 methods, the calculated scattering length densities of the core and of the shell are,
39 respectively, -0.58 and $2.24 \cdot 10^{-6} \text{ \AA}^{-1}$ for h-bicelles and 7.39 and $5.05 \cdot 10^{-6} \text{ \AA}^{-1}$ for d-bicelles.
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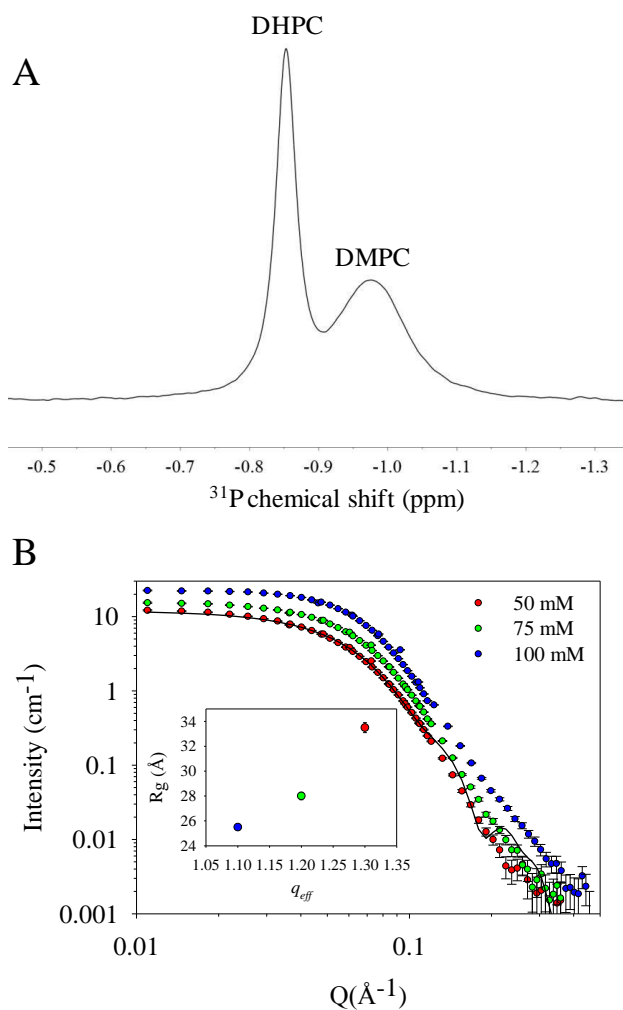


Figure 2: (A) ^{31}P NMR spectrum of $q = 1$ h-bicelles at 200 mM, typical of a bicelle organization. (B) SANS intensities measured for $q = 1$ h-bicelles at 20°C for a total DMPC/DHPC concentration of 100 ($q_{\text{eff}} = 1.1$) (blue), 75 ($q_{\text{eff}} = 1.2$) (green), and 50 ($q_{\text{eff}} = 1.3$) (red) mM in 100% D_2O d-TNE buffer. Data at 50 mM were fitted (black line) with the “core-shell cylinder” model. The thickness (t) is 40 \AA and the radius (R) is 42 \AA , in line with the dimensions of a disk. Insert: R_g as a function of q_{eff} showing the increased bicelle size with q_{eff} .

