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Magnetic Nano-composite Micelles and Vesicles

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Abstract: Novel magnetic nano-composites are obtained by the self-assembly in water of polypeptide based diblock copolymers polybutadiene-\(b\)-poly(glutamic acid) combined with hydrophobically modified \(\gamma\)-Fe\(_2\)O\(_3\) nanoparticles. These hybrid supramolecular objects are either filled micelles (3-d) or hollow vesicles with a magnetic membrane (2-d), which deformation under an applied magnetic field has been evidenced.

Graphic for TOC:

[Artist view of a magnetic vesicle made of a diblock copolymer membrane confining magnetic nanoparticles. The shape is cut in half so that one sees the monolayer of nanoparticles (red) embedded in the hydrophobic blocks (yellow) of the bilayer, with its hydrophilic blocks (blue) exposed to both the outer and inner media. The interior is free for encapsulation of aqueous species.]
High or low molecular weight synthetic amphiphiles of many types have been revealed as interesting building blocks that can lead to highly ordered self-assembled structures in water.\cite{1} Depending on their molecular architecture and on external parameters such as temperature or pH, numerous morphologies have now been caught up including micelles, vesicles, tubules and more complicated morphologies reminiscent of internal compartments of biological cells.\cite{1,2} The bilayer formation obtained by the self-assembly of certain lipids in water is certainly the most interesting morphology both for materials science and biological perspectives.\cite{2,3} Nevertheless, applications involving these so-called liposomes are generally limited by their lack of stability related to the small hydrophobic layer thickness forming the core of the liposome membrane.\cite{4} Different groups reported on the vesicle formation from block copolymers in water (called polymersomes) and discussed on the improvement of the mechanical properties of the membrane obtained.\cite{5} Recently, vesicles have also been obtained from peptide-based diblock copolymers in our group\cite{6} and are called peptosomes.\cite{7}

In this context, the introduction of nanoscopic inorganic objects in self-organized structures is still a challenge for physico-chemists, even if Nature produces and uses frequently these hybrids. For example, colloidal particles of calcium carbonate or phosphate\cite{8} or iron oxides\cite{9} are commonly observed in organisms, where they are often confined in vesicles. In addition to the biomimetic aspect, the introduction of nanometric inorganic compounds in a self-assembled organic matrix raises the question of the resulting structure of the hybrids. Two extreme cases are possible: the inorganic nanoparticles can be either diluted and used as local probes for scattering methods, or they can really participate to the supramolecular structure and modify the resultant properties. A pioneering work on the entrapment of inorganic particles in vesicles was done by Fendler et coll.\cite{10} Previous studies on polymers and nanoparticles have dealt with the stabilization of these colloidal particles with low or high molecular weight polymers\cite{11} or with their introduction into macroscopic
polymeric gels.[12] The topic here is somehow different, for we are interested in the inclusion of magnetic nanoparticles into supramolecular objects at the mesoscopic scale and in the study of the resulting structures and properties. Recently, such vesicles and nanocomposites of inorganic moieties and peptide-type amphiphiles have been developed respectively by Katagiri et al.,[13] Stupp et al.,[14] and Deming et al.[15]. The closest approach to ours concerns extremely well defined clusters of magnetic nanoparticles confined in micelles of diblock copolypeptides,[16] except that we are looking here for larger objects exhibiting a more complex magnetic response that mere magnetization.

We report indeed on the preparation of a novel type of hybrid colloids, based on the association of several polymeric systems and ferrofluids. On the one hand, we use inorganic nanoparticles made of a magnetic iron oxide that respond to a magnetic field of low intensity. On the other hand, the organic part is made of mesoscopic structures (vesicles and micelles) self-assembled from amphiphile polybutadiene-\(b\)-poly(glutamic acid) diblock copolymers. Those PB-\(b\)-PGA copolymers bearing a rod-coil structure with a cross-linkable hydrophobic block and a hydrophilic peptidic block have been synthesized recently by combining anionic polymerization and ring-opening polymerization.[6] Due to a helix-coil transition of the PGA block, such peptosomes behave as stimuli-responsive nanocapsules as they respond to a pH-change by a variation of their hydrodynamic diameter as large as 50%.[6,17] As for the magnetic nanoparticles, they come from ferrofluids which are colloidal suspensions of nanometric magnetic grains stabilized either by electrostatic charges in aqueous media,[18] or by appropriate tensioactives in organic solvents.[19] Embedded in supra-macromolecular objects formed by diblock copolymers, these inorganic nanoparticles are used as structural probes in small angle neutron scattering (SANS) experiments. Benefiting from the strong contrast of iron oxide compared to polymer and solvent, we measure the structure factor inside the aggregates \(S_{\text{intra}}(q)\). On the one hand, this curve reflects the interactions between the
nanoparticles and possibly the mean inter-particle distance, hence their local concentration.\[20\]

