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Mechanical properties of membranes composed of gel-phase or fluid-phase phospholipids probed on liposomes by atomic force spectroscopy

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3 1 **Mechanical properties of membranes composed of gel-phase or fluid-phase**
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5 2 **phospholipids probed on liposomes by atomic force spectroscopy**
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38
39 14 Key words: small unilamellar vesicle; Phase state; Bending modulus; Young's modulus;
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41 15 Atomic force microscopy
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43

44 16 **Abstract**
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46
47 17 In many liposome applications, the nanomechanical properties of the membrane envelope are
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49 18 essential to ensure e.g. physical stability, protection or penetration into tissues. Of all factors,
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51 19 the lipid composition and its phase behavior are susceptible to tune the mechanical properties
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53 20 of membranes. To investigate this, small unilamellar vesicles (SUV; diameter <200 nm),
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3 21 referred to as liposomes, were produced using either the unsaturated 1,2-dioleoyl-sn-glycero-
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5 22 3-phosphocholine (DOPC) or the saturated 1,2-dipalmitoyl-sn-glycero-3-phosphocholine
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8 23 (DPPC) in aqueous buffer at pH 6.7. The respective melting temperatures of these
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11 24 phospholipids were -20°C and 41°C. X-ray diffraction analysis confirmed that at 20°C DOPC
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14 25 was in the fluid phase and DPPC was in the gel phase. After adsorption of the liposomes onto
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17 26 flat silicon substrates, atomic force microscopy (AFM) was used to image and probe the
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20 27 mechanical properties of the liposome membrane. The resulting force-distance curves were
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22 28 treated using an analytical model based on the shell theory to yield the Young's modulus (E)
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25 29 and the bending rigidity (k_C) of the curved membranes. The mechanical investigation showed
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28 30 that DPPC membranes were much stiffer ($E = 116 \pm 45$ MPa) than those of DOPC ($E = 13 \pm$
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31 31 9 MPa) at 20°C. The study demonstrates that the employed methodology allows
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34 32 discrimination of the respective properties of gel- or fluid-phase membranes when in the
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37 33 shape of liposomes. It opens perspectives to map the mechanical properties of liposomes
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40 34 containing both fluid and gel phases or of biological systems.

35 I. INTRODUCTION

36 In the recent years, small unilamellar vesicles of phospholipids ($< \mu\text{m}$ diameter) have gained
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39 37 increasing interest in various liposome technologies, e.g. as drug delivery systems in
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42 38 pharmacy or as protective cargo capsules for cosmetics, nutraceuticals or for food design^{1,2}.
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45 39 They are also interesting models to investigate and possibly control *in vivo* biological
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48 40 signaling mechanisms involving extracellular vesicles such as the so-called exosomes^{3,4}. The
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3 41 composition of the phospholipids, i.e. the acyl chains' saturation/unsaturation and length,
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5 42 impacts their phase state in the liposomes at ambient temperatures⁵. This, in turn, is likely to
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8 43 affect the mechanical properties of the liposomes, which are of paramount importance for
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11 44 these applications. Mechanical properties direct the stability, size, shape and fusion of the
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14 45 liposomes⁶⁻⁸ as well as the membrane fluidity/rigidity or permeability and hence their loading
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17 46 capacity or their ability to penetrate tissues⁹⁻¹³. For example, the permeability of various
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20 47 saturated polar lipid membranes to glucose strongly increases as the lipid undergoes gel-to-
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23 48 liquid disordered (l_d) phase transition¹⁴. As another example, milk sphingomyelin exhibited
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26 49 maximum permeability at temperatures where gel-to- l_d phase transition occurs¹³. This has
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29 50 important consequences on the formulation and design of liposomes, as the desirable
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32 51 mechanical properties will differ depending on the application and temperature, e.g. storage at
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35 52 ambient temperature, transdermal delivery of drugs, etc. Experimental techniques to measure
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38 53 the mechanical properties of liposomes, such as the pipette aspiration technique, osmotic or
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41 54 mechanical compression, shear-induced or optical tweezers deformation⁷ are often designed
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44 55 for large objects of the μm length scale. For liposomes at the nanoscale, such as small
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47 56 unilamellar vesicles, indentation measurement using atomic force microscopy (AFM) has
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50 57 proven a valuable and sensitive approach¹⁵⁻²⁰. The liposomes, adsorbed onto a flat substrate,
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53 58 are indented by the AFM with low penetration distances (i.e. the order of magnitude of the
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56 59 membrane thickness) and at as low force values as down to the pN. The major advantage of
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59 60 AFM over a nano-indenter is also its imaging capacity, which allows to locate liposomes and
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3 61 to indent them centrally, even though not perfectly normally¹⁶. In a pioneering study, Laney et
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5 62 al.¹⁷ extracted synaptic liposomes (~110 nm diameter) from the electric organ of the electric
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8 63 ray *Torpedo californica*, immobilized them by adsorption onto mica and measured elastic
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11 64 moduli values in the range 0.2 – 1.3 MPa. Liang et al.^{19,21,22} followed the same approach on
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14 65 small (40-160 nm) liposomes of egg phosphatidylcholine (mainly
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17 66 stearyloleoylphosphatidylcholine, SOPC) and found that the elastic modulus increased with
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20 67 the addition of up to 50 mol % cholesterol from ~2 to 13 MPa. However, these authors used
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23 68 an adaptation of the Hertz model for their calculations that assumed the liposomes to be
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26 69 homogeneous filled spheres. By implementing the shell deformation theory, Delorme and
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28 70 Fery¹⁶ obtained higher elasticity values of ~110 MPa by indenting DPPC
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31 71 (dipalmitoylphosphatidylcholine) and proposed that the Hertz model underestimated the
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34 72 mechanical properties of the liposomes. However, due to its mono-unsaturation, egg PC has a
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37 73 melting temperature (T_m) of -15°C and is therefore in the l_d or liquid crystalline ($L\alpha$) phase at
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40 74 20°C, while the fully saturated DPPC is in the gel or solid-ordered (s_o) phase (T_m =
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42 75 41.7°C)^{22,23}. Therefore, it is yet to assess whether the AFM indentation measurement
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45 76 combined with the shell theory interpretation is sensitive enough to discriminate liposomes
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48 77 with presumably different mechanical properties of their membranes, e.g. by comparing
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51 78 saturated and unsaturated phospholipids in different physical phases at 20°C.
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54 79 When the membranes are spread as two-dimensional supported lipid bilayers (SLBs),
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56 80 measurement of the rupture force of membranes using AFM indentation has proven to fully
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3 81 resolve mechanical differences between phospholipids with various chain lengths, saturation
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5 82 degrees or head groups²⁴. However, calculation of the SLBs' effective elasticity using
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8 83 indentation of the membrane and the Hertz model requires careful and narrow experimental
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11 84 conditions not to be affected by the support²⁵. Furthermore, direct measurement on small
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14 85 liposomes would encompass possible curvature effects. Indeed, their high membrane
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17 86 curvature may affect their mechanical properties, as lateral intermolecular distances and
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20 87 forces vary across the bilayer's thickness^{26,27}. However, indentation measurements on 3D
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23 88 vesicles may be either impossible²⁸, require high deformation of the objects¹⁹ or somewhat
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26 89 depend on the composition of the internal medium, if different from the surrounding
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29 90 medium²⁹. For these reasons, there is interest to assess indentation measurement of the
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32 91 mechanical properties of membranes directly on volumetric objects such as liposomes.
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34 92 In the present study, the respective elasticities and bending rigidities of membranes of either
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37 93 the unsaturated phospholipid dioleoylphosphatidylcholine (DOPC) or the saturated
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40 94 phospholipid dipalmitoylphosphatidylcholine (DPPC) as shaped in the form of liposomes,
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43 95 were measured at 20°C using AFM indentation and the shell theory. The results were
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46 96 discussed in light of the respective phase states of the two phospholipids at 20°C and showed
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49 97 that AFM indentation is a sensitive method to assess the mechanical properties of 3D
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52 98 membrane objects at the nanoscale.
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100 II. EXPERIMENTAL METHODS

101 **2.1. Materials**

102 Pure phospholipids 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC; 18:1; >99%) and 1,2-
103 dipalmitoyl-sn-glycero-3-phospholcholine (DPPC; 16:0; >99%) were purchased from Avanti
104 Polar Lipids (Alabaster, AL). PIPES (1,4-piperazinediethanesulfonic acid) buffer was
105 prepared as: PIPES 10 mM (purity \geq 99%; Sigma-Aldrich, Milwaukee, WI, USA), NaCl 50
106 mM (Sigma), and CaCl₂ 10 mM (Sigma) were dissolved in Milli-Q water and adjusted to pH
107 6.7 using NaOH 5 M.

