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# Exploring the water/oil/water interface of phospholipid stabilized double emulsions by micro-focusing synchrotron SAXS

Ilaria Clemente, pab Kristian Torbensen,†c Emanuela Di Cola,d Federico Rossi, pe Sandra Ristori to\*b and Ali Abou-Hassan to\*c

Surfactant stabilized water/oil/water (w/o/w) double emulsions have received much attention in the last years motivated by their wide applications. Among double emulsions, those stabilized by phospholipids present special interest for their imitation of artificial cells, allowing the study of the effect of confining chemical reactions in biomimetic environments. Upon evaporation of the oil shell, phospholipid stabilized double emulsions can also serve as templates for giant vesicles. In this context, general assumptions have been made on the self-assembly and structural organization/arrangement of amphiphilic molecules, at the aqueous/oil liquid interface. However, to the best of our knowledge, no detailed evidence of the interfacial structuring have been reported. In this paper, w/o/w double emulsions formulated using the phospholipid 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC) and a mixture of chloroform and cyclohexane as the oil phase were produced using a microfluidic device. To obtain information on the phospholipid arrangement, the w/o/w interface was investigated by spatially resolved micro-focusing SAXS. We observed that (i) the basic units forming both the w/o and o/w interfaces were oil-swollen DMPC bilayers, arranged into a substantially disordered shell of ~45 µm thickness surrounding the internal oil phase; (ii) the evaporation process was slow, i.e. in the order of one hour at 50 °C and (iii) oil evaporation led to a shrinkage of the interfacial shell, but not to an increase of the ordering of the lipid bilayers. Interestingly, no stacked DMPC bilayers were observed during the evaporation process, as shown by the absence of Bragg's peaks in the SAXS intensity profiles.

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## Introduction

Complex biological systems such as living cells have been the subject of research for more than a century, resulting in a profound understanding of transport phenomena and metabolism. These systems often exhibit nonlinear responses to external and internal stimuli; thus, the handling of living cells has led to the search for simpler and more manageable models. For this purpose, liposomes, giant unilamellar vesicles (GUVs), water-in-oil (w/o) and water-in-oil-in water (w/o/w) double emulsions (DEs) with interfacial boundary layers

composed of self-assembled amphiphiles have proven invaluable tools to mimic cellular systems, allowing the investigation of the compartmentalization effect.1 In particular, phospholipids<sup>2,3</sup> constitute an important class of molecules to build membrane models, and have been extensively used in the study of signal transduction and transport phenomena.4 Moreover, soft self-organized structures not only serve as synthetic models to mimic biological cells,5 but also play a key role for developing carriers in drug delivery6-8 or serve as encapsulating agents in the food industry. 1-3,9 Such versatile applications have motivated researchers to develop different methodologies for controlling their architecture. Traditionally, GUVs and layered emulsions are formed by hydration of dry lipid films or directly in bulk solution, followed by extrusion. 10,11 However, these methods often fail to provide scaling-up, monodispersity and reproducibility. Techniques such as pulsed jet flow12,13 and microfluidics have been enabling the scale-up formation of uniform lipid systems.4 In particular, excellent procedures have been developed using microfluidics for templating liposomes from w/o/w DEs, allowing artificial cells to be constructed with an increasing control over size, architecture, etc. Likewise, encapsulation of active principles or chemical reactions to be carried out in confined environments has been obtained in this

<sup>&</sup>lt;sup>a</sup>Department of Biotechnology, Chemistry and Pharmacy, University of Siena, 53100, Siena, Italy

<sup>&</sup>lt;sup>b</sup>Department of Chemistry Ugo Schiff & CSGI, University of Florence, 50019, Sesto Fiorentino, Italy. E-mail: ristori@unifi.it

<sup>&</sup>lt;sup>c</sup>Sorbonne Université, CNRS, Physico-chimie des Electrolytes et Nanosystèmes InterfaciauX, F-75005 Paris, France. E-mail: ali-abou\_hassan@sorbonne-universite.fr <sup>d</sup>Department of Medical Biotechnology and Translational Medicine (Biometra), Università degli Studi di Milano, 20090, Segrate, MI, Italy

<sup>&</sup>lt;sup>e</sup>Department of Earth, Environmental and Physical Sciences, DEEP Sciences, University of Siena, Pian dei Mantellini 44, 53100 Siena, Italy

<sup>†</sup> Current address: Laboratoire d'Electrochimie Moléculaire, Université Paris Diderot, France.