On the other hand, \( S_{\text{intra}}(q) \) can be identified at low \( q \) to the global form factor \( P_{\text{agg}}(q) \) of the object that they decorate.\[21\] Another guiding idea of this mineralization is to bring a magnetic response to the whole object, either an induced shape change towards an ellipsoid or a more complex shape or to trigger the delivery of an active substance by the application of a field. It has been soon predicted indeed that liposomes exhibit an ellipsoidal deformation in strong magnetic fields.\[22\] Depending on the sign of the magnetic susceptibility anisotropy \( \Delta \chi = \chi_{||} - \chi_{\perp} \) of the bilayer, the shape is either an elongated (prolate) or a flattened (oblate) ellipsoid. However, this phenomenon requires field intensities \( B \) of 1 Tesla \( (10^4 \text{ G}) \) at least, due to a too small natural anisotropy \( \Delta \chi < 10^{-7} \) for a classical lipid (lecithin) bilayer.\[23\] Better magnetically alignable membranes can be obtained by adsorption of paramagnetic ions like lanthanides.\[24\] But theory predicts that the greatest enhancement of the magnetic response would be obtained by filling the interiors of the vesicles with magnetic nanoparticles.\[25\] Anisotropic SANS patterns have been reported for pure lipid vesicles in strong magnetic fields \( B \) from 1 to 4 Tesla.\[26\] Using the same spectrometer (PAXY, LLB, CEA-Saclay, France), we show here by SANS not only the orientation but also the deformation of copolymer membranes stuffed with magnetic nanoparticles.

This work starts by the verification that those organic and inorganic systems can effectively be combined together to generate well defined and properly dispersed objects. Therefore we examine a series of PB-\( b \)-PGA copolymers which differ by the length of their polypeptide PGA block. We present a short series of samples representative of the micellar and vesicular structures confining magnetic nanoparticles in either 3 or 2 dimensions respectively. We start from the following original structures as building bricks:

- PB_{48-}\( b \)-PGA_{114} and PB_{48-}\( b \)-PGA_{145} which self-assemble in water as micelles of hydrodynamic diameters \( d_{\text{H}}=60 \text{ nm} \) and \( d_{\text{H}}=70 \text{ nm} \) respectively as determined by DLS, the
internal diameter of the hydrophobic core measured by SANS being $d_{\text{int}}=14$ nm for both.\footnote{27}

- PB$_{48}$-b-PGA$_{56}$ which forms closed membranes in water, \textit{i.e.} vesicles characterized by an outer diameter $d_H=100$ nm and a hydrophobic thickness $\delta_{\text{PB}}=14$ nm inside the bilayer measured by SANS matching the contrast of PGA blocks.\footnote{27}

- a ferrofluid in dichloromethane denoted S2-CH$_2$Cl$_2$, consisting of maghemite nanoparticles with a characteristic diameter $d_0=7.6$ nm (see Supplemental Information I for the methods to measure and to decrease their polydispersity in sizes and to graft an appropriate surfactant).

\textit{Table 1: Short summary of samples and self-assembled hybrid objects that have been prepared.}

<table>
<thead>
<tr>
<th>Copolymer</th>
<th>Ferrofluid</th>
<th>Eq. ferro</th>
<th>Solvent</th>
<th>System</th>
<th>$\Phi_{\text{ferro}}$</th>
<th>$d_H$</th>
<th>$d_{\text{int}}$</th>
<th>morphology</th>
<th>$N_{\text{part}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>PB$<em>{48}$-b-PGA$</em>{145}$</td>
<td>S2-CH$_2$Cl$_2$</td>
<td>1</td>
<td>H$_2$O</td>
<td>A</td>
<td>0.26</td>
<td>429</td>
<td>123</td>
<td>Micelles</td>
<td>80000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>A</td>
<td>0.043</td>
<td>225</td>
<td>95</td>
<td>Micelles</td>
<td>12000</td>
</tr>
<tr>
<td>PB$<em>{48}$-b-PGA$</em>{114}$</td>
<td>S2-CH$_2$Cl$_2$</td>
<td>1</td>
<td>H$_2$O</td>
<td>A</td>
<td>0.14</td>
<td>333</td>
<td>108</td>
<td>Micelles</td>
<td>38000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>A</td>
<td>0.021</td>
<td>260</td>
<td>110</td>
<td>Micelles</td>
<td>18000</td>
</tr>
<tr>
<td>PB$<em>{48}$-b-PGA$</em>{56}$</td>
<td>S2-CH$_2$Cl$_2$</td>
<td>1</td>
<td>H$_2$O</td>
<td>B</td>
<td>0.30</td>
<td>624</td>
<td>256</td>
<td>Vesicles</td>
<td>12000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td></td>
<td>B</td>
<td>0.26</td>
<td>208</td>
<td>104</td>
<td>Vesicles</td>
<td>2700</td>
</tr>
</tbody>
</table>
Micelles of PB_{48-b}-PGA_{114} and PB_{48-b}-PGA_{145} mixed with surfacted ferrofluid S2-CH2Cl2 (A) lead to a stable (over months) dispersion of magnetic micelles in water, as attested by the structure factor deduced from SLS and SANS. Combined with 1 mass equivalent of ferrofluid, both diblock copolymers lead to fully dispersed suspensions. On the contrary at 2 mass equivalents, a coagulum is still present after several weeks, leading to much lower concentrations of dispersed objects (Table 2). The radii of gyration of the objects are of the order of 100 nm and smaller than (half of) the hydrodynamic diameters dH which are ranging from 225 nm to 430 nm (Table 2). The ratios \(2R_g/d_H\) between 0.6 and 0.8 are close to the hard sphere value \((3/5)^{0.5} \approx 0.77\). There is a large (several-fold) increase of the outer diameters compared to unloaded micelles (dH=60 nm and 70 nm for pure PB_{48-b}-PGA_{114} and PB_{48-b}-PGA_{145} respectively), which means the self-assembly of the diblocks is significantly modified by the presence of the nanoparticles. The position of the structure peak around \(q_{\text{max}} \approx 8 \times 10^{-2} \text{ Å}^{-1}\) (Figure 1) corresponds to a short inter-particle distance d_{max} \approx 8 nm. The hydrophobic cores are thus filled with nanoparticles at a volume fraction