108 **2.2. Preparation of liposomes**

109 Samples were prepared by dissolving appropriate quantity of the lipid powder of DOPC or
110 DPPC in glass vials with chloroform/methanol (4:1 v/v). The organic solvent was then
111 evaporated at 40 °C under a stream of dry nitrogen. The dried lipid films were hydrated with
112 PIPES-NaCl-CaCl₂ buffer at 70°C to reach a final concentration of 0.1 wt. % lipids then
113 thoroughly vortexed. Small unilamellar vesicles (SUV) were produced at 65°C by sonication
114 using a Q700 equipment (Q-sonica, Newtown, CT, USA) and a microtip operating at 50%
115 amplitude (~400 W) for 30 min. After sonication, the SUV suspension was left to cool and
116 equilibrate at room temperature (20°C). The SUV produced according to this protocol will be
117 designated as “liposomes” throughout this report.

118 **2.3. Dynamic light scattering (DLS)**

119 The size distribution and the average hydrodynamic diameter (D_h) of the vesicles were
120 measured in PIPES-NaCl-CaCl₂ buffer at 20°C by dynamic light scattering (DLS) on a

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3 121 Zetasizer Nano ZS (Malvern Instruments, Worcestershire, United Kingdom). Measurements
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5 122 were carried out at a scattering angle of 173° and a wavelength of 633 nm. The average D_h (\pm
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8 123 5 nm) was calculated from the intensity distribution using conversion into an autocorrelation
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11 124 function which is then analyzed with the Stokes-Einstein relation, assuming that particles had
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14 125 a spherical shape. The viscosity of the solution was 1.003 mPa.s at 20°C and the refractive
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17 126 index of the solvent was 1.33.

127 **2.4. Differential scanning calorimetry (DSC)**

128 The thermotropic properties of DOPC or DPPC were measured on multilamellar vesicles
129 using a differential scanning calorimetry (DSC) Q1000 apparatus (TA Instruments,
130 Newcastle, USA). Multilamellar vesicles (MLV) were produced by rehydration of the lipid
131 films with PIPES-NaCl-CaCl₂ buffer at 65°C to reach a final concentration of 20 wt. % lipids,
132 then thorough vortex mixing. MLV are preferred over unilamellar vesicles in order to
133 accommodate the high bilayer concentration. They also allow higher resolution of the DSC
134 thermograms thanks to higher cooperativity of the molecules³⁰. The samples were introduced
135 in 20 μL aluminum pans that were then hermetically sealed. An empty pan was used as a
136 reference. The samples were heated at $2^\circ\text{C}\cdot\text{min}^{-1}$ from -40°C to 70°C . The calibration of the
137 calorimeter was performed with indium standard (melting point = 156.66°C , ΔH melting =
138 $28.41 \text{ J}\cdot\text{g}^{-1}$). The thermal measurements were performed in triplicate. Standard parameters
139 were calculated by the TA software (Universal Analysis 2000, v 4.1 D).

140 **2.5. Temperature-controlled X-ray diffraction (XRD)**

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3 141 X-ray scattering experiments were performed on the home-made Guinier beamline at IPR³¹ .
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6 142 A two-dimensional Pilatus detector with sample to detector distance of 232 mm allowed the
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9 143 recording of XRD patterns in the range 0.013 \AA^{-1} to 1.742 \AA^{-1} , thus covering both the small
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12 144 and wide-angles regions of interest to characterize the lamellar structures and to identify the
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14 145 packing of the acyl chains, respectively. Diffraction patterns displayed series of concentric
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17 146 rings as a function of the radial scattering vector $q = 4 \pi \sin\theta / \lambda$, where 2θ is the scattering
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19
20 147 angle and $\lambda = 1.541 \text{ \AA}$ is the wavelength of the incident beam. The channel to scattering vector
21
22
23 148 q calibration of the detector was carried out with silver behenate³². Small volumes (around 10
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26 149 μL) of samples containing DOPC or DPPC vesicles were loaded in thin quartz capillaries of
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29 150 1.5 mm diameter (GLAS W. Muller, Berlin, Germany) and inserted in the set-up at a
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32 151 controlled temperature.

33 152 **2.6. Transmission electron microscopy (TEM)**

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36 153 The observation of DOPC and DPPC Liposomes by cryo-TEM was realized as described in
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39 154 previous work³³. The samples were prepared using a cryoplunge cryo-fixation device (Gatan,
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42 155 Pleasanton, CA, USA) in which a drop of the aqueous suspension was deposited on to glow-
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45 156 discharged holey-type carbon-coated grids (Ted Pella Inc., Redding, CA, USA). The TEM
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48 157 grid was then prepared by blotting the drop containing the specimen to a thin liquid layer
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51 158 remained across the holes in the support carbon film. The liquid film was vitrified by rapidly
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54 159 plunging the grid into liquid ethane cooled by liquid nitrogen. The vitrified suspension of
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57 160 liposomes was mounted in a Gatan 910 specimen holder that was inserted in the microscope
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3 161 using a CT-3500-cryotransfer system (Gatan, USA) and cooled with liquid nitrogen. TEM
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5 162 images were then obtained from liposomes suspension preserved in vitreous ice and
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8 163 suspended across a hole in the supporting carbon substrate. The samples were observed under
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11 164 low dose conditions ($< 10 \text{ e}^-/\text{Å}^2$), at -178 °C , using a JEM 1230 'Cryo' microscope (Jeol,
12
13
14 165 Japan) operated at 80 keV and equipped with a LaB6 filament. All the micrographs were
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16
17 166 recorded on a Gatan 1,35 K \times 1,04 K \times 12 bit ES500W CCD camera.