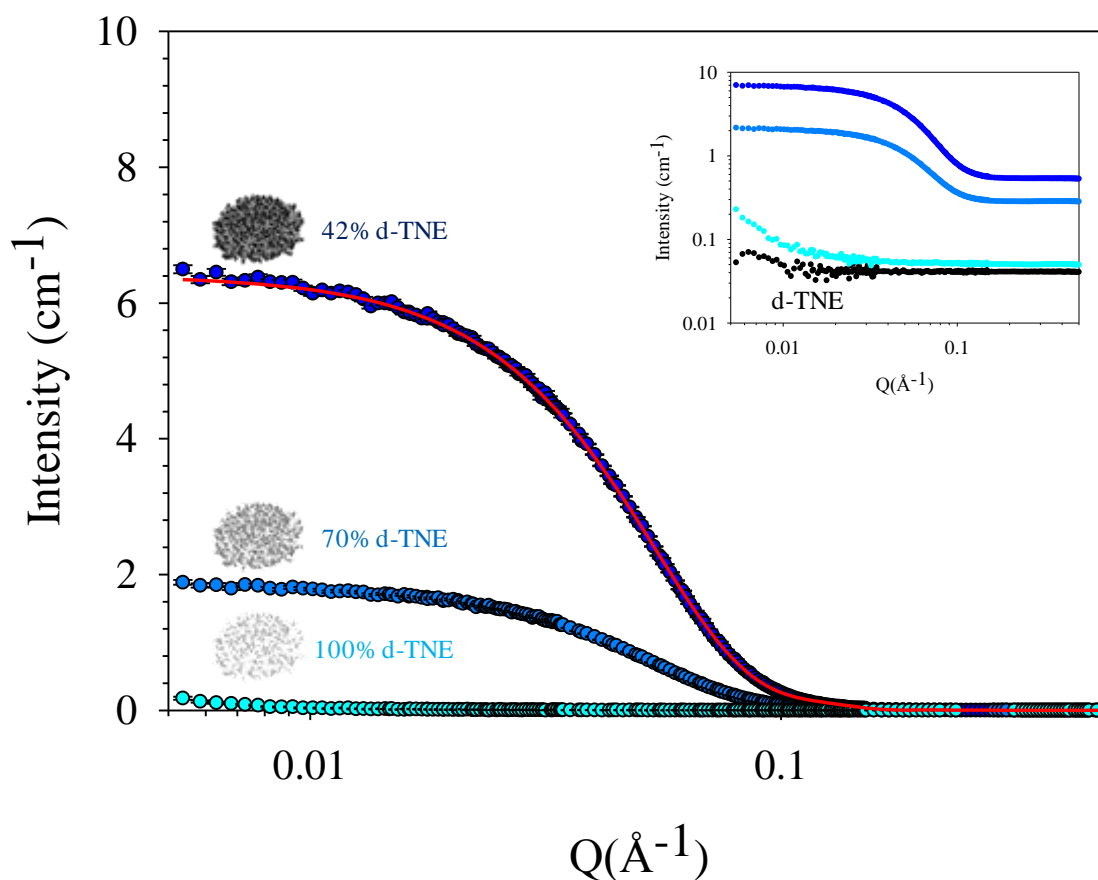


Figure 3: SANS intensities measured for $q_{eff} = 1.3$ d-bicelles at 50 mM lipids in 42% (dark blue), 70% (blue), and 100% (cyan) D₂O in d-TNE buffer (black) at 20°C. The d-bicelles in 42% d-TNE were fitted with the “core-shell cylinder” model (red line), giving 40 and 42 Å for the thickness (t) and the radius (R), respectively. The incoherent SANS signal was subtracted. Insert: the same data, without incoherent subtraction in a log/log scale, highlighting the small residual signal of d-bicelles at very small Q -values and compared to d-TNE buffer prepared in 100% D₂O (black).