\[
\Phi_{\text{ferro}} = \pi \left( \frac{d_0}{d_{\text{max}}} \right)^3
\]

about 45%, of the order of the dense packing value. This high encapsulation yield together with the global 3-d shape of the aggregates are confirmed by a look at the TEM (Figure 2) and AFM images (Supplemental Information II) in the case of PB_{48-b}-PGA_{114}, which show large and thick baggies of inorganic particles of sizes comparable to the determinations of Table 2. The nanoparticles appear compactly confined in localized volumes. Those objects do not resemble at all to the tenuous fractal clusters observed when magnetic nanoparticles

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[a] mass equivalent of iron oxide compared to copolymer in the preparation mixture.
[b] code corresponding of the system related to Table 1 and to the text.
[c] volume fraction of ferrofluid in the final suspension (measured by titration of iron).
[d] hydrodynamic diameter measured by multi-angle DLS with CONTIN analysis.
[e] radius of gyration measured from a Guinier fit of the low angle light scattering.
[f] approximate number of nanoparticles per object (estimated by formulae in the text).
spontaneously aggregate. The over-concentration in these baggies compared to the very low global concentration of nanoparticles in the suspension (Φ\text{ferro}≈0.1%) means that the nanoparticles inside the micelles are almost in contact to each other, the remaining space inside the bags being taken by the PB blocks (about 55vol.%). In these peptide-based micellar systems, the average number of magnetic nanoparticles per micelle can be estimated by

\[ N_{\text{part}} = \frac{\pi}{6} \left( \frac{d_{\text{nl}}}{d_{\text{max}}} \right)^3 \] (see Table 2). Each micelle contains a few 10⁴ nanoparticles, which is two orders of magnitude larger than in aggregates of magnetic nanoparticles obtained by the addition of coagulating homopolymers, or other self-assembling diblock copolymers.

Figure 1: Intra-aggregate structure factor \( S_{\text{ intra}}(q) \) of the hydrophobic nanoparticles S2-CH2Cl2 associated to PB₄₈-b-PGA₁₁₄ and PB₄₈-b-PGA₁₄₅ copolymers. The dash line corresponds to a slope \( q^{-4} \). The square dots represent simulations of the SANS intensities for spherical micelles of the given radii. The few points at lowest q come from SLS and enable an estimate of the radii of gyration by a standard Guinier fit. The inset shows the SANS curve of the dilute ferrofluid fitted by the form factor \( P_{\text{ferro}}(q) \) of polydisperse spheres (\( d_0=7.6 \) nm and \( \sigma=0.25 \)).

Figure 2: TEM picture of hybrid copolymer micelles loaded with magnetic nanoparticles obtained by combining PB₄₈-b-PGA₁₁₄ and 1 mass equiv. of surfacted ferrofluid S2-CH2Cl2. The scale bar measures 400 nm.
Vesicles of PB$_{48}$-b-PGA$_{56}$ still form in water even in the presence of S$_2$-CH$_2$Cl$_2$ (B), as attests the structure factors deduced from SLS and SANS (Figure 3). Their outer diameter is significantly increased by the presence of the nanoparticles (d$_H$ equal to 620 nm and 210 nm for 1 and 2 mass equivalents respectively). The ratios 2R$_g$/d$_H$ of the radii of gyration (Table 2) over half of the hydrodynamic diameters are close to 1, the theoretical value for a hollow sphere. Due to their hydrophobic coating, the nanoparticles are confined in 2-d between the two leaflets of the copolymer bilayer, as proved together by SANS measurements, TEM pictures and AFM imaging (which in addition contains a valuable topographical information).