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manner.14-16 As such, the w/o/w DE drops constitute both a template for liposome production, and can serve as a fairly good model for studying cell-cell communication.<sup>17,18</sup> It was proposed that by controlling the thickness of the oil film in the double emulsion drops, the final lamellar properties of vesicles can be tuned, leading to the formation of uni- or multilamellar shells or shells comprised of aggregated material. 18-21 From a molecular point of view, though it is established that in dispersed systems amphiphiles are mainly localized at the interface between polar and nonpolar domains, the detailed representation of their arrangement is a difficult task, especially when the interfacial region extends over lengths that are hundreds of times larger than the molecular size.5 Highresolution techniques, such as small angle scattering, are valuable tools for probing the organization of amphiphiles at the o/w interface of emulsions<sup>6-9</sup> and can be used to unravel the interfacial structure.

In this context, we have recently undertaken a systematic study of the effect of compartmentalization on chemical communication among diffusively coupled chemical oscillators.22,23 In particular, by taking advantage of microfluidic techniques, and in a multi-scale interfacial approach, the effect of the compartment organization (GUVs, w/o and w/o/w emulsions) on the global network dynamics (in-phase and anti-phase oscillations, pulse transmission, etc.) has been assessed. 22,24-26 In this perspective, we herein studied w/o/w DEs as microreactors formulated with a known chemical composition which was investigated previously in the chemical communication context24 in order to get in-depth information on the interfacial lipid structures including during solvent evaporation, as this fundamentally impacts the chemical communication. 17,27,28 DEs with two different oil thicknesses were first generated using a microfluidic device. Next, the w/o/w interface was investigated during the oil evaporation process with micro-focusing SAXS in order to reach structural information on the phospholipid arrangement during the oil evaporation process.

## Experimental

#### Materials and methods

Emulsions preparation. All chemicals were purchased from Sigma Aldrich and used as received. 1,2-Dimyristoyl-sn-glycero-3-phosphocholine (DMPC) phospholipids were purchased from Avanti Polar Lipids. Double emulsions (Table 1) were formed using a microfluidic device described elsewhere.29 In general, the microfluidic device consisted of coaxially aligned glass tubes of various dimensions and geometry. First, a single polyimide coated 150/50 μm (outer diameter (OD)/inner diameter (ID)) glass capillary tubes (Polymicro) was inserted into a cylindrical 1/0.5 mm (OD/ID) borosilicate tube (Drummond Scientific). The borosilicate tube was narrowed in one end for enhanced flow focusing by exposing the tube orifice to a butane flame while rotating the tube. The inner wall and the orifice of this tube were rendered hydrophobic using 2% (v/v) trichloro(1H,1H,2H,2H-perfluorooctyl)silane (Sigma Aldrich) in toluene, rinsed with toluene and dried overnight at 70 °C. These tubes are in turn inserted into a square borosilicate tube 1.25 imes