18 19 20 167 **2.7. Atomic force microscopy (AFM)**

21 22 168 **2.7.1. Indentation of liposomes**

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25 169 Simple open liquid sample cells were fabricated by gluing small ($\sim 0.5 \times 1 \text{ cm}^2$) pieces of
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28 170 silicon substrate (molecular orientation 100) onto diagnostic glass slides (Thermo Scientific,
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31 171 Waltham, MA, USA). After thorough cleaning of the cell with ethanol, water and UV/O₃, the
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34 172 liposome suspension equilibrated at 20°C was deposited onto the clean silicon surface then
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37 173 left to incubate at 20°C for 30 min. The droplet was then gently exchanged with PIPES-NaCl-
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40 174 CaCl₂ buffer at 20°C to remove the un-adsorbed liposomes. The sample was then imaged in
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43 175 contact mode using an MFP-3D Bio AFM (Asylum Research, Santa Barbara, CA, USA), with
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46 176 a typical scan rate of 1 Hz for $20 \times 20 \text{ }\mu\text{m}^2$ and 256×256 pixels images, silicon MLCT
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48
49 177 probes (nominal spring constant $k \sim 0.03 \text{ N.m}^{-1}$ – Bruker Nano Surfaces, Santa Barbara, CA,
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52 178 USA) calibrated extemporaneously using the thermal noise method, and loading forces
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55 179 typically below 1 nN. Upon adsorption onto the flat surface, the spherical liposomes deform
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58 180 into spherical cap geometry. AFM imaging of large area (typical size) allows localization of
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3 181 the adsorbed liposomes, then closer images (typically $2 \times 2 \mu\text{m}^2$ or less) were recorded and
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6 182 sections were drawn across the images in order to measure the individual liposome's height
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9 183 (H) and base width (W). The AFM probe was then positioned above the center of each
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11 184 liposome and individual force curves ($n > 60$) were recorded with a set point of 200 pN, a
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14 185 distance of 100 nm and a Z-piezo speed of $2 \mu\text{m}\cdot\text{s}^{-1}$. In these conditions, indentation of the
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17 186 AFM tip into the liposome did not exceed ~ 5 nm, thereby allowing measurement of the
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19
20 187 mechanical properties in the elastic regime of the membrane. The approach curves were then
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22 188 treated using the shell theory as described in the result section.
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25 189 For data visualization and analysis, Gwyddion 2.47 software was also used, as a means to
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28 190 deduce the local radius of curvature (R_c) at the top of individual adsorbed liposomes over a
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31 191 distance of ~ 130 nm both sides of the apex (130 nm being the estimated radius of the
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34 192 membrane area affected by indentation by a 20-nm radius MLCT probe). The R_c values of
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37 193 individual liposomes were input in the calculation of the Young modulus (see below).
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39 194 Measurement of the local R_c values at the apex of individual liposomes, instead of for the
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42 195 complete objects, avoided errors due to imperfect (e.g. flattened) spherical cap geometry of
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45 196 the liposome upon adsorption, and/or errors due to convolution by the AFM tip. The latter
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47
48 197 may indeed overestimate lateral dimensions of the liposomes by as much as the tip diameter,
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50
51 198 but can be avoided as long as near-normal contact is maintained between the tip and the
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53 199 liposomes, which is the case when measuring R_c at their apex.

200 **2.7.2. Measurement of the bilayer thickness**

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3 201 Immediately after sonication at 65°C, 10 µg of the hot lipid suspension were deposited onto
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6 202 freshly cleaved mica in an Asylum Research open liquid cell, then incubated at 65°C for 60
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9 203 min. Slow cooling of the samples was performed using a programmed incubator as in Murthy
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11 204 et al.³⁴ to yield supported lipid bilayers (SLB). Once equilibrated at 20°C, the bilayers were
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14 205 extensively rinsed and exchanged with PIPES-NaCl-CaCl₂ buffer. AFM imaging of the
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17 206 bilayers was then performed in the same buffer and in contact mode using MSNL probe
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20 207 (nominal spring constant $k \sim 0.03 \text{ N.m}^{-1}$ – Bruker Nano Surfaces, Santa Barbara, CA, USA)
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22
23 208 with the same imaging parameters already cited. The probes were calibrated
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25
26 209 extemporaneously using the thermal noise method. Force spectroscopy curves were then
27
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29 210 acquired at 20°C by using force-volume imaging of the bilayers (typically $10 \times 10 \text{ µm}^2$ or
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31 211 less) with a typical set point of 20 nN and a piezo speed of 2 µm.s^{-1} .

33 212 **2.8. Statistical analysis**

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36 213 The results are presented as mean value \pm standard deviation. Analysis of variance was
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39 214 performed using the General Linear Model procedure of Statgraphics Plus version 5
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41
42 215 (Statistical Graphics Corp., Englewood Cliffs, NJ). Differences were significant for $p < 0.05$.

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47 217 **III. RESULTS AND DISCUSSION**

50 218 **3.1. Phase state of DOPC and DPPC at 20°C**

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53 219 The thermal phase behavior of unsaturated (18:1) DOPC and saturated (16:0) DPPC
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56 220 phospholipids (chemical structures shown in Fig1.A) was examined using DSC and the
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3 221 results were correlated with the structural analysis performed by XRD. Fig1.B shows the
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6 222 respective DSC heating thermograms of both lipids in the same conditions. For DOPC, the
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9 223 thermogram shows a very low temperature of L_{β} to L_{α} (gel to fluid) phase transition at $T_m = -$
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11 224 20°C , in good agreement with previous reports ^{11,35,36}. Meanwhile, the heating of DPPC
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13
14 225 revealed two endotherms, characteristic of the L_{β} (gel) to P_{β} (ripple) transition at
15
16
17 226 $T_m = 37.09^{\circ}\text{C}$ and of the P_{β} (ripple) to L_{α} (fluid) transition at $T_m = 41.01^{\circ}\text{C}$, also in good
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19
20 227 agreement with previous reports ^{34,35,37}. XRD experiments allowed identification of the lipid
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22 228 phases at 20°C (Fig. 1C). For this, the MLV are interesting not only to investigate the lateral
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25 229 packing of the acyl chains (at large q), but also to confirm the lamellar organization of the
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28 230 phospholipid (at small q). For DOPC, the absence of diffraction peak at wide angles and a
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30
31 231 lamellar organization characterized at small angles, confirmed the L_{α} phase of DOPC at 20°C .
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33 232 DPPC multilamellar vesicles exhibited a single broad peak at $q \sim 1.5 \text{ \AA}^{-1}$ corresponding to the
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35
36 233 formation of ordered phase packed in pseudo-hexagonal lattice and a lamellar organization at
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39 234 small angles, showing the formation of a L_{β} organization of DPPC at 20°C . The tilt of the
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41
42 235 acyl chains may have been induced by the multilamellar organization of the DPPC molecules
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45 236 in the vesicles. Since AFM experiments on DPPC molecules were performed in SUV, the
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47
48 237 ordered phase state of DPPC molecules in SUV at 20°C was also proved by XRD
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51 238 experiments performed on SUV of DPPC. As previously reported in literature³⁸, the high
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54 239 curvature of expected SUV of DPPC do not allow the recording of a peak at wide angles.
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56 240 Nevertheless, the Small Angles X-ray Scattering could be performed upon heating of the
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3 241 SUV. Both high and low temperature spectra can be fitted as an individual membrane using
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5 242 equation (1):
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12 244
$$I(q) = \frac{1}{q^2} \left[\frac{\sin(q.e_{HG}/2)}{q.e_{HG}} . e^{-(q.r)^2} + \frac{\sin(q.e_{CH2}/2)}{q.e_{CH2}} \right]^2 \quad (1)$$

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16 245 with e_{HG} , the headgroup thickness, e_{CH2} , the aliphatic chain thickness and r the rugosity of the
17
18 246 headgroup/water interface.
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22 247 The full analysis demonstrates a contraction of 2Å of both e_{HG} and e_{CH2} at the gel-fluid
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24 248 transition at 40-44°C (Fig. 1D). This showed that the DPPC molecules in SUV are sensitive to
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26 249 temperature and consequently in the gel phase at room temperature. In conclusion, DSC and
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28 250 XRD confirmed that DOPC (unsaturated; $T_m = -20^\circ\text{C}$) and DPPC (saturated; $T_m = 41^\circ\text{C}$)
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30 251 were respectively present in fluid and gel phases at 20°C.
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34 252 **3.2. Morphology of the liposomes at 20°C**

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38 253 Electron microscopy, dynamic light scattering (DLS) and AFM imaging were used to
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40 254 characterize the morphology (size and shape) of the liposomes obtained after sonication (Fig
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42 255 2). Cryo-TEM images showed that DOPC and DPPC vesicles were essentially unilamellar
43
44 256 and spherically shaped. However, while DOPC liposomes consistently exhibited circular
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46 257 cross-sections (Fig 2A), DPPC could show both rounded and/or somewhat faceted
47
48 258 membranes (Fig 2D). The presence of angle facets was attributed to the physical gel state of
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50 259 DPPC at 20°C³⁹. Cryo-TEM images revealed variable liposome diameters (from tens to
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1
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3 260 hundreds of nm) which corresponded to the size distribution measured by DLS (Fig 2B, E).