Figure 3 represents the intra-aggregate structure factor S$_{\text{intra}}$(q) measured by SANS for the hydrophobic nanoparticles associated to PB$_{48}$-b-PGA$_{56}$ diblock copolymer forming vesicles. In the small q-regime, S$_{\text{intra}}$(q) is identical to the form factor of the aggregate P$_{\text{agg}}$(q) and follows a power law with a slope approximately q$^{-2}$, completely different from the q$^{-4}$ slope of Figure 1. This q$^{-2}$ slope could be associated to 3-d aggregation of the nanoparticles under Diffusion or Reaction Limited Aggregation processes (DLA or RLA). However, both TEM pictures of Figure 4 and AFM measurements (which are discussed in Supplemental Information II) do not correspond to such a fractal cluster formation. On the contrary, one can calculate the thickness $\delta$ of the layer forming the vesicle membrane from the asymptotic Kratky-Porod approximation in this q-region for the form factor of a flat infinite sheet of thickness $\delta$,$^{[31]}$ 

$$\ln[S_{\text{intra}}(q)q^2] \approx \ln(I_0) - q^2\delta^2/12,$$

where $I_0$ is proportional to the surface area.

We get $\delta_{1\text{eq}}=8.6$ nm and $\delta_{2\text{eq}}=17.7$ nm respectively for the system with 1 equivalent and 2 equivalents of ferrofluid. Note that here the SANS contrast is given by the nanoparticles only and that $\delta$ is an inorganic thickness which do not take into account the copolymer. As the typical diameter $d_0$ of the nanoparticles is 7.6 nm, one can conclude that this membrane contains either one or two layers of the magnetic colloids depending on the equivalent ratio at
preparation. Instead of the Kratky-Porod approximation, one can also simulate the form factor of vesicles with radius $R$ (taken as $d_{H}/2$) and membrane thickness $\delta$. The good adjustment of the curves confirms unambiguously the presence of flat sheets of thickness $\delta$, but their radius of curvature $R$ cannot be measured precisely in this q-range. At wider angle, the scattered intensities are dominated by the magnetic nanoparticles and thus $S_{\text{intra}}(q)$ reaches 1. The absence of correlation distance $d_{\text{max}}=2\pi/q_{\text{max}}(=8\text{nm})$ between neighboring nanoparticles for the preparation at 1 equivalent of ferrofluid corroborates the fact that the inorganic colloids form at most one monolayer inside the polymeric membrane. Assuming a close-packing of the nanoparticles within the shell, one estimates the maximum number of nanoparticles per vesicle (see Table 2) as

$$N_{\text{part}} \approx \frac{\pi}{\sqrt{3}} n_{\text{layers}} \left( \frac{d_{H}}{d_{0}} \right)^2$$

where $n_{\text{layers}}$ is the number of layers, identical to the number of equivalents (1 or 2). The bidimensional character of the objects is also visualized directly by TEM (Figure 4) and AFM (Supplemental Information II). Those patterns look neither like spontaneously aggregated nanoparticles\textsuperscript{[28]} nor to clusters induced by other (co)polymers\textsuperscript{[16,29,30]}. 
Figure 3: SANS measurements of the intra-aggregate structure factor $S_{\text{intra}}(q)$ of hydrophobic nanoparticles S2-CH$_2$Cl$_2$ (1eq and 2eq) associated to PB$_{48}$-b-PGA$_{56}$. The solid lines represent simulations of the SANS intensities in the same conditions as the experiments for hollow vesicles with following dimensions: $R_{1\text{eq}}=312$ nm and $\delta_{1\text{eq}}=8.7$ nm, $R_{2\text{eq}}=104$ nm and $\delta_{2\text{eq}}=17.7$ nm, respectively. Data for 1eq obtained in light water by full angular averaging of the 2-d PAXY spectrometer under zero field are added to show $S_{\text{intra}}(q)$ at lower $q$ (the overlap with data acquired on PACE is not perfect as the solvent is not contrast matched to polymer in that case). They lead however to the same $q^{-2}$ behavior. The few points at lowest $q$ come from static light scattering and give estimates of the radii of gyration from a Guinier fit. The inset shows Kratky-Porod plots, which slope give the values $\delta_{1\text{eq}}$ and $\delta_{2\text{eq}}$ of membrane thickness given above. A thickness $\delta_{\text{PAXY}}=10$ nm somehow larger than $\delta_{1\text{eq}}$ is found from the data measured on PAXY, presumably due to the contribution of the copolymer to the neutron scattering contrast.