1 mm (outer/inner width, Vitrocom). A second cylindrical borosilicate tube (1/0.5 mm (OD/ID)), likewise narrowed in one end, was inserted from the opposite end of the square tube and the two cylindrical tubes were positioned with the narrowed orifices in close position. A sketch of the device used in this work is reported in Scheme 1. The inner aqueous phase, containing 2% (wt/wt) polyvinyl alcohol (PVA, average molecular weight of 18 kDa), was injected into the 150/50 µm tube with a volumetric flow rate  $Q_1 = 15 \mu L \text{ min}^{-1}$ . The middle oil phase was injected into the 1/0.5 mm cylindrical tube, engulfing the smaller tube, with a volumetric flow rates of  $Q_2 = 10 \mu L \text{ min}^{-1}$ and was composed of a solution of the phospholipid 1,2dimyristoyl-sn-glycero-3-phosphocholine (DMPC; 1% (wt/wt)) dissolved in a mixture of the volatile solvents cyclohexane/ chloroform (volume ratio 2:1). The boiling points of cyclohexane and chloroform are 81.0 °C and 61.2 °C, respectively. The microfluidic device was operated at 20 °C, thus maintaining the DMPC in the gel state. The outer phase was an aqueous solution containing 2% (wt/wt) PVA and was injected into the 1.25/1.0 mm square tube engulfing both the cylindrical tubes at a volumetric flow rates of  $Q_3 = 25 \mu L \text{ min}^{-1}$ . In this sequence, a w/o emulsion is formed first, which is subsequently embedded in the aqueous phase flowing in the square tube, forming the w/o/w emulsion. To change the thickness of the oil shell, the distance between the orifices of the two cylindrical tubes was varied. All volumetric flows were controlled using syringe pumps (KD Scientific). The PVA serves both to increase the viscosity for enhanced flow focusing and to increase the stability of double emulsions. The obtained DEs were collected in quartz capillary tubes with an inner diameter of 300 μm, and a wall thickness of 10 μm. These tubes were sealed at both ends with paraffin wax, as to reproduce the experimental setup employed for experiments on diffusively coupled chemical oscillators.22

Microfocus small angle X-ray scattering (μ-SAXS). Microfocus-SAXS was performed at the ID13 beamline of ESRF (Grenoble, France). A monochromatic beam ( $\lambda = 0.095372 \text{ nm}$ ) was focused down to a 1.5  $\times$  1.5  $\mu$ m<sup>2</sup> spot at the desired sample position using a combination of beryllium-compound refracting lenses (Be-CRL transfocator). The sample-to-detector distance was 0.941 m, corresponding to a q range of 0.2 < q <8 nm<sup>-1</sup>, where q is the scattering vector defined as  $q = (4\pi/\lambda)$  $\sin(\theta/2)$  and  $\theta$  the scattering angle. One- or two-dimensional beam raster scans with a step size of 2 µm covering the emulsion interfaces were performed in transmission geometry. At each raster step, a scattering pattern was collected by the DECTRIS EIGER 4 M single-photon-counting detector, providing frames of 2070  $\times$  2167 pixels (75  $\times$  75  $\mu$ m<sup>2</sup> pixel size). Exposure times were limited to 0.1 s per raster point to prevent beam damage.

**Optical microscopy imaging.** The structural features of the interface were studied by fabricating DEs with different starting oil thickness, which was measured from the half width of the scan profile obtained by optical microscopy at room temperature (RT). The behavior of DEs with regards to evaporation was also investigated by heating the samples at 50 °C for 30–240 min. The obtained values for the oil thickness are reported

Table 1 w/o/w double emulsions (DE) prepared using the microfluidic device and investigated using the μ-SAXS

Preparation conditions	Oil thickness (μm)	Sample name
Kept at room temperature before irradiation	44	$\mathrm{DE}_{1;44}$
Heated at 50 °C for 30 minutes before irradiation	24	$\mathrm{DE}_{1;24}$
Heated at 50 °C for 60 minutes before irradiation	20	$\mathrm{DE}_{1:20}$
Kept at room temperature before irradiation	57	$\mathrm{DE}_{2:57}$
Heated at 50 °C for 120 min before irradiation	54	$\mathrm{DE}_{2:54}$
Heated at 50 $^{\circ}\mathrm{C}$ for 240 min before irradiation	50	$ ext{DE}_{2;50}$

in Table 1. In this paper, two indexes were used for labelling samples, the first one (*i.e.* 1 or 2) indicates the series, while the second index indicates the measured oil thickness (before or after thermal treatment).