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5 261 For DOPC, the size distribution was monomodal with a mean diameter D_h at 120 ± 69 nm

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7
8 262 (Fig 2.B). Whereas, DPPC liposomes exhibited a shouldered size distribution with a mean

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10
11 263 diameter D_h of 132 ± 73 nm for the whole distribution (Fig 2.E; Table1).

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13
14 264 In the next step, the liposomes were immobilized by adsorption onto silicon to be first

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17 265 imaged then indented using an AFM probe. Images recorded in contact mode showed that

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20 266 DOPC and DPPC liposomes were perfectly stable on the flat substrate. However, it was

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22
23 267 not possible to prevent non-destructive deformation due to adsorption, as previously

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25 268 reported^{19,40}. The AFM images were also used to measure the height H and basal width W

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28 269 of adsorbed individual liposomes in order to prove that objects observed in AFM images

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31 270 corresponded truly to liposomes. To do this, the adsorbed liposomes were considered to

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34 271 adopt a spherical cap geometry and their volume was calculated as:

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$$V = \frac{\pi H}{6} \left(\frac{3}{4} W^2 + H^2 \right) \quad (2)$$

272 From (Eq.2), the mean diameter of a sphere of equivalent volume was inferred and compared

273 with DLS data (Table 1; Figures 2B and E). The mean diameter values were of similar orders

274 of magnitude in both methods; and whatever the method used, DOPC liposomes always

275 exhibited smaller diameters than DPPC ones. This confirmed that the adsorbed objects visible

276 on the AFM images truly were liposomes. However, the mean diameter results measured by

277 AFM were higher than DLS probably because larger liposomes were preferably chosen out of

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3 278 large-area images (such as shown on Figure 2) to perform close-up views and indentation
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5 279 measurements (Table1).
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8 280 **3.3. Mechanical properties of liposomes as a function of phase state at 20°C**

9 281 **3.3.1. Bilayer thickness**

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14 282 The average bilayer thickness d was measured in the conditions used in this work:
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17 283 temperature 20°C, pH=6.7, 50 mM NaCl and 10 mM CaCl₂. This measurement was
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20 284 performed using atomic force spectroscopy on SLB of DOPC or DPPC spread onto freshly
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23 285 cleaved mica. Average values of d were obtained from measurements performed on various
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26 286 regions of different samples (n=50). In a typical force spectroscopy experiment, the AFM tip
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29 287 approaches the surface until a mechanical contact with the SLB is established. Then, the
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32 288 bilayer is elastically deformed by the AFM probe until the tip ruptures (breaks through) the
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35 289 membrane, thereby coming into contact with the substrate. The indentation *vs* force curves
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38 290 exhibited breakthrough events^{23,34,41} where the jump-through distance (noted d) was
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41 291 assimilated as the bilayer's thickness³⁴. The limitations of this measurement were discussed
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44 292 elsewhere⁴² and efforts were taken to regard bilayer compression. Fig.3A and C show three
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47 293 examples of force-distance curves for each type of lipid bilayer. The d mean values obtained
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50 294 were 3.91 ± 0.46 nm, and 4.93 ± 0.47 nm for DOPC and DPPC bilayers, respectively. This
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53 295 thickness difference between fluid and gel phases was perfectly detected by AFM
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56 296 spectroscopy, in agreement with previous works^{25,43}. In the literature, Nagles and Tristram-
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59 297 Nagles⁴⁴ also determined the thicknesses of DOPC and DPPC fully hydrated bilayers using
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3 298 XRD and obtained a value of 3.6 nm for DOPC in the fluid phase and of 4.4 nm for DPPC in
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6 299 the gel phase. It should be noted that DOPC has two unsaturated 18 hydrocarbon chains and
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9 300 DPPC has two fully saturated 16 hydrocarbon chains (Fig1A). Unsaturation is the main
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11 301 reason that adversely affected chain elongation and molecular packing⁴⁵, making DPPC
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14 302 bilayers more ordered and thicker than DOPC bilayers at room temperature.
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16
17 303 Noteworthy, the value of the force at which the SLB ruptured (i.e., the breakthrough force)
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19 304 was higher for DPPC (2-3 nN) than for DOPC (<1 nN), in agreement with their respective
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22 305 phase states⁴⁶⁻⁴⁸. The lower absolute values found in the present study may be accounted for
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25 306 the lower ionic strength and the sharper AFM tip (radius of MNSL tip ~2 nm) in comparison
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28 307 to previous works.

308 **3.3.2. Mechanical properties of DOPC or DPPC liposomes:**

309 In this work, we show that AFM spectroscopy could be used to discriminate and compare the
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311 mechanical properties of very small liposomes (~150 nm) in different phase states.
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313 Calculations of the Young modulus E and bending modulus k_C , based on the shell theory
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315 model, were done on DOPC or DPPC liposomes respectively in fluid or gel phase. In order to
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317 limit the plastic deformation, force distance curves were recorded over 100 nm distance with a
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319 set point of 200 pN maximal force. Typical examples are shown in Fig. 3 B and D. In these
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321 conditions, the approaching and retracting curves were superimposed, demonstrating the
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323 elastic behavior of the membrane (not shown). The tip-membrane contact was defined as the
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325 point where significant positive slope appeared. According to the shell theory developed by

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3 318 Reissner⁴⁹ then Fery et al.^{16,50}, the Young modulus E of thin-shelled spherical micro-capsule
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6 319 under a point load scales with the bilayer's stiffness, k , which was deduced from the slope of
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9 320 the linear region of each force curve after the tip-membrane contact, i.e. in the small
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11 321 deformation region (Fig 3.B and D). As k strongly depends on the size of the individual
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14 322 liposomes²¹, it has to be normalized by the local radius of curvature R_c of the individual
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17 323 liposomes¹⁶ to describe their mechanical properties. Calculation of E requires the bilayers'
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20 324 stiffness k , the local radius of curvature R_c (see section 2.7.1), the bilayer's thickness d
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22 325 (Table1) and the Poisson ratio ν , taken as 0.5:

$$E = \frac{1}{C} \frac{kR_c \sqrt{3(1-\nu^2)}}{4d^2} \quad (3)$$

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29 326 In equation 3, C is a coefficient that accounts for the double deformation of an adsorbed shell
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32 327 object, such as liposomes, upon indentation by the AFM tip. Indeed, whereas membrane
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35 328 deformation occurs at the contact point between the tip and the liposome, simultaneous
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38 329 deformation also occurs at the contact area between the liposome and the substrate⁵¹. In their
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41 330 recent paper, Bery et al.⁵² produced calculations for a correction factor, C , to be applied as a
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44 331 function of the relative radii of the tip (20 nm) and liposome, and of the shell thickness
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47 332 relative to the radius of the liposomes (Table 1). Since that dimensions were similar for both
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50 333 the DOPC and DPPC liposomes, C was ~ 0.55 for both types of liposomes.