Figure 4: a) TEM picture of a magnetic membrane made of PB$_{48}$-b-PGA$_{56}$ and 1 equiv. of S2-CH$_2$Cl$_2$. The scale bar measures 333 nm. b) 3-d view built from an AFM picture of the same sample (see Supplemental Information II) showing a flat copolymer membrane stuffed with magnetic iron oxide nanoparticles. Strong spreading on mica opens holes in the central part, whereas some pure copolymer bilayer is visible at the periphery.
The magnetic response of vesicles of PB_{48-b}-PGA_{56} with the membrane filled with 1eq S2-CH_{2}Cl_{2} (B) is studied using anisotropic SANS under an applied magnetic field. The system is dispersed in light water (H_{2}O) to work in (almost) pure nuclear contrast under field.\[32] The scattering patterns become clearly anisotropic when a magnetic field is applied to this sample (Figure 5a). The iso-intensity curves in the 3 \times 10^{-3}–2 \times 10^{-2} \text{ Å}^{-1} q-range are ellipses elongated perpendicularly to the field direction, with aspect ratios increasing from 1.1 to 1.3 when the field intensity B increases from 290 to 1450 G, almost independently of the value of q. The scattering patterns are also analyzed at a constant q, as a function of the angle θ with the field direction. The intensity profile I_{B=1050G}(θ) normalized by the isotropic intensity I_{B=0}(θ) is plotted in Figure 5b for several values of q between 4 \times 10^{-3} \text{ Å}^{-1} and 9 \times 10^{-3} \text{ Å}^{-1}. The ratio is maximum in the direction perpendicular to the field (I_{⊥}/I_{B=0}=1.1–1.15) and minimum in the direction parallel to the field (I_{∥}/I_{B=0}=0.6–0.7). We deduce an anisotropy parameter (I_{⊥}-I_{∥})/I_{B=0}≈0.5 for this field value B=1050 G.

However, it would be a mistake to consider that such a ratio measured on the scattering patterns represents directly the shape anisotropy of the vesicles under a magnetic field. One has indeed to remember that the window 3 \times 10^{-3}–5 \times 10^{-2} \text{ Å}^{-1} of scattering vectors q explored by these SANS measurements under field lies far above the Guinier regime, which is in the domain of small angle light scattering. Instead of the global shape of the vesicles, these experiments tell us about the deformation at the length scale of the membrane thickness δ.

Besides, we checked that this anisotropy under field does not originate from magnetic dipolar interactions between nanoparticles.\[32,33] The nuclear SANS pattern under field of the same ferrofluid at high concentration in water exhibits no anisotropy at all indeed. An anisotropy due to magnetic scattering being also excluded in light water,\[32] we deduce that the
anisotropy of the SANS patterns described here reflects a sensitivity to the magnetic field of the spatial distribution of the nanoparticles within the membranes. An exhaustive analysis of the field-induced anisotropy in the two principal directions || and ⊥ relatively to the magnetic field is presented in Supplemental Information III. It shows that either the membrane becomes stretched in the portions of shell crossed by the magnetic field (decrease of the apparent membrane thickness $\delta_{||}$), or almost equivalently that the magnetic nanoparticles move away from the magnetic poles. The remaining parts of the membrane remain unaffected ($\delta_{⊥}$ almost equal to the initial bilayer thickness at $B=0$ G). Until now, the deformation under field of hollow objects with a magnetic material confined in 2 dimensions has only been examined theoretically.$^{[34,35]}$ Even though more investigations with high resolution microscopy like cryo-TEM could help by directly imaging those vesicles “frozen” under field, the anisotropic SANS presented here already provides an unambiguous evidence of their shape sensitivity to the magnetic field. To the authors’ knowledge, this is the first described example of vesicles with a deformable magnetic membrane. Other magnetic shells have been indeed mentioned in literature,$^{[36,37]}$ but none as thin and floppy as these.

![2-dimensional SANS patterns under a magnetic field (directed horizontally). The anisotropy is observed in the q range $3 \times 10^{-3} - 2 \times 10^{-2}$ Å$^{-1}$. b) Intensity profile for $B=1050$ G normalized by the intensity for $B=0$ G vs. angle $\theta$ with the field direction for several values of q. The dashed intervals correspond to the angular regroupings respectively || and ⊥ (see Supplemental Information III).](https://example.com/sans_patterns_image.png)

*Figure 5*: Deformation of hollow magnetic shells (system B 1eq) studied by anisotropic SANS.

a) 2-dimensional SANS patterns under a magnetic field (directed horizontally). The anisotropy is observed in the $q$ range $3 \times 10^{-3} - 2 \times 10^{-2}$ Å$^{-1}$. b) Intensity profile for $B=1050$ G normalized by the intensity for $B=0$ G vs. angle $\theta$ with the field direction for several values of $q$. The dashed intervals correspond to the angular regroupings respectively || and ⊥ (see Supplemental Information III).
Conclusions

To summarize, the main challenge addressed in this paper is to build supra-macromolecular objects from magnetic nanoparticles and diblock copolymers. At first, the nanoparticles are used as structural probes to elucidate the transformation of the micelles and vesicles by the insertion of the inorganic material itself. Spherical micelles in water (samples A) are filled with a hydrophobic ferrofluid at volume fraction as high as 45%. With the same nanoparticles and another length of the PGA blocks of the copolymers, hollow fluid magnetic spheres are obtained (samples B). The organic amphiphile copolymers confine the inorganic nanoparticles coated by surfactants within the thin layer of PB blocks. Furthermore, anisotropic SANS data bring the experimental evidence of the capability to modify the shape of these mineralized membranes in response to a magnetic field of intensity as low as B=290 G.