#### Results and discussion

#### The w/o/w DEs

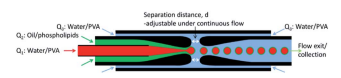
Optical images of the double emulsions (DE $_1$  and DE $_2$ ) obtained in the microfluidic device and transferred in the glass capillary are shown in Fig. 1 at RT and after heating. The middle oil phase of the DEs can be easily identified as a darker region, due to its higher refractive index with respect to the aqueous core and bulk phase. Panel 1A shows sample DE $_1$  kept at room temperature after preparation, while panel 1B and 1C show the same sample (DE $_1$ ) stored at 50 °C for 30 and 60 minutes, respectively. At close look, these optical images confirmed that a decrease in the outer diameter took place with increasing heating time.

The scan plot profiles, displayed in Fig. 1D, obtained by using the ImageJ software, clearly evidenced this trend. In addition, it showed that the width of the core size was practically constant, thus indicating that oil evaporation mainly occurs in the surrounding aqueous phase.

From the cross sections, the outer and inner diameter of the oil phase could be estimated, and the thickness and volume of the oil phase were calculated by simple geometry:

$$V_{
m oil\ shell} = rac{4}{3}\pi (R_{
m outer}^3 - R_{
m inner}^3)$$

The values obtained for heated samples showed a decrease in the oil shell thickness for DE<sub>1</sub> from 44  $\mu$ m at RT to 24  $\mu$ m and 20  $\mu$ m after heating to 50 °C for 30 and 60 minutes respectively, corresponding to a decrease in volume of the shell thickness of 20% and 29% relative to the sample stored at RT. Considering that the vapor pressure and the boiling point of chloroform are substantially lower than those of cyclohexane, for simplicity, we



Scheme 1 Asset of the microfluidic device used in this work.

assumed that the evaporated solvent was only chloroform. Thus, a 29% decrease in the oil shell volume was equivalent to a loss of 29 volume percentage chloroform. An original content of 33.3% implies that the residual chloroform in the oil phase was approximately 4% after heating the sample at 50 °C for 60 minutes. Concerning DE $_2$  (Fig. 1, panels A'–D'), the samples heated at 50 °C during 120 and 240 minutes showed a decrease in the oil thickness from 57  $\mu m$  (RT), to 54 and 50  $\mu m$  respectively, corresponding to a decrease in volume of 12% and 20%, respectively, relative to the sample stored at RT. Assuming, as previously, that all the evaporated solvent was chloroform, the corresponding residual chloroform in the oil phase was approximately 13% after heating the sample to 50 °C for 240 minutes.

#### μ-SAXS of DE

Thanks to the capability of the SAXS apparatus used in this work, the self-assembly properties of DMPC could be investigated by scanning with focused X-ray each of the three phases of the w/o/w systems, *i.e.* external water, intermediate hydrophobic layer and internal water. The one-dimensional intensity plot (I(q) vs. q) showed that some structuring occurred in the oil/DMPC, which was evidenced by a large excess scattering. The intensity decay was consistent with the SAXS pattern of bilayers,

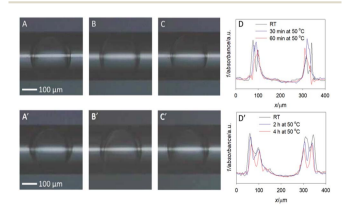


Fig. 1 Optical images of the w/o/w double emulsions studied in this paper: DE<sub>1</sub> (A) taken at the exit of the microfluidic device (RT), (B) after storage at 50 °C for 30 min and (C) 50 °C for 60 min, (D) plot profile scans obtained from the optical images (A), (B) and (C) of DE<sub>1</sub>. DE<sub>2</sub> (A') as obtained at the exit of the microfluidic device (RT), (B') after storage at 50 °C for 120 min and (C') 50 °C for 240 min, (D') plot profile scans obtained from the optical images (A'), (B') and (C') of DE<sub>2</sub>.