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52 334 In another way, the mechanical properties of liposomes can be also represented by the
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55 335 bending rigidity k_C , which is expressed in terms of the same parameters and is also common in
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58 336 the literature:

$$k_C = \frac{Ed^3}{12(1-\nu^2)} \quad (4)$$

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337 Calculation of the Young's modulus, regardless of liposome size, showed that the DPPC
338 liposome membranes (L_β) $E= 116 \pm 45$ MPa were significantly more elastic at 20°C than
339 DOPC liposome membranes (L_α) which exhibited a lower value of $E = 13 \pm 9$ MPa (Fig.4.A
340 and B; $p < 0.05$). Accordingly, the liposome membranes composed of DPPC were also stiffer
341 with $k_C = (15.5 \pm 6) \times 10^{-19}$ J (360 k_B T) than those composed of DOPC with $k_C = (0.9 \pm 0.6)$
342 $\times 10^{-19}$ J (22 k_B T) (Fig.4.C and D; $p < 0.05$). To verify that electrostatic repulsion was not
343 implicated in the force curves near the contact point, we compared the results acquired on
344 DPPC liposomes with varying surface charge, taken as the zeta potential and measured as
345 described in Makino et al.⁵³. The DPPC liposomes were positively charged with a zeta
346 potential of 18 mV in PIPES buffer (ionic strength $I=0.09$); and barely charged with a zeta
347 potential of 0.11 mV in PBS buffer (14 mM KH_2PO_4 , 200 mM Na_2HPO_4 , NaCl 1.36 M, KCl
348 20 mM; $I=1.99$; $\text{pH}= 7.2$). The Young's modulus showed a mean value of $E= 120 \pm 39$ MPa
349 in PBS buffer and no significant difference was found between the Young moduli values
350 obtained in the two buffers ($p > 0.05$). This result showed that force measurement of DPPC
351 membrane was not influenced by electrostatic interaction between the AFM probe and the
352 liposomes.

353 The Young modulus was also estimated using the R_c of the whole liposome, which involves
354 the two parameters H and W (equation A.1). The results showed that the elasticity of DPPC

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3 355 was higher (183 MPa) than that obtained by local R_c (116 MPa), whereas E was not
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6 356 significantly changed for DOPC. With either methods of R_c calculation, the results therefore
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9 357 showed that AFM indentation was able to detect significant differences in the elasticities of
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11 358 the two membranes depending on their fluid or gel phase state. However, the quantification of
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14 359 the mechanical properties was found very sensitive to calculation of R_c and to the liposome
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17 360 geometry. For these reasons, we chose to compare the liposomes using the method based on
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20 361 the local R_c using Gwyddion in order to avoid any effect due to the form adopted by the
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22 362 liposomes upon adsorption.

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25 363 In conclusion, the structural differences between the fluid-phase DOPC and gel-phase DPPC
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28 364 bilayers as evidenced by DSC and XRD, induced by the unsaturation of the acyl chains
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31 365 (Fig1), resulted in significant difference in their respective mechanical responses. Hence,
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33 366 AFM force spectroscopy proved a sensitive method to compare the mechanical properties of
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36 367 small liposomes with different lipid compositions and phase states. Only few reports exist that
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39 368 have evaluated these mechanical parameters for similar systems of lipid membrane in 3D,
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42 369 especially for DOPC or DPPC using AFM spectroscopy (Table 2). Of all these studies, only
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45 370 Liang et al.²¹ used AFM spectroscopy comparatively, on liposomes with increasing addition
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48 371 of cholesterol. Liang et al.²¹ obtained Young's modulus of 1.97 ± 0.75 MPa, for liposomes in
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51 372 the fluid phase composed of egg PC (mixture of saturated and unsaturated polar lipids; mostly
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54 373 in the form of SOPC – stearyl-oleoyl-phosphatidylcholine). Using an optical method,
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56 374 Meleard et al.⁵⁴ and Duwe et al.⁵⁵ respectively obtained bending rigidities of $1.27 \pm 0.09 \times 10^{-7}$

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3 375 ^{19}J ($31 k_{\text{B}}\text{T}$) or $1.15 \times 10^{-19}\text{J}$ ($28 k_{\text{B}}\text{T}$) for liposomes composed of the saturated polar lipid
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5 376 DMPC (1,2-dimyristoyl-sn-glycero-3-phosphocholine) in fluid phase at 40°C ($T_{\text{m}} = 24^{\circ}\text{C}$).
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8 377 Rawicz et al.⁴⁵ found values of $\sim 0.9 \times 10^{-19}$ J for bilayers of various synthetic 18:1
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11 378 phospholipids. Another group, Hantz et al.⁵⁶ used the osmotic swelling method and found
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14 379 values of the Young's modulus of 15 MPa for DOPC liposomes in the fluid phase at 20°C .
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17 380 Delorme and Fery¹⁶ obtained Young's moduli values of 110 ± 15 MPa for DPPC liposomes in
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20 381 the gel phase. Taking only account of the phase state conditions, the values available in the
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22 382 literature (Table 2) are in the same order of magnitude in comparison to the results showed in
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25 383 this work. Differences between studies could be attributed mainly to the nature of the lipids
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28 384 (carbon chain length, number of unsaturation), the temperature, the chosen technique or, for
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31 385 indentation studies, the mathematical model chosen for the calculation of R_{c} or E ^{57,58}. The
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34 386 presented results show that AFM indentation of liposome is a sensible method for comparison
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37 387 between different lipid membranes.
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40 388 For the sake of comparison with the SUV (tridimensional organization), the elasticity of
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43 389 DPPC or DOPC membranes was calculated on the SLB (two dimensional organization) using
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46 390 AFM load curves performed in the same conditions as for the breakthrough force
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49 391 measurement. Only a larger MLCT probe was used for better sensibility. By attempting the
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52 392 fit of equation (5), the Young's moduli (noted E_{SLB}) and bending modulus (noted $k_{\text{C-SLB}}$)
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55 393 were calculated using the classical Hertz model:
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$$F = \frac{4E_{SLB}\sqrt{R_{tip}}\delta^{\frac{3}{2}}}{3(1-\nu^2)} \text{ or } E_{SLB} = \frac{3k(1-\nu^2)}{4\sqrt{R_{tip}}\delta} \quad (5)$$

394 For which corresponds a bending modulus:

$$k_{C-SLB} = \frac{E_{SLB}\delta^3}{24(1-\nu^2)} \quad (6)$$

395

396 Where the parameters are the same already cited for liposome membranes. In these equations,

397 δ is the indentation distance and R_{tip} is the nominal radius of AFM MLCT tip (~ 20 nm).

398 The use of contact mechanics using the Hertz model on SLB is limited by the effect of

399 confinement of the sample between the tip and the underlying substrate^{59,60}. Furthermore,

400 lipid bilayers are not anisotropic materials. However, previous investigations suggested that

401 this calculation yet has comparative interest^{25,61}. On SLBs, the Young modulus (E) of DOPC

402 was found to be 27 ± 8 MPa with a corresponding bending modulus $k_C = (0.88 \pm 0.25) \times 10^{-19}$ J,

403 while the E and k_C of DPPC were found to be 31 ± 12 MPa and $(2.03 \pm 0.79) \times 10^{-19}$ J,

404 respectively. Therefore, the E and k_C parameters of DPPC and DOPC were less distinct when

405 measured on SLB than on SUV (Table 2). Meanwhile, more dispersed E values were reported

406 in the literature for SLBs than for liposomes, thereby indicating the higher sensitivity of this

407 measurement upon experimental conditions in the case of SLB^{25,61,62} (Table 2). To conclude,

408 it is showed that membranes in the gel phase were more elastic and stiffer than the

409 membranes in fluid phase regardless the lipid organization (2D or 3D). However, the values

410 of E and k_C were more robust when measured on liposomes than on SLBs, where the presence

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3 411 of the solid support affected the results depending on the indentation distance²⁵. These
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5 412 comparisons show that liposomes are adequate systems to determine the elastic properties of
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8 413 lipid membranes. On the other hand, measurement of the breakthrough force is relevant for
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11 414 SLBs.

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14 415 The structure of molecules and the intermolecular interactions that lead lipid molecules to
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16 416 self-assembly in bilayers, have a significant impact on the rigidity of these bilayers^{63,64}. The
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18 417 elasticity of the membrane allows it to accommodate strain without failure, which is essential
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20 418 in many applications where the membrane need to resist shear stress, e.g. in transdermal
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22 419 application, in blood vessels, the epithelial cells of the gastrointestinal tract, etc. The
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24 420 hydrophobic interactions between the lipid molecules, in particular Van der Vaal interactions,
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26 421 are the major responsible of the fluidity and the rigidity of the membrane. The double bond in
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28 422 cis conformation interferes with hydrocarbon chain packing and destroys the cooperativity of
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30 423 the chain interactions in the bilayer^{65,66}.