At last, one can imagine potentialities of these superparamagnetic micelles and vesicles in biomedicine and biotechnology, related the specific properties of magnetic iron oxide: manipulation by an external magnetic field gradient, radio-frequency heating for cancer therapy,\textsuperscript{[38]} labeling of organs in Magnetic Resonant Imaging.\textsuperscript{[39]} Due to their small dimensions, these objects are potential candidates for intravenous injection. The possible toxicity for cells to contact iron oxide directly would not be an issue because it is embedded in copolymer. The soft magnetic shells are especially promising for drug delivery, their internal compartment being available for encapsulation of water soluble species. A pH change or the application a magnetic field could trigger the transient opening of the bilayer and the release of an encapsulated content.
Experimental

The copolymer – nanoparticles self-assemblys were prepared by mixing 20 mg of each copolymer with either 1 or 2 mass equivalents of surfacted ferrofluid S2-CH2Cl2 (Table 1). The mixture was stirred for 2 h in the open air to eliminate dichloromethane. The resulting pellet was slowly redispersed in 2 mL of aqueous solvent. Addition of 50 µL of NaOH 2 mol/L enabled to deprotonate ≈75 mol% of the acidic groups of the copolymer, raising pH up to ≈7. After three weeks of vigorous stirring at T=37°C, all suspensions were analyzed by static and dynamic light scattering (SLS and DLS) and small angle neutrons scattering (SANS). Thus we could check the dispersion state and study the equilibrium shapes and sizes. The isotropic SANS experiments were performed on the PACE spectrometer of the Laboratoire Léon Brillouin (CEA-Saclay, France). We used two configurations with sample-to-detector distance D=4.75m and two neutrons wavelengths λ=17 Å and λ=8 Å to cover q-ranges 5 10⁻³–5 10⁻² Å⁻¹ and 2 10⁻²–2 10⁻¹ Å⁻¹ respectively. The calculated contrast of neutrons scattering length densities is Δρ=5.0x10¹⁰ cm⁻² between γ-Fe₂O₃ and the solvent matching the copolymers, which is a 36:64 w/w mixture of D₂O and H₂O. The scattered intensity curves of the samples held in 1 mm quartz cells were corrected from incoherent background and normalized by the signal of water. Following a method used for other kinds of colloids – micelles nano-composites, the intensity curves of the copolymer – nanoparticles complexes were divided by the volume fractions Φ_ferro and by the form factor P_ferro(q) of the nanoparticles measured independently (see inset of Figure 1). This procedure yields the intra-aggregate structure factor S_intra(q) of the nanoparticles associated to the block copolymers. The form factors of hollow vesicles taking into account the polydispersity of radii and the experimental resolution of PACE were calculated with a program written by Dr. J. Oberdisse.
The anisotropic SANS experiments were performed on the PAXY spectrometer of the Laboratoire Léon Brillouin (CEA-Saclay, France) equipped with a 2-d detector made of 128 x 128 cells. The sample was placed between the poles of an electromagnet producing a homogeneous magnetic field, which intensity was measured with a Gaussmeter (Walker Scientific, Worcester, MA). We used two configurations with sample-to-detector distance D=6.775m and two neutrons wavelengths $\lambda=12$ Å and $\lambda=6$ Å to cover q-ranges $3 \times 10^{-3}$–$3 \times 10^{-2}$ Å$^{-1}$ and $7 \times 10^{-3}$–$5 \times 10^{-2}$ Å$^{-1}$ respectively. Due to complete isotopic exchange with atmosphere, the solvent of this PB$_{48}$-b-PGA$_{56}$/1eq S2-CH$_2$Cl$_2$ sample under magnetic field was pure H$_2$O, thus not matching the contrast of the copolymer, but insuring a negligible magnetic scattering of iron oxide.$^{[32]}$ Nevertheless, the nuclear contrast of the nanoparticles $\Delta \rho=7.5 \times 10^{10}$ cm$^{-2}$ was still three times larger than the contrast of the copolymer $\Delta \rho=2.5 \times 10^{10}$ cm$^{-2}$. A full angular averaging of the cells at constant q was realized for the isotropic scattering patterns. An anisotropic analysis was applied to the scattering patterns under magnetic field. By masking the appropriate cells of the 2-d detector, the intensity was averaged in angular sectors either (-15°,15°) along the field direction and thus called $I_{||}(q)$, or (75°,105°) around the perpendicular direction and denoted $I_{\perp}(q)$.

Static and dynamic light scattering was performed on a Macrotron spectrometer (Amtec, France) equipped with a He/Ne laser and a Malvern 7132 correlator. Aliquots of the samples (1 mL in cylindrical vials) diluted 20–100 times in 0.2 µm filtered water were immersed in a filtered toluene bath with temperature regulated at 25°C. For DLS, the autocorrelation functions of the intensity scattered at various angles from 45° up to 155° every 5° were recorded using a 0.5 µs sampling time for a duration of 15 min per angle. Using the standard program CONTIN, the dominant decay rate $\Gamma_{\text{max}}$ was extracted and plotted vs. $q^2$ to get the hydrodynamic diameters $d_H$ from the slope extrapolated in the low-q limit. For SLS, the light
intensity was time-averaged and corrected from geometrical and polarization effects so that the signal of filtered toluene appears independent of the angle.