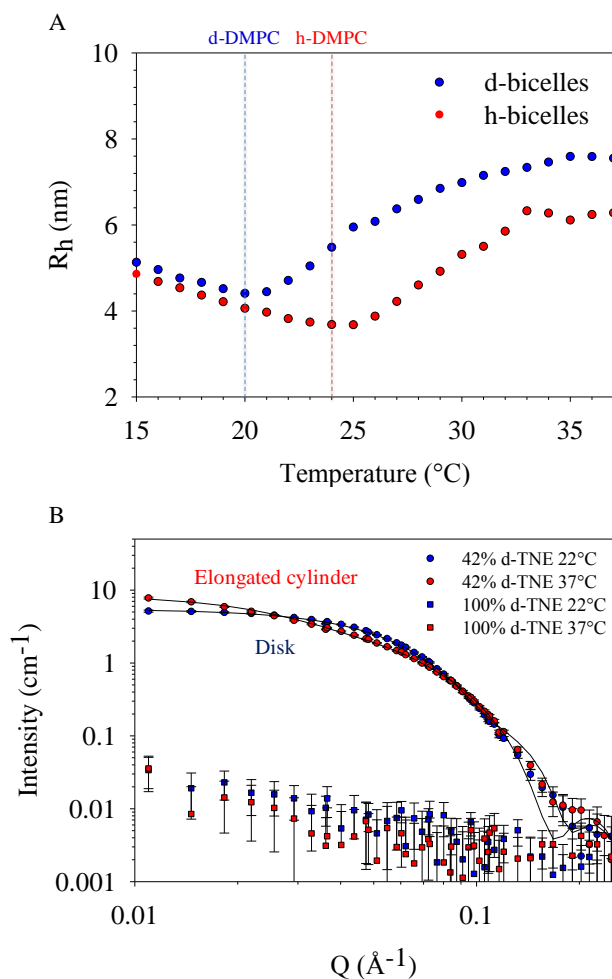


Figure 4: (A) DLS analysis of h- (red) and d-bicelles (blue) with $q_{eff} = 1.3$ bicelles at 50 mM from 15 to 37 $^{\circ}\text{C}$ in d-TNE. Dotted lines indicate the temperature transition for d- and h-DMPC (blue and red, respectively). (B) SANS intensities measured for $q_{eff} = 1.3$ d-bicelles (50 mM) at 22 (blue) and 37 $^{\circ}\text{C}$ (red) in 42% (circles) and 100% (squares) d-TNE. The data were fitted (black lines) with the “core-shell cylinder” model. At 22 $^{\circ}\text{C}$, the thickness (t) is 40 \AA and the radius (R) is 42.5 \AA , in line with the dimensions of a disk, while at 37 $^{\circ}\text{C}$, the radius is 26 \AA and the thickness is 192 \AA , corresponding to an elongated cylinder.

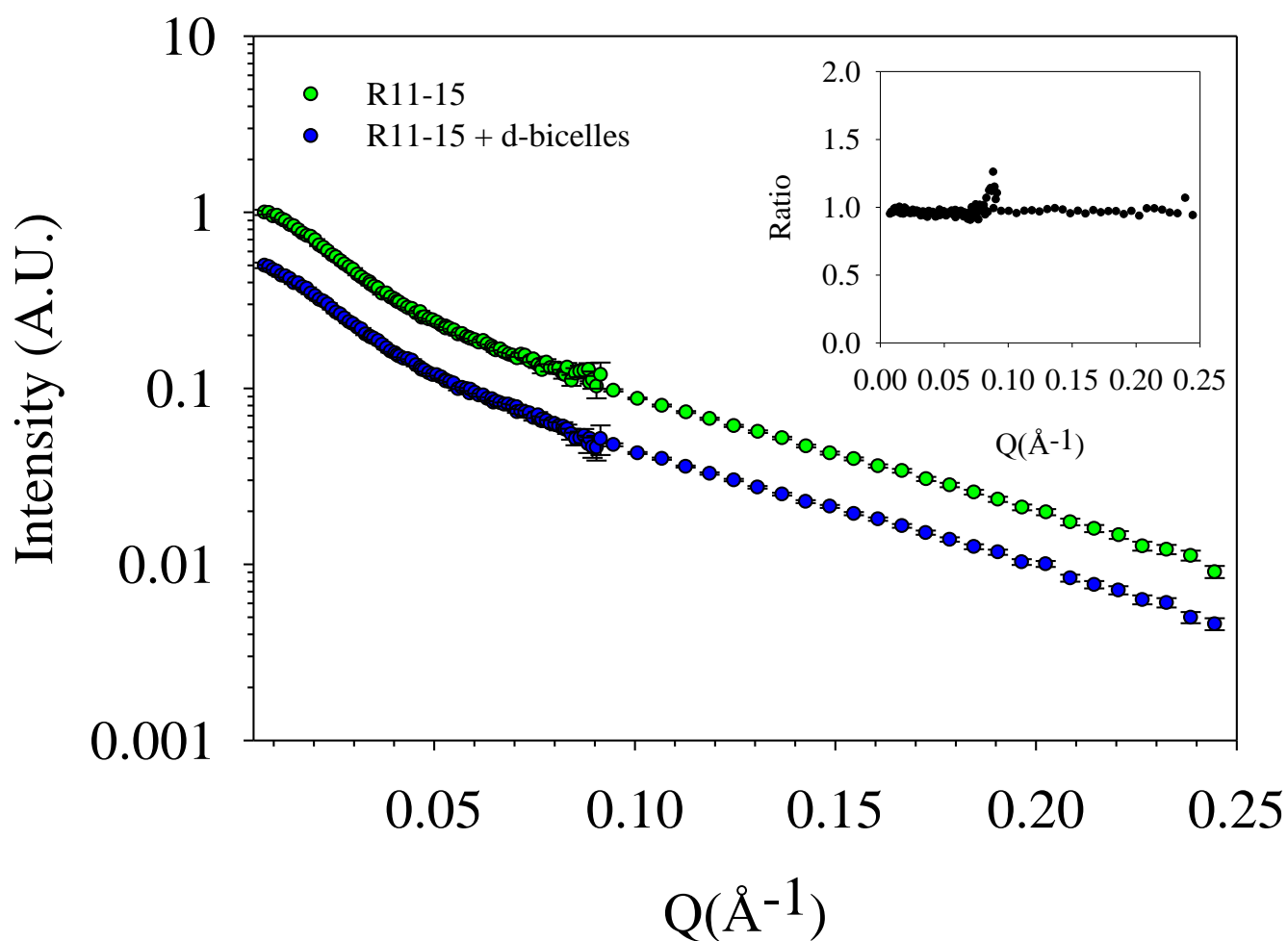


Figure 5: Normalized SANS intensities measured for DYS R11-15 at 93 μM (5.6 g/L) alone (green) or in the presence of the d-bicelles (blue) in 100% D_2O d-TNE at 18°C. Insert: the ratio of the two scattering curves. The blue curve is shifted for clarity since the two curves superimpose, as shown by the ratio of the raw data in the insert.