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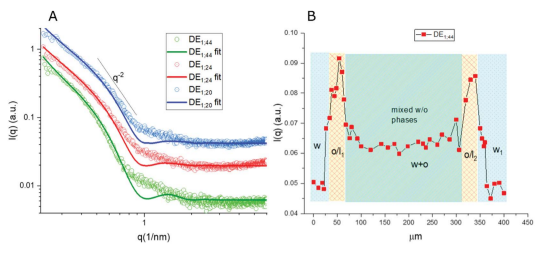


Fig. 2 (A) 1D μ-SAXS profile of the w/o/w double emulsion (DE<sub>1</sub>) interface as obtained at the exit of the microfluidic device (RT; green), after storage at 50 °C for 30 min (red) and for 60 min (blue) respectively, (B) SAXS intensity at q = 0.74 nm<sup>-1</sup> recorded along an axial line covering a whole DE<sub>1</sub> double emulsion unit. This q value was chosen to minimize the contribution of single bilayers to the overall scattering intensity.

as also reported in a previous study on w/o simple emulsions. The change of slope at 0.74 nm<sup>-1</sup> (Fig. 2, green curve) corresponded to the first minimum in the form factor of flat objects extending over large distances. However, while in the case of simple emulsions a true minimum was observed, in agreement with the presence of well-defined bilayers, here the minimum was broadened by local roughness and polydispersity in thickness (see the Table 2 for fitting results), suggesting disorder in the organization at higher level. For samples heated at 50 °C during 30 and 60 minutes the minimum position was shifted toward larger q (q = 0.85 nm<sup>-1</sup> and q = 0.88 nm<sup>-1</sup>, respectively), meaning that the bilayers thickness shrunk when heated due to solvent removal.

Noticeably, in these systems the heating process didn't induce bilayer stacking within the DMPC/oil domain but rather enhanced de-structuring. A contribution to disorder in the bilayer could also originate from partial mixing with PVA, which might have acted as a structural breaker if it diffused from the aqueous phase toward the adjacent amphiphilic layer. To correlate optical images with structural data, one-dimensional intensity diagrams along a single DE were plotted as a function of the horizontal scan (Fig. 2B), using the intensity values at a chosen wave vector ( $q = 0.74 \text{ nm}^{-1}$ ).

In agreement with optical cross sections shown in Fig. 1, the peaks of SAXS intensity were localized in the oil/lipid layers (o/l) and the lowest intensity was found in the external aqueous phases (w). Nevertheless, SAXS data indicated that the internal water/PVA phase had an intermediate intensity, caused by diffusing molecules from the interface and forming a mixed aqueous/oil domain (w + o). Similar considerations could be applied to micro-SAXS plots of DE $_2$  samples either stored at RT or heated at 50 °C for 120 and 240 min (Fig. 3).

These intensity profiles still corresponded to the form factor of loosely inter-correlated and polydisperse bilayers, as in the case of the DE $_1$  samples. However, the diagrams of the DE $_2$  didn't show any appreciable difference between RT and heated samples, meaning that samples prepared with a thicker oil layer possessed an external shell which was more resistant to evaporation. Unfortunately, the DE $_2$  samples were also prone to a marked mixing of the layers, which caused extended destructuring. Indeed, smaller differences in intensity could be observed in the scans along the capillary with respect to the DE $_1$  samples. The only visible features were the presence of excess scattering and the broad minimum at 0.9 nm $^{-1}$ , *i.e.* at lower q values than for the DE $_1$  samples, indicating that even if the oil shell was thicker, the single bilayers were less swollen. The