The molar content of iron [Fe] (mol/L) was titrated by atomic absorption spectroscopy with a Perkin Elmer Analyst 100 apparatus after degrading the nanoparticles within the nanocomposites in boiling HCl (35%). The volume fraction of iron oxide was deduced from formula weight and mass density of \( \gamma \)-Fe\(_2\)O\(_3\) as: \( \Phi_{\text{ferro}}(\text{vol\%}) = 1.577 \times [\text{Fe}] \).

**TEM pictures** were recorded on a JEOL JEM100S microscope working at 100 kV. Samples were prepared by spraying a 1 g.L\(^{-1}\) solution of the block copolymer associated to ferrofluid onto a TEM grid using a homemade tool.

**AFM pictures** of dilute samples deposited on mica (5 \( \mu \)L at 0.1 g.L\(^{-1}\)) were obtained with a Nanoscope III multimode scanning probe microscope (Digital Instruments, Santa Barbara, CA) in the Tapping Mode in air at a 512*512 pixels resolution and a 0.9 Hz scan rate.

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**References:**


Supplemental Information

I. Preparation and characterization of the surfacted ferrofluid S2-CH₂Cl₂

In order to get a ferrofluid stable in dichloromethane, we first need to prepare a precursor aqueous ferrofluid by alkaline co-precipitation of FeCl₂ and FeCl₃ salts. This ionic ferrofluid consists of negatively charged superparamagnetic Fe₃O₄ (magnetite) nanocrystals in an aqueous medium at high pH. Acidification and surface sign reversal with HNO₃ is followed by complete oxidation by Fe(NO₃)₃, leading to γ-Fe₂O₃ (maghemite) nanoparticles at pH≈1. After several washing steps to remove ions in excess, this acidic ferrofluid can be used “as is”, resisting against sedimentation even in strong magnetic fields or field gradients. For dispersion in non aqueous media, further functionalization has to be performed by appropriate ligands, for example by oleic acid for most alkanes. In the present study, the surface of the nanoparticles has been grafted by a phosphoric di-ester type tensioactive called Beycostat NE.
This surfactant is a commercial product provided by CECA (Courbevoie, France) used to disperse inorganic pigments in aromatic and chlorinated oils (but insoluble in aliphatic solvents).

The magnetic nanoparticles prepared by the method briefly described above exhibit a rather broad polydispersity of diameters, ranging from 5 to 12 nm (from TEM images). The distribution of sizes of the magnetic cores is estimated quite accurately by measuring the magnetization curve with a vibrating sample magnetometer. This curve M(H) corresponds to giant paramagnetism and can be fitted by the first Langevin’s law weighted by a Log-normal distribution of the diameters. The typical synthesis of an ionic ferrofluid usually yields an average diameter $d_0^{\text{mag}}=7$ nm with a large standard deviation $\sigma=0.4$. Prior to the surfactant grafting, a size-sorting procedure is achieved on this raw acidic ferrofluid. As the iron oxide surface is coated simply by H⁺ protons providing electrostatic repulsions between the nanoparticles, a liquid-gas like phase separation is induced by an increase of ionic strength by adding excess HNO₃. The more concentrated liquid-like phase enriched in larger nanoparticles is separated from the supernatant by magnetic sedimentation. After two cycles of this fractionated phase separation, the second supernatant denoted S2-HNO₃ exhibits a distribution of diameters of parameters $d_0^{\text{mag}}=6.8$ nm and $\sigma^{\text{mag}}=0.24$, which remains unchanged after grafting with Beycostat NE to get the surfacted ferrofluid S2-CH₂Cl₂. Another characterization of sizes is given from the SANS curve of the pure dilute ferrofluid (inset of Figure 1). The oscillations of the well known form factor of spheres is here smeared out by polydispersity. We obtain from the Log-normal fit $d_0^{\text{SANS}}=7.6$ nm and $\sigma^{\text{SANS}}=0.25$, which corresponds to a slight shift of 0.8 nm towards larger diameters compared to the previous estimate of $d_0$. This gap is attributed to the core-shell magnetic structure of the nanoparticles, the atomic spins near the surface being disordered and thus not contributing to
II. AFM study of the two types of nano-composite objects

Unlike TEM which shows only the nanoparticles because of the low electron density of polymer compared to iron oxide, AFM is sensitive to both the copolymer and the nanoparticles. The magnetic micelles (system A) appear as filled bulky objects (Figure S1a) with a height of several diameters $d_0$ (Figure S1b).