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33 424 The presence of this double bonds reduces the hydrophobic interactions by increasing the
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35 425 distance between the hydrophobic moieties which decreases the stiffness of membrane⁴⁵.

36 426 **IV. CONCLUSION:**

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39 427 The nano-indentation of DOPC and DPPC liposomes by AFM probe at low force load was
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41 428 able to provide local and discriminant information on the elastic properties of bilayer
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43 429 membranes in 3D organization without plastic deformation. The Young's moduli E and
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45 430 bending rigidity values k_C of gel phase DPPC membranes is significantly higher than that of
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3 431 fluid phase DOPC ones at 20°C, in agreement with their different phase state. The
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6 432 perspective of this work is to investigate the mechanical properties of biological membranes
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9 433 with complex chemical composition and fluid/gel phase coexistence. Thanks to the high
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11 434 lateral resolution of AFM, it is expected that phase separation and correlated nanomechanical
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14 435 contrast may be measured directly on model liposomes or even biological vesicles.

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21
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29 440 CEPIA and Région Bretagne under the grant ARED 8806.

32 441 **VI. APPENDIX:**

35 442 The Young modulus values were also obtained using radius of curvature R_C of the whole
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38 443 individual liposomes. It was calculated using H and W of the individual liposomes⁶⁷
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41 444 , as:

$$44 R_C = \frac{0.25W^2 + H^2}{2H} \quad (A.1)$$

48 445 The mechanical data obtained from literature were expressed either as the Young modulus E ,
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51 446 the bending rigidity k_C or both. To complete and compare literature information in table 2; we
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53
54 447 used the following equations to provide both parameters for each cited reference:

56
57 448 For liposome (shell model):

$$k_C = \frac{Ed^3}{12(1-\nu^2)} \Rightarrow E = \frac{12(1-\nu^2)k_C}{d^3} = \frac{9k_C}{d^3} \quad (A.2)$$

449 For supported lipid bilayer (Hertz model):

$$k_{C-SLB} = \frac{E_{SLB} d^3}{24(1-\nu^2)} \Rightarrow E_{SLB} = \frac{24(1-\nu^2)k_{C-SLB}}{d^3} = \frac{18k_{C-SLB}}{d^3} \quad (A.3)$$

450 E : Young modulus

451 k_C : bending rigidity

452 ν : Poisson coefficient (0.5)

453 d : membrane thickness

454 For the publication where the membrane thickness not shown, the following values were used

455 to calculate the elasticity or the bending rigidity:

456 $d(\text{DMPC}) = 3.6 \text{ nm}$ ⁴⁴

457 $d(\text{DOPC}) = 3.9 \text{ nm}$ ^(present work)

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657 TABLES

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659 Table1: Geometrical parameters of dioleoylphosphatidylcholine (DOPC) or
660 dipalmitoylphosphatidylcholine (DPPC) liposomes and their corresponding membrane
661 thicknesses. The height (H) and width (W) mean values were measured using the cross section
662 of AFM images of the liposomes and used to calculate the volume of the adsorbed liposomes,

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3 663 from which the mean diameter of a sphere of equivalent volume was deduced (Eq.2). The
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5 664 mean values of DOPC and DPPC membrane thicknesses were measured using the jump
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8 665 distance d from breakthrough force curves ($n=30$) recorded in PIPES/NaCl/CaCl₂ buffer, pH
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11 666 6.7 at 20°C.
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Lipid	Mean height H of liposomes by AFM (nm)	Mean width W of liposomes by AFM (nm)	Mean diameter of equivalent sphere by AFM (nm)	Mean diameter of liposomes by DLS (nm)	Membrane thickness by AFM (nm)
DOPC	59 ± 27	310 ± 97	150 ± 90	120 ± 69	3.91 ± 0.41
DPPC	151±48	448 ± 154	296 ± 88	132 ±73	4.93 ± 0.47

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670 Table 2: Comparison of the bending modulus (k_C) and Young modulus values (E) of

671 liposomes or of supported lipid bilayers in the fluid or gel phase reported in the literature

672 using different techniques as indicated. Abbreviations stand for: EggPC = mixture of

673 unsaturated (54.8 wt. %) and saturated (45.2 wt. %) phosphatidylcholine; DMPC = 1,2-

674 dimyristoyl-sn-glycero-3-phosphocholine; ESM = egg sphingomyelin; Chol = cholesterol.

675 Values in italics were calculated by the authors using calculations described in the Appendix

676 (eq.A.2; A.3)

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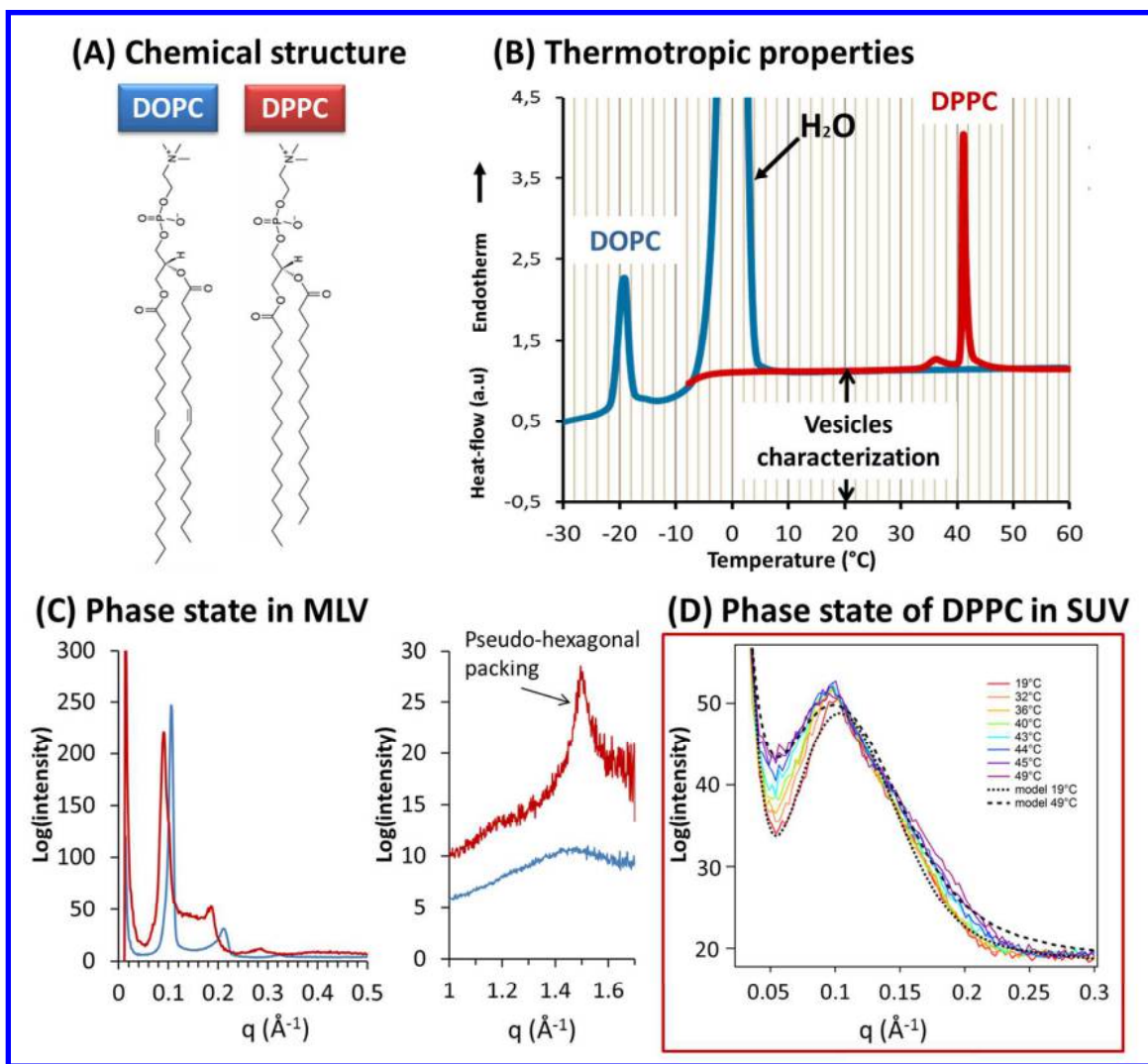
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Reference	Lipid	Technique	Phase	Bending modulus k_C ($\times 10^{-19} \text{J}$)	Young modulus E (MPa)
Liposomes (3D)					
Present work	DOPC	AFM spectroscopy	Fluid	0.9	13
	DPPC	AFM spectroscopy	Gel	15.50	116
16	DPPC	AFM spectroscopy	Gel	13.54	110
21	Egg PC	AFM spectroscopy	Fluid	0.27	1.97
45	synthetic phospholipids (18:1)	Micropipette pressurization	Fluid	0.90	12.70
54	DMPC	Phase contrast microscopy	Fluid	1.27	26
55	DMPC	Phase contrast microscopy	Fluid	1.15	24
56	DOPC	Osmotic swelling	Fluid	1.06	15
Supported bilayers (2D)					
Present work	DOPC	AFM spectroscopy	Fluid	0.88	27
	DPPC	AFM spectroscopy	Gel	2.03	31
28	DOPC	AFM spectroscopy	Fluid	1.69	19.3
	DPPC	AFM spectroscopy	Gel	2.33	28.1
61	DOPC	AFM spectroscopy	Fluid	4.61	80
	ESM/Chol	AFM spectroscopy	Liquid ordered	12.94	140
62	DOPC	AFM spectroscopy	Fluid	8.65	150
	ESM/Chol	AFM spectroscopy	Liquid ordered	27.72	300