Table 2 Fitting parameters of SAXS 1D profiles

Sample	Single tail thickness (Å)	Head thickness (Å)	SLD head $(\mathring{A}^{-2})$	SLD tails ( $\mathring{A}^{-2}$ )	Tail polydispersity (lognormal distribution)
$DE_{1;44}$	$25.0 \pm 0.7$	$5.9\pm0.4$	$12\pm1$	$3.6\pm0.4$	$1.6\pm0.2$
$DE_{1:24}$	$25.0\pm0.5$	$6.0\pm0.6$	$11.0\pm0.5$	$4.0\pm0.3$	$1.6\pm0.1$
$DE_{1;20}$	$25.0\pm0.5$	$5.7\pm0.4$	$11.0\pm0.5$	$4.0\pm0.3$	$1.6\pm0.1$
$DE_{2:57}$	$21.5\pm0.5$	$5.0 \pm 0.3$	$11.0\pm0.2$	$4.0\pm0.5$	$1.4\pm0.3$
$DE_{2;54}$	$21.5\pm0.4$	$5.0\pm0.4$	$11.0\pm0.2$	$4.0\pm0.5$	$1.4\pm0.3$
$\mathrm{DE}_{2;50}$	$21.5\pm0.4$	$5.0\pm0.4$	$11.0\pm0.2$	$4.0\pm0.5$	$1.4\pm0.3$

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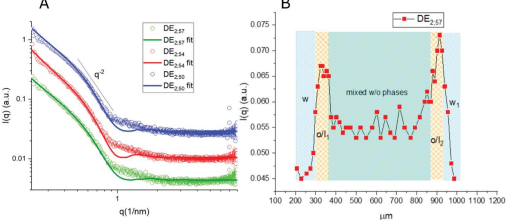


Fig. 3 (A) 1D μ-SAXS profile of the w/o/w double emulsion (DE<sub>2</sub>) interface as obtained at the exit of the microfluidic device (RT; green), after storage at 50 °C for 120 min (red) and for 240 min (blue) respectively, (B) SAXS intensity at  $q = 0.9 \text{ nm}^{-1}$  recorded along an axial line covering a whole DE2 double emulsion unit.

fitting of the SAXS diagrams of the DE1 samples (Fig. 2A) shown as solid lines superimposed to the corresponding experimental profiles confirmed the presence of polydisperse bilayers with a typical  $q^{-2}$  decay of locally flat object, and total bilayer thickness of about 60 Å. For the fitting of the DE<sub>2</sub> samples (Fig. 3A), the same model was used obtaining a total bilayer thickness of about 55 Å, confirming the presence of less swollen bilayers. The results of the fittings for DE<sub>1</sub> and DE<sub>2</sub> are summarized in Table 2.

In both DE<sub>1</sub> and DE<sub>2</sub> systems there was a localized region of scattering intensity that could not be fitted using the polydisperse lamellar model (SASView package). Therefore, a certain level of correlation was assumed. To account for this hypothesis, a Gaussian peak (Fig. 4, dashed line) centered at 0.9 nm<sup>-1</sup> could be considered in addition to the form factor of uncorrelated bilayers, as the signature of a loose organization occurring at intermediate distances. It could also be evidenced that some

residual intensity was present in the internal water/PVA phase  $(w_1 \text{ and } w_2)$ . As it is well-known, SAXS is extremely sensitive to the presence of molecules, or groups with high electron density and can be used to monitor the variations of chemical composition in the environment probed by the beam. 30,31 Indeed, due to partial mixing, molecules able to provide contrast and produce scattering could penetrate the boundary layer and, as a consequence, such region appeared less dark than expected for a pure isotropic phase with low SAXS intensity. This was in perfect agreement with the increased background intensity

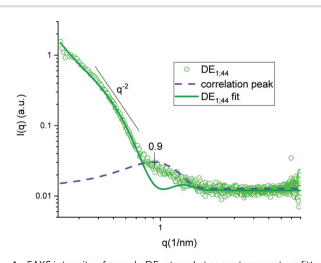


Fig. 4 SAXS intensity of sample DE<sub>1</sub> stored at room temperature fitted by a polydisperse bilayer model (green), with Gaussian peak (violet) superposition to account for inter-bilayer correlation.