In the case of the magnetic vesicles (system B), the nanoparticles are also locally confined, this time in a quite unusual way. The observed closed loops of nanoparticles around a central hole are quite strange indeed, looking like pieces of lace (Figure S2a). Nevertheless, this shape is extremely reproducible as the low magnification AFM picture shows many of such objects (Figure S2b). In this context, the uniform film visible around the objects is ascribable to the pure copolymer bilayer spreading onto the mica substrate (Figure S2c and Plot S2d). However, the apparent height of this film $\delta_{\text{AFM}}=1.2–1.4$ nm is much smaller than the true bilayer thickness (larger than $\delta_{\text{PB}}=14$ nm). Such an artifact of tapping mode AFM is well known for soft materials like block copolymer vesicles, which in addition are very sensitive to the force exerted by the tip and to the nature of the substrate. On the opposite, iron oxide is a very tough material. That explains why membrane portions containing nanoparticles lead to a measured height quite equal to the characteristic diameter $d_0$ of the nanoparticles. This observation confirms the result given by SANS that vesicles B doped with 1 equiv. of ferrofluid confine the magnetic nanoparticles within their membranes as monolayers. Besides, vesicles appear holey on the pictures presumably due to the strong adhesion of those soft shells on high energy surfaces (freshly cleaved mica for AFM or graphite for TEM). The ripping of the bilayers is indeed necessary to enable the spreading of initially curved
membranes onto strongly adhesive planar substrates (a sort of like pealed orange skins!). The opening of pores due to strong adhesion is a phenomenon known for lipid vesicles.\(^{[S7,S8]}\) By comparison, red blood cells also burst on adhesive substrates,\(^{[S9]}\) even though their membrane is thicker than pure lipid bilayers and exhibits a viscoelastic behavior due to a connected protein network. The pictures here show that pores due to strong adhesion occur for polymersomes as well, even though they require a larger surface tension to nucleate.\(^{[S10]}\)

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figureS1.png}
\caption{a) AFM picture of hybrid copolymer micelles loaded with magnetic nanoparticles. PB\(_{48}\)-b-PGA\(_{114}\) was combined with 1 mass equiv. of hydrophobic ferrofluid S2-CH\(_2\)Cl\(_2\). The scale bar measures 400 nm. b) Plot showing the cross-section along the solid line drawn on the image.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figureS2.png}
\caption{AFM pictures of magnetic membranes made of PB\(_{45}\)-b-PGA\(_{56}\) and 1 equiv. S2-CH\(_2\)Cl\(_2\). a) Typical lace shape (bar length is 500 nm). b) Wide scan field (bar is 1 \(\mu\)m). c) Tight view (bar is 150 nm). d) Height profile along the solid line.}
\end{figure}
III. Analysis of the anisotropy of the SANS patterns under magnetic field

In order to study the anisotropy quantitatively, the 2-d scattering patterns are averaged in angular sectors ±15° around the two perpendicular directions || and ⊥ defined by the magnetic field (see Figure 5b). After dividing by the known form factor $P_{\text{ferro}}(q)$ of the nanoparticles themselves, we obtain two different curves for the intra-aggregate structure factor $S_{\text{intra}}(q)$, which are plotted in Figure S3 for the field intensity $B=1050$ G and compared to the isotropic case $B=0$ G. The inset of Figure S3 is a plot of the same data as $\ln[S_{\text{intra}}(q)q^2]$ vs. $q^2$ in order to point out the Kratky-Porod regime of flat objects. Concerning now the anisotropy in both representations $S_{\text{intra}}(q)$ vs. $q$ and $\ln[S_{\text{intra}}(q)q^2]$ vs. $q^2$, no significant difference is observed between the measurements perpendicularly to the field (⊥) and the isotropic case without field, as could already be expected from the profile plots at constant $q$ of Figure 5b. The behavior is totally different in the direction parallel to the field (||), where $S_{\text{intra}}(q)$ is significantly modified compared to the isotropic case for values of $q$ vectors up to about $2 \times 10^{-2}$ Å⁻¹. Thus we deduce that the portions of membrane mostly affected by the magnetic field are those with their normal vector parallel to the field, in other words the magnetic poles. More precisely, two important features arise on the Kratky-Porod plot in the parallel direction: the slope (identified as $-\delta_{||}^2/12$) becomes nearly flat and the intercept (noted $\ln(I_{0||})$ as previously) is shifted to lower values by about 0.4 (in logarithmic units). Those observations are interpreted respectively as an important decrease of the apparent thickness $\delta_{||}$ near the poles and as an escape of the nanoparticles from membrane patches that cross the field direction. We remind once again that for both reasons of contrast and $q$ range, this evolution of $S_{\text{intra}}(q)$ denotes the variation of the spatial distribution of the magnetic nanoparticles thus probing indirectly the deformation of the vesicles.
Figure S3: Intra-aggregate structure factor $S_{\text{intra}}(q)$ of the magnetic nanoparticles imbedded in copolymer shells (system B 1eq) in the isotropic case at zero field (crosses) and under a magnetic field $B=1050$ G after angular regrouping in a sector ±15° around the direction of the field (circles, noted $||$) or ±15° around its normal direction (triangles, noted $\perp$). The inset shows the Kratky-Porod plot $\ln[S_{\text{intra}}(q) \times q^2]$ vs. $q^2$, which slope $-\delta^2/12$ (measured at $B=0$G) gives a membrane thickness $\delta=10$ nm.

References