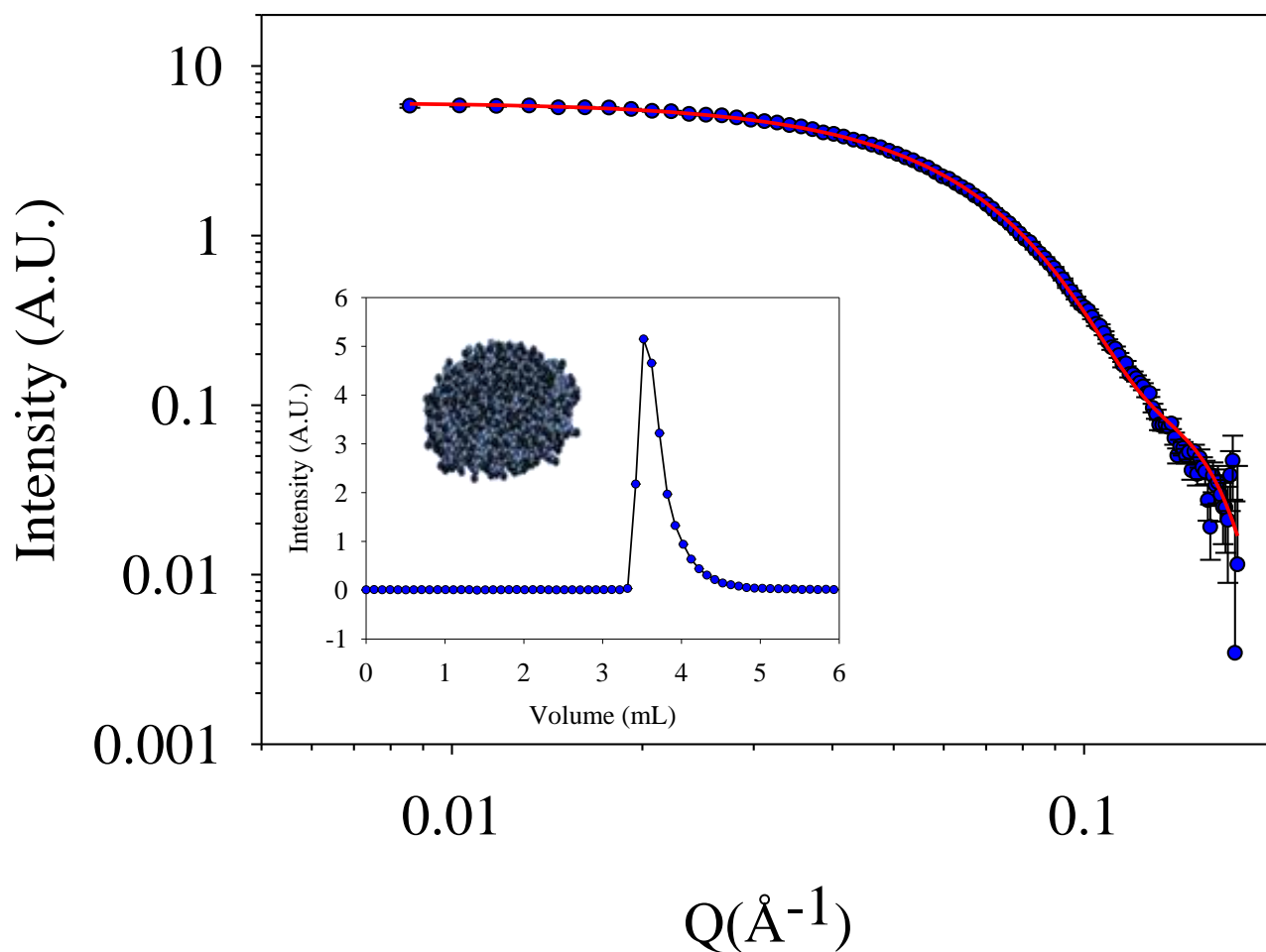


Figure 6: HPSEC-SANS intensities measured for h-bicelles fitted (red line) with the expected “core-shell cylinder” model as for the same sample in the Hellma cell environment. Insert: the associated scattergram (SANS detector) as a function of elution volume, showing one unique peak corresponding to h-bicelles.