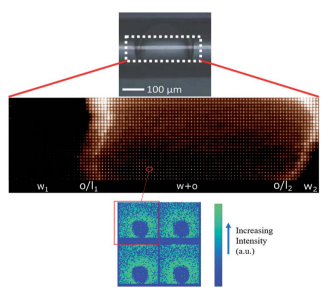


Fig. 5 Upper panel: two-dimensional intensity mesh of a typical DE<sub>1</sub> sample stored at room temperature showing the water/PVA mixture which constitutes the internal (w<sub>1</sub>) and external (w<sub>2</sub>) phases, both interfaces (oil/l), and the oil shell named w + o because of partial mixing (see text for further details). In this image the signals of glass capillary and water have been subtracted. Lower panel: typical 2D patterns of the detector in blue-to-green intensity scale expressed in arbitrary units

observed in the profile shown in Fig. 2 and 3 and discussed above. Such insights from 1D data were supported by the two-dimensional diagrams (Fig. 5), where pixels given from the 2D scattering collected on the detector are reported. The observed patterns evidenced that the systems have no orientational anisotropy, which could possibly be induced by the glass walls. Moreover, the meshes obtained by subsequent recording of 2D patterns at 0.1 s per frame interval showed the presence of a film at the oil/DMPC interface, appearing as the brightest region (o/l in Fig. 5), in contrast with the internal and external water/PVA phases ( $w_1$  and  $w_2$ ), which appear darker due to lower scattering intensity in these regions.

## Conclusions

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Liquid-liquid confined microdroplets stabilized by lipids are complex systems with wide scientific and applicative potential. Even though many techniques have been developed to control their formation, there is still some uncertainty about the fine molecular arrangement of self-assembling amphiphiles at the interface during such process. In this work the compartmentalization of organic and aqueous domains into soft matter microstructures was achieved by fabricating w/o/w double emulsions with precisely controlled geometry and size. The boundary layers were formed by phospholipids (DMPC), which self-assembled from a solution of organic volatile solvents, i.e. a mixture of cyclohexane/chloroform 2:1 v/v. An ad hoc engineered microfluidic platform coupled to a microfocus-SAXS apparatus allowed to obtain information on the internal structure of each compartment and the extent of molecular transmigration between adjacent micro-phases. In particular, solvent evaporation and its effects on the lipid arrangement were investigated by spatially resolved SAXS. This analysis revealed that the basic units of the phospholipid domains at the w/o interface were oil-swollen DMPC bilayers, which were arranged into a substantially disordered shell of  ${\sim}45~\mu m$ thickness surrounding the internal oil phase. Upon heating at 50 °C for 30 and 60 minutes, a large fraction of the initial solvent volume was removed, and the thickness of the lipid boundary was reduced to 20-25 µm. Concurrently, the single bilayers shrunk by about 20% of their initial thickness and the interbilayer correlation slightly increased. However, the heating process did not induce stacking of DMPC bilayers, evidenced by the absence of Bragg's peaks in the SAXS intensity profiles of the heated samples. A further interesting finding was that the thickness of the boundary layer was able to modulate oil evaporation, since the solvent removal was more pronounced in samples with lower interfacial thickness. We think that, in the chemical communication context, the arrangement of the phospholipids at the liquid/liquid interfaces and in the oil shell, its evolution with time upon solvent evaporation as well as the mixing observed in the water phase (as revealed by microfocusing SAXS) should be taken into account as they may play an relevant role in the partition and transfer of chemical species between double emulsions compartments, thus influencing the dynamics of the system.

## Conflicts of interest

There are no conflicts to declare.

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