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694 FIGURES:



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697 Figure 1: (A) Molecular structures of the unsaturated DOPC (dioleoylphosphatidylcholine)
 698 and saturated DPPC (dipalmitoylphosphatidylcholine). (B) Differential scanning calorimetry
 699 thermograms of DOPC (blue trace) and DPPC (red trace) multilamellar vesicles recorded on
 700 heating at $2^{\circ}\text{C}\cdot\text{min}^{-1}$. (C) X-ray diffraction patterns of DOPC and DPPC fully hydrated
 701 multilamellar vesicles recorded at 20°C at small (left) and wide (right) angles. (D) Small
 702 angle X-ray Scattering of DPPC SUV recorded on heating. Unilamellar SAXS model at low

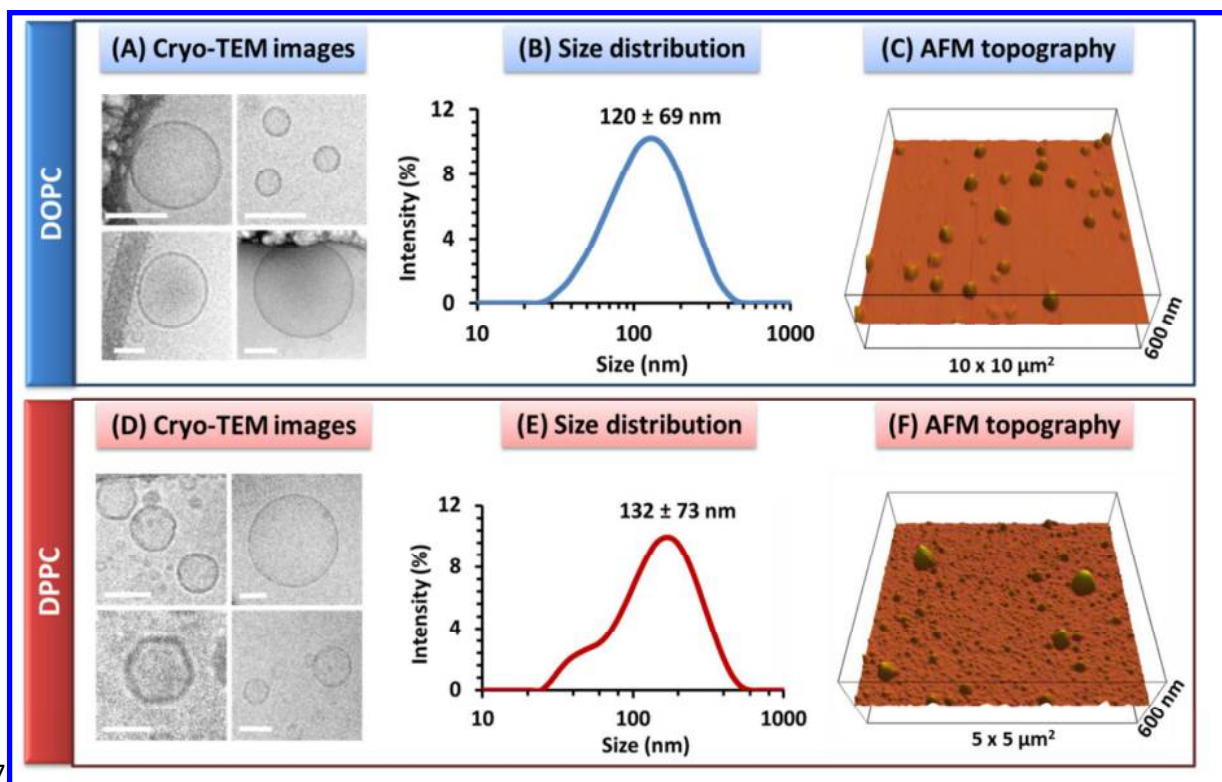
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3 703 and high temperature are superimposed. All experiments were performed in aqueous
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6 704 PIPES/NaCl/CaCl₂ medium at pH = 6.7.
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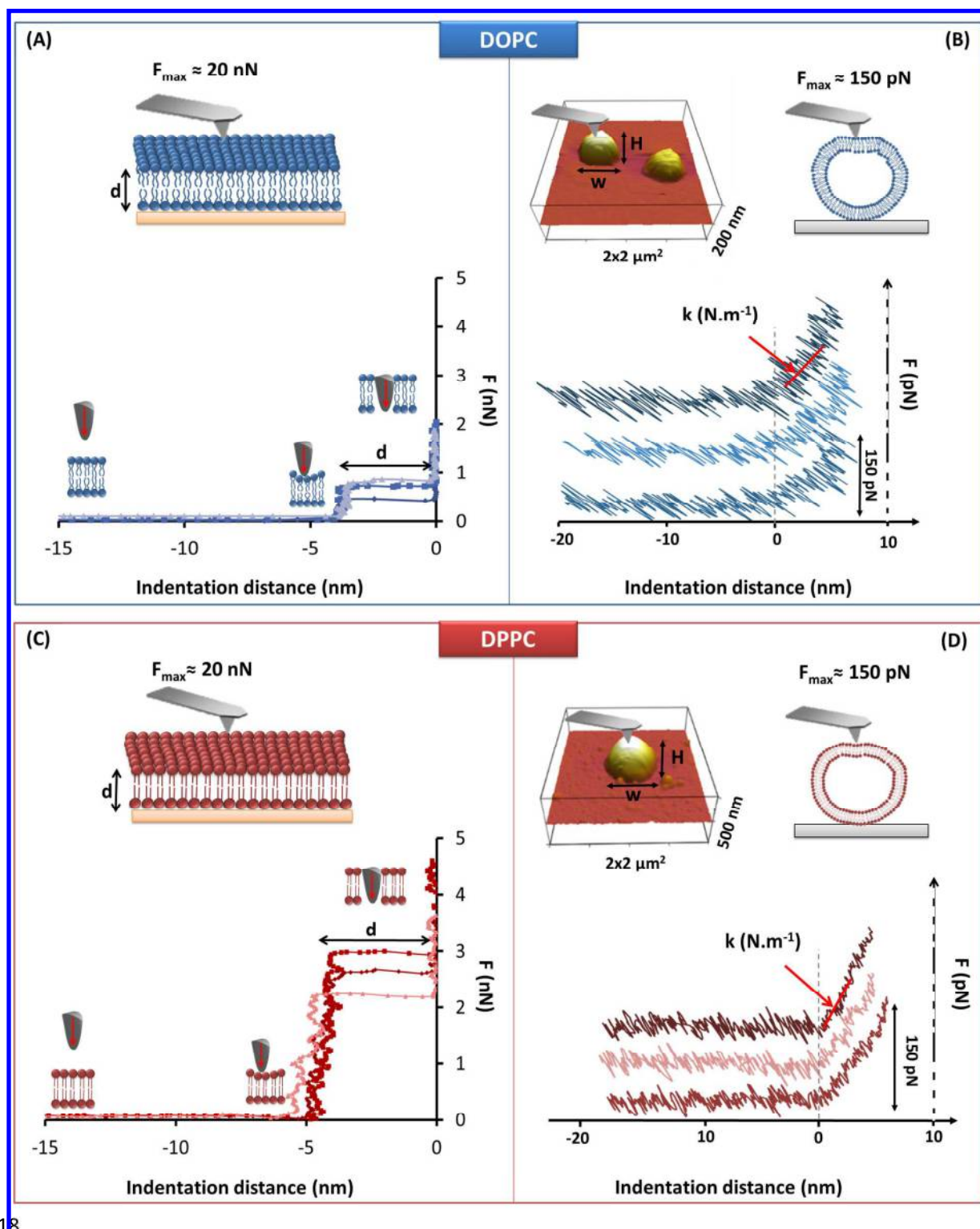
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708 Figure 2: Characterization of the shape and size of DOPC (dioleoylphosphatidylcholine) or
 709 DPPC (dipalmitoylphosphatidylcholine) liposomes. (A) and (D) Cryo-TEM images of the
 710 DOPC and DPPC liposomes, respectively; scale bars are 100 nm. (B) and (E) Dynamic light
 711 scattering size distribution in intensity of the DOPC and DPPC liposome suspensions,
 712 respectively; and (C) and (F) typical AFM 3D images of DOPC and DPPC liposomes,
 713 respectively. All experiments were performed in aqueous PIPES/NaCl/CaCl₂ medium at pH =
 714 6.7 and at 20°C.

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720 Figure 3: AFM indentation measurement on DOPC (dioleoylphosphatidylcholine, in blue) or

721 DPPC (dipalmitoylphosphatidylcholine, in red) liposomes adsorbed on silicon substrate. (A)

722 and (C) show the breakthrough force curves obtained as a result of tip penetration into DOPC

723 or DPPC supported lipid bilayers, respectively. The average bilayer thickness value d was724 measured from the jump-through distance ($n=50$). (B) and (D) show typical force curves

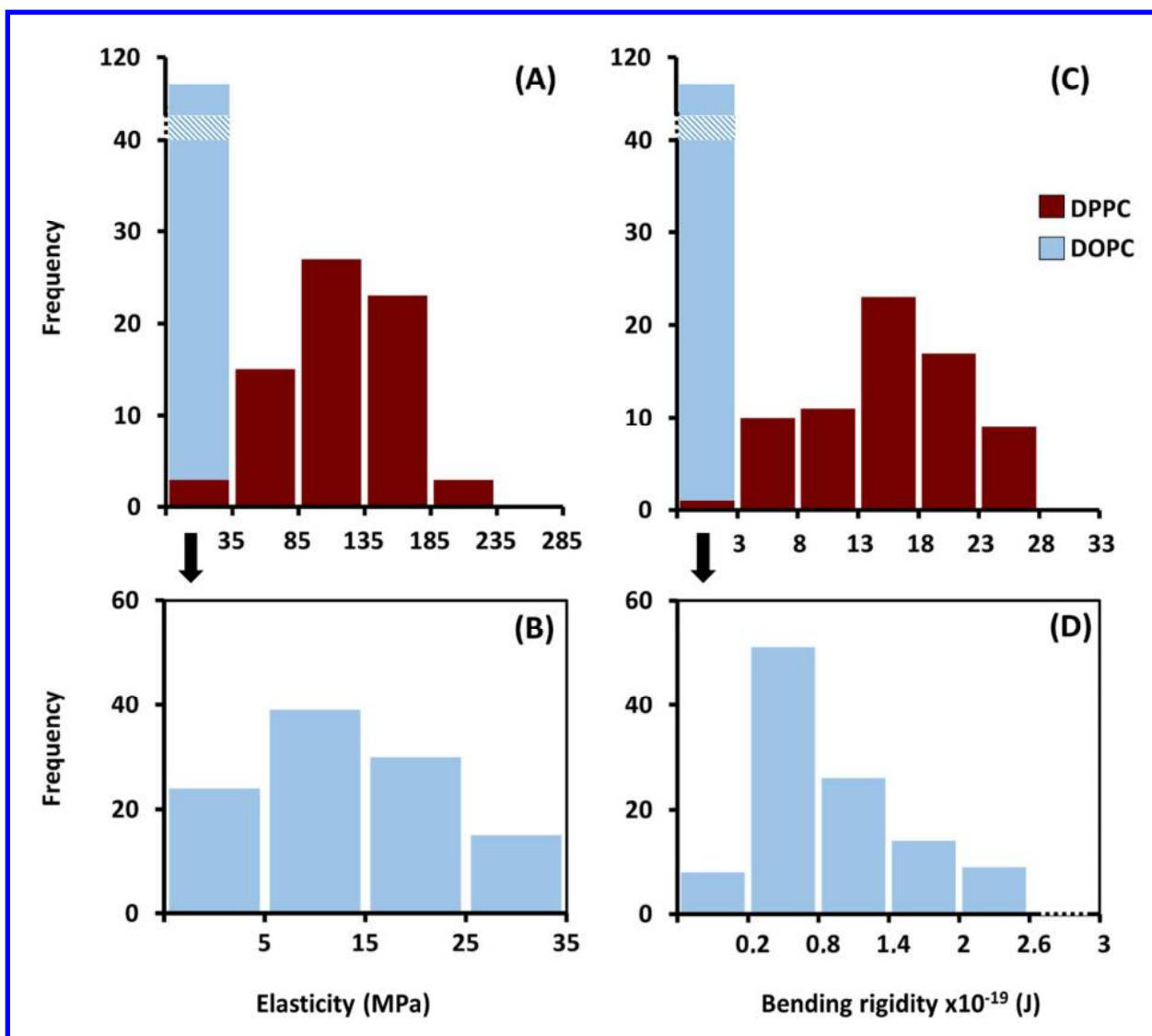
725 acquired during indentation of, respectively, DOPC or DPPC liposomes in the elastic regime;

726 in order to infer the bilayer stiffness k from the slope after tip-membrane contact. For the sake

727 of clarity, the force curves are shifted along the Y axis. All measurements were recorded in

728 aqueous PIPES/NaCl/CaCl₂ medium at pH = 6.7 and at 20°C.

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734 Figure 4: Distributions of the Young moduli E and of the bending rigidity k_C , independent of735 liposome size, of DOPC ($n=108$) and DPPC ($n=71$). (A) and (C) show the superposition of736 both DOPC and DPPC frequency distributions for E and k_C , respectively. (B) and (D) show

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3 737 respective enlargements of the E and k_C frequency distributions for the DOPC liposomes. All

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6 738 measurements were recorded in aqueous PIPES/NaCl/CaCl₂ medium at pH = 6.7 and at 20°C.

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741 **Table of contents graphic**